

MCRA Documentation

Release 10

Biometris, Wageningen University and Research

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USER GUIDE

Ι	Use	r Guide	3		
1	Introduction to MCRA				
	1.1	User interface	5		
		1.1.1 Welcome page	5		
		1.1.2 Registering	7		
		1.1.3 Logging in	8		
		Setting up two-factor authentication	8		
		Authenticator apps	10		
		Request an authenticator OR code	10		
		114 Main user interface	10		
	12	Data and calculation model	10		
	1.2	1.2.1 Modular design	10		
		1.2.1 Would design	14		
		1.2.2 Nonlined full did uncertainty dialysis	14		
		1.2.5 Variability diagnostics	14		
		1.2.4 Retain & Retain and uered approaches	13		
			18		
			18		
			19		
	1.0	Uncertainty due to modelling approach	19		
	1.3		19		
		1.3.1 Creating and uploading data files	20		
		1.3.2 Moving repositories and data sources	21		
		1.3.3 Repository access levels	21		
		1.3.4 Linking remote data repositories	21		
	1.4	Workspaces and actions	22		
		1.4.1 Workspace browser	24		
		1.4.2 Workspace overview page	24		
		1.4.3 Action area	24		
		Scoping: entity selection	26		
		1.4.4 Standard action area	29		
		Creating a standard action	29		
		Standard action reports	30		
		Converting a standard action to a full action	30		
		1.4.5 Action zip files	30		
	1.5	MCRA web and core	32		
		1.5.1 MCRA core	32		
		1.5.2 MCRA web	33		
		1.5.3 Running MCRA core using the command line interface	33		
	1.6	Results panel	33		
2 Examples					
	2.1	Cumulative dietary exposure assessment	35		
		2.1.1 Introduction	35		
		2.1.2 Preparation	35		

	2.1.3	Example 1
	2.1.4	Example 2
	2.1.5	Example 3
2.2	TDS-ba	sed exposure and risk assessment
	2.2.1	Standard actions
2.3	Aggrega	ate exposure assessment
	2.3.1	Introduction
	2.3.2	Preparation
	2.3.3	Example 1
	2.3.4	Example 2
2.4	Hazard	characterisations from PoDs
	2.4.1	Introduction
	2.4.2	Preparation
	2.4.3	Example 1
2.5	Health i	mpact estimates
	2.5.1	Introduction
	2.5.2	Preparation
	2.5.3	Example 1
2.6	Assessn	nent group membership probabilities 44
	2.6.1	Introduction
	2.6.2	Preparation
	2.6.3	Example 1
	2.6.4	Example 2

II Reference Manual

3	Mod	ules	
	3.1	Prima	ry entity modules
		3.1.1	Effects
			Effects from data
			Effects calculation
			Effects data formats
			Effects settings
			Effects as data
		3.1.2	Foods
			Foods from data
			Foods data formats
			Foods as data
			Food coding systems
			FoodEx2
			Food hierarchies
			Food unit weights
		3.1.3	Consumer products
			Consumer products from data
			Consumer products data formats
			Consumer products as data
		3.1.4	Non-dietary exposure sources
			Non-dietary exposure sources from data
			Non-dietary exposure sources data formats
			Non-dietary exposure sources as data
		3.1.5	Populations
			Populations from data
			Populations calculation
			Populations data formats
			Populations settings
			Populations as data
			Calculation of populations

	3.1.6	Responses
		Responses from data 76
		Responses data formats
		Responses as data
	3.1.7	Substances
		Substances from data
		Substances data formats
		Substances settings
		Substances as data
	3.1.8	Test systems
		Test systems from data
		Test systems data formats
		Test systems as data
3.2	Consu	mption modules
	3.2.1	Consumptions
		Consumptions from data
		Consumptions calculation
		Consumptions data formats
		Consumptions settings
		Consumptions tiers
		Consumptions uncertainty
		Consumptions as data
	3.2.2	Dust exposure determinants
		Dust exposure determinants from data
		Dust exposure determinants data formats
		Dust exposure determinants as data
	3.2.3	Soil exposure determinants
		Soil exposure determinants from data
		Soil exposure determinants data formats
		Soil exposure determinants as data
	3.2.4	Air exposure determinants
	0.20	Air exposure determinants from data
		Air exposure determinants data formats
		Air exposure determinants as data
	325	Consumer product exposure determinants 103
	0.2.0	Consumer product exposure determinants from data 103
		Consumer product exposure determinants data formats
		Consumer product exposure determinants as data
	326	Consumer product use frequencies 104
	5.2.0	Consumer product use frequencies from data
		Consumer product use frequencies data formats
		Consumer product use frequencies as data 108
	327	Food recipes 108
	5.2.7	Food recipes from data 108
		Food recipes data formats
		Food recipes as data
	328	Market shares
	5.2.0	Market shares from data 109
		Market shares data formats
		Market shares as data
		Market shares and brand lovalty 110
	320	Single value consumptions
	3.4.7	Single value consumptions from data 111
		Single value consumptions calculation 111
		Single value consumptions data formats
		Single value consumptions settings
		Single value consumptions as data
		Calculation of single value consumptions
		carculation of single value consumptions

2.2	0000	range modules 114
5.5		Concentration distributions
	3.3.1	Concentration distributions from late
		Concentration distributions from data
		Concentration distributions data formats
		Concentration distributions as data
	3.3.2	
		Concentration limits from data
		Concentration limits data formats
		Concentration limits as data
	3.3.3	Concentration models
		Concentration models calculation
		Concentration models settings
		Concentration models tiers
		Concentration models uncertainty
		Calculation of concentration models
	3.3.4	Concentrations
		Concentrations from data
		Concentrations calculation
		Concentrations data formats
		Concentrations settings 184
		Concentrations tiers 186
		Concentrations uncertainty 195
		Concentrations as data 195
	335	Deterministic substance conversion factors
	5.5.5	Deterministic substance conversion factors from data
		Deterministic substance conversion factors data formats
		Deterministic substance conversion factors as data
	336	Dust concentration distributions
	5.5.0	Dust concentration distributions from data 107
		Dust concentration distributions data formate 107
		Dust concentration distributions as data 107
	227	Soil concentration distributions
	5.5.7	Soil concentration distributions from data
		Soli concentration distributions from data
	220	Soli concentration distributions as data
	3.3.8	Consumer product concentrations
		Consumer product concentrations from data
		Consumer product concentrations data formats
		Consumer product concentrations as data
	3.3.9	Indoor air concentrations
		Indoor air concentrations from data
		Indoor air concentrations data formats
		Indoor air concentrations as data
	3.3.10	Outdoor air concentrations
		Outdoor air concentrations from data
		Outdoor air concentrations data formats
		Outdoor air concentrations as data
	3.3.11	Focal food concentrations
		Focal food concentrations from data
		Focal food concentrations data formats
		Focal food concentrations settings
		Focal food concentrations as data
	3.3.12	Food extrapolations
		Food extrapolations from data
		Food extrapolations data formats
		Food extrapolations as data
	3.3.13	Modelled foods
		Modelled foods calculation

Modelled foods settings	204
Calculation of modelled foods	204
3.3.14 Occurrence frequencies	205
Occurrence frequencies from data	205
Occurrence frequencies calculation	205
	205
Occurrence requencies data formats	203
Occurrence frequencies Settings	206
Occurrence frequencies as data	206
Calculation of occurrence frequencies	207
3.3.15 Occurrence patterns	207
Occurrence patterns from data	207
Occurrence patterns calculation	207
Occurrenza patterns data formats	207
	200
Occurrence patterns settings	209
Occurrence patterns tiers	210
Occurrence patterns as data	236
Calculation of occurrence patterns	236
3.3.16 Processing factors	236
Processing factors from data	236
Processing factors calculation	236
Drocessing factors data formate	230
	237
Processing factors settings	239
Processing factors tiers	239
Processing factors uncertainty	243
Processing factors as data	244
3.3.17 Single value concentrations	244
Single value concentrations from data	244
Single value concentrations calculation	244
Single value concentrations data formata	244
	244
Single value concentrations settings	245
Single value concentrations as data	245
Calculation of single value concentrations	246
3.3.18 Substance authorisations	246
Substance authorisations from data	246
Substance authorisations data formats	246
Substance authorisations as data	246
2.2.10. Substance automotions as data	240
5.5.19 Substance approvals	247
Substance approvals from data	247
Substance approvals data formats	247
Substance approvals as data	247
3.3.20 Substance conversions	247
Substance conversions from data	248
Substance conversions data formats	248
Substance conversions as data	249
3.3.21 Total diet study sample compositions	240
Tetal diet study sample compositions from data	249
Total diet study sample compositions from data	250
Total diet study sample compositions data formats	250
Total diet study sample compositions as data	250
3.3.22 Unit variability factors	250
Unit variability factors from data	250
Unit variability factors data formats	250
Unit variability factors	252
Unit variability factors tiers	252
	252
	200
Hyposure modules	255
3.4.1 Consumptions by modelled food	255
3.4.1 Consumptions by modelled food	255 255

3.4

	Calculation of consumptions by modelled food
3.4.2	Dietary exposures
	Dietary exposures calculation
	Dietary exposures settings
	Dietary exposures tiers
	Calculation of dietary exposures
3.4.3	Dust exposures
	Dust exposures calculation
	Dust exposures settings
	Calculation of dust exposures
3.4.4	Soil exposures
	Soil exposures calculation
	Soil exposures settings
	Calculation of soil exposures
3.4.5	Air exposures
	Air exposures calculation
	Air exposures settings
	Calculation of air exposures
3.4.6	Consumer product exposures
	Consumer product exposures calculation
	Consumer product exposures settings
	Calculation of consumer product exposures
3.4.7	Individuals
	Individuals calculation
	Individuals settings
	Calculation of individuals
3.4.8	High exposure food substance combinations
	High exposure food substance combinations calculation
	High exposure food substance combinations settings
	Calculation of high exposure food-substance combinations
3.4.9	Internal exposures
	Internal exposures calculation
	Internal exposures settings
	Calculation of internal exposures
3.4.10	Exposure mixtures
	Exposure mixtures calculation
	Exposure mixtures settings
	Calculation of exposure mixtures
3.4.11	Food conversions
	Food conversions calculation
	Food conversion settings
	Food conversions tiers
	Calculation of food conversions
3.4.12	Biological matrix concentration comparisons
	Biological matrix concentration comparisons calculation
	Biological matrix concentration comparisons settings
	Calculation of biological matrix concentration comparisons
3.4.13	Exposure biomarker conversions
	Exposure biomarker conversions from data
	Exposure biomarker conversions data formats
	Exposure biomarker conversions
	Exposure biomarker conversions as data
3.4.14	HBM analysis
	HBM analysis calculation
	HBM analysis settings
	Calculation of hbm analysis
3.4.15	HBM concentrations
	HBM concentrations from data

		HBM concentrations data formats
		HBM concentrations settings
		HBM concentrations as data
	3.4.16	Non-dietary exposures
		Non-dietary exposures from data
		Non-dietary exposures data formats
		Non-dietary exposures settings
		Non-dietary exposures uncertainty
		Non-dietary exposures as data 486
	3417	Single value dietary exposures 487
	5.1117	Single value dietary exposures calculation 487
		Single value dietary exposures data formats 490
		Single value dietary exposures settings 402
		Calculation of single value dietary exposures 403
	3/18	Single value non dietary exposures
	5.4.10	Single value non-dictary exposures from data
		Single value non-dictory exposures colculation 404
		Single value non-dietary exposures calculation
		Single value non-dietary exposures data formats
		Single value non-dietary exposures settings
		Single value non-dietary exposures as data
~ ~		Calculation of single value non-dietary exposures
3.5	Hazaro	d modules
	3.5.1	Active substances
		Active substances from data
		Active substances calculation
		Active substances data formats
		Active substances settings
		Active substances as data
		Calculation of active substances
	3.5.2	AOP networks
		AOP networks from data
		AOP Networks calculation
		AOP networks data formats
		AOP networks settings
		AOP networks as data
	3.5.3	Burden of disease
		Burden of disease from data
		Burden of disease data formats
		Burden of disease as data
	3.5.4	Dose response data
		Dose response data from data
		Dose response data data formats
		Dose response data settings
		Dose response data as data 516
	355	Dose response models 516
	0.0.0	Dose response models from data 516
		Dose response models calculation 516
		Dose response models data formats 517
		Dose response models settings 520
		Dose response models as data 521
		Calculation of dose response models 521
	356	Effect representations 521
	5.5.0	Effect representations from data 521
		Effect representations data formate
		Effect representations
		Effect representations as data 522
	257	Encer representations as uata
	5.5.7	Exposure response functions from deta
		Exposure response functions from data

		Exposure response functions data formats
		Exposure response functions
		Exposure response functions as data
	3.5.8	Hazard characterisations
		Hazard characterisations from data
		Hazard characterisations calculation
		Hazard characterisations data formats
		Hazard characterisations settings
		Hazard characterisations tiers 540
		Hazard characterisations as data 543
		Calculation of hazard characterisations 543
	359	Inter-species conversions 543
	5.5.7	Inter-species conversions from data 543
		Inter species conversions data formate 543
		Inter species conversions settings 544
		Inter species conversions as data 545
	2 5 10	Intro species conversions as data
	5.5.10	
		Intra species factors from data
		Intra species factors calculation
		Intra-species factors data formats
		Intra species factors settings
		Intra species factors as data
	3.5.11	Points of departure
		Points of departure from data
		Points of departure data formats
		Points of departure settings
		Points of departure as data
	3.5.12	Relative potency factors
		Relative potency factors from data
		Relative potency factors calculation
		Relative potency factors data formats
		Relative potency factors settings
		Relative potency factors as data
		Calculation of relative potency factors
3.6	In-silic	co modules
	3.6.1	Molecular docking models
		Molecular docking models from data
		Molecular docking models data formats
		Molecular docking models
		Molecular docking models as data 556
	3.6.2	OSAR membership models 556
		OSAR membership models from data 556
		OSAR membership models data formats 556
		OSAR membership models 558
		OSAR membership models as data 558
37	Kineti	c modules
5.7	271	Absorption factors 550
	5.7.1	Absorption factors from data 550
		Absorption factors data formats 550
		Absorption factors data formats
		Absorption factors as data
	272	Absorption factors as data
	5.1.2	Kinetic conversion factors
		Kinetic conversion factors from data
		Kinetic conversion factors data formats
		Kinetic conversion factors settings
	0 7 0	Kinetic conversion factors as data
	3.7.3	PBK model parametrisations
		PBK model parametrisations from data

		PBK model parametrisations data formats	65
		PBK model parametrisations settings 56	67
		PBK model parametrisations as data	67
		Available kinetic models	68
		27.4 DBK model definitions	06
		DPK model definitions from data	07
)/ 07
		PBK model definitions data formats	J/
		PBK model definitions as data	37
	3.8	Risk modules)7
		3.8.1 Environmental burden of disease)8
		Environmental burden of disease calculation)8
		Environmental burden of disease settings	98
		Calculation of environmental burden of disease	10
		3.8.2 Risks	11
		Risks calculation	11
		Risks settings	15
		Risks tiers 61	17
		Calculation of ricks 77	72
		2.9.2 Cincle value risks	23 NA
		Single value risks	24 24
			24
			28
		Single value risks tiers	30
		Calculation of single value risks	40
	~		
4	Stand	dard actions 80	57
	4.1	Chronic mixture risk assessment of metals	57
	4.2	Chronic cumulative exposure assessment PA	59
	4.3	Chronic cumulative exposure assessment PFAS	59
	4.4	Demo acute cumulative risk assessment	59
	4.5	Acute single substance dietary exposure assessment of carbofuran or chlorpyrifos	70
	4.6	Chronic single substance dietary exposure assessment of lead or atropine	71
	4.7	Demo Human Monitoring Analysis bisphenols	72
	4.8	TDS-based long term dietary exposure and risk assessment	73
	4.9	Long-term dietary exposure and risk of nickel for the Belgian population	73
	4.10	TDS-based long-term exposure and risk assessment of methylmercury for German children 87	74
		410.1 Introduction 87	74
		4 10 2 Standard action options	, . 75
		4.10.2 Standard action data	75
	1 11	Acute Cumulative Rick Assessment Craniofacial Alterations (EESA 2022)	75
	4.12	ELL aguta cumulative exposure assessment (2018) Tier Land Tier II	70
	4.12	EU acute cumulative exposure assessment (2018) Tier L and Tier I.	70
	4.15	EU Chilomic cumulative exposure assessment (2016) Tiel Land Tiel II	19 70
	4.14	EU Prospective Dietary Cumulative Risk Assessment (2023) $\dots \dots \dots$	/9
	4.15	EU Prospective Dietary Cumulative Risk Assessment (2024)	51
	4.16	EU Retrospective Dietary Cumulative Risk Assessment (2023)	51
	4.17	EU Retrospective Dietary Cumulative Risk Assessment (2024)	32
	4.18	Risk steatosis from imazalil	33
	4.19	Training prospective risk assessment acute Tier II	34
	4.20	Training prospective risk assessment chronic Tier II	35
	4.21	Training substance prioritisation acute neuro	86
_	_		
5	Туре	and Unit definitions 88	89
	5.1	Adjustment factor distribution method types	39
	5.2	Age alignment method	8 9
	5.3	Application amount unit	90
	5.4	Assessment group membership calculation methods	90
		Panchmark response type	91
	5.5		
	5.5 5.6	Biological matrix	93

5.8	Biomarker conversion distribution type
5.9	BoD Indicator
5.10	Body weight unit
5.11	Boolean type
5.12	Burden of disease approach
5.13	Cluster method type
5.14	Combination method membership info and PoD presence types
5.15	Concentration limit value type
5.16	Concentration model types
5.17	Concentration unit
5.18	Concentration value type
5.19	Concentration volume unit
5.20	Consumer product application amount distribution types
5.21	Consumer product concentration uncertainty distribution types
5.22	Consumption intake unit
5.23	Consumption unit
5.24	Consumption value type
5.25	Covariate model types
5.26	Dietary exposures details level types
5.27	Dose response model type
5.28	Dose unit
5.29	Dust adherence amount uncertainty distribution types
5.30	Dust availability fraction uncertainty distribution types
5.31	Dust body exposure fraction uncertainty distribution types
5.32	Dustingestion uncertainty distribution types
5.33	Effect metric
5.34	Estimates nature types
5.35	Exposure approach types
5.36	Exposure calculation method
5.37	Exposure events and dose generation
5.38	Exposure grouping method
5.39	Exposure method types
5.40	Exposure path type
5.41	Exposure response type
5.42	Exposure route
5.43	Exposure source
5.44	Exposure type
5.45	Expression type
5.46	External exposure unit
5.47	Focal commodity replacement method types
5.48	Function types
5.49	Harvest application type
5.50	Hazard characterisation type
5.51	Hazard characterisations calculation method types
5.52	Hazard dose imputation method types 921
5.53	Health effect types
5.54	Individual generation method
5.55	Individual generation method
5.56	Individual generation method
5.57	Individual property type
5.58	Individual subset types
5.59	Intake model types
5.60	Internal model type
5.61	Isced type
5.62	Job task type
5.63	Kinetic conversion factor uncertainty distribution types
5.64	Left-censored data handling methods
5.65	Mean value correction types

	5.66	Measurement result type	928
	5.67	Missing value imputation method types	929
	5.68	Modelled foods calculation source types	929
	5.69	Month type	930
	5.70	Multiple substance handling method types	930
	5.71	Network analysis type	931
	5.72	Nondetect imputation method types	931
	5.73	PBK model compartment type	931
	5.74	PBK model parameter distribution types.	933
	5.75	PBK model parameter type	933
	5.76	PBK model species type	934
	5 77	PBK simulation methods	934
	5 78	Point of departure type	934
	5 79	Point of departure types	335
	5.80	Population alignment method	335
	5.80	Dopulation characteristic distribution types	135
	5.01	Depulation characteristic distribution types.	126
	J.02	Population characteristic type	200
	5.85		230
	5.84		931
	5.85	Response type	937
	5.86	Risk characterisation ratio	937
	5.87	Riskmetric calculation types	938
	5.88	Selected exposure routes	938
	5.89	Sex type	938
	5.90	Single value dietary exposures calculation method types	939
	5.91	Single value risk calculation method types	939
	5.92	Soilingestion uncertainty distribution types.	940
	5.93	Standardise blood methods	940
	5.94	Standardise/normalise urine concentration method	940
	5.95	Substance group selection method types	941
	5.96	Substance translation allocation method types	942
	5.97	Target dose selection method types	943
	5.98	Target level type	943
	5.99	Test system type	944
	5.100	Testing method types	944
	5.101	Time unit	945
	5.102	Transform types	945
	5 103	Uncertainty types	946
	5 104	Unit variability correlation types)46
	5 105	Unit variability model types	240 247
	5 106	Unit variability types) //7
	5 107	Unit weight value type)/Q
	5 100	Value qualifier	140
	5 100	Value qualifier	140
	5.109		740
6	Appli	cation Programming Interface (API)	949
7	Com	nand Line Interface (CLI))51
	7.1	Introduction)51
	7.1	Action template structure	951
	7.2	Create a new action template) 57
	7 /	Run an action	,52)52
	7. 4 7.5	Output files and folder structure	,55)5/
	1.5		,54
8	Appe	ndices	955
	8.1	NOEL collection of Munro et al.	955
	8.2	Box-Cox power transformation	955
	8.3	Gauss-Hermite Integration)55
		8.3.1 One-dimensional Gauss-Hermite integration)55

		8.3.2	Two-dimensional Gauss-Hermite integration	956
		8.3.3	Maximum likelihood for the LNN model with two-dimensional Gauss-Hermite integration	956
	8.4	Distribu	itions	956
		8.4.1	Uniform distribution	956
		8.4.2	Lognormal distribution	957
		8.4.3	Beta distribution	957
		8.4.4	InverseUniform distribution	957
9	Gloss	ary		959
III	Bi	bliogra	aphy	967
10	Publi	cations u	using MCRA	969
11	Color	hon		981
11	11 1	Contrib	utors to MCR A	981
	11.1	Contino		701
12	Chan	ge Log		983
	12.1	Version	10.2.0 (2025-07-07)	983
		12.1.1	Added	983
		12.1.1	Fixed	083
	12.2	12.1.2	10.1.10 (2025.05.09)	002
	12.2		10.1.10(2023-03-08)	900
		12.2.1	Added	983
		12.2.2	Changed	983
		12.2.3	Fixed	983
	12.3	Version	10.1.9 (2025-04-28)	983
		12.3.1	Added	983
		12.3.2	Changed	984
		1233	Fixed	084
	12.4	Varcian	10.1.9 (2025.04.09)	004
	12.4		$10.1.6 (2023-04-06) \dots \dots$	904
		12.4.1	Changed	984
		12.4.2	Fixed	984
	12.5	Version	10.1.7 (2025-03-24)	984
		12.5.1	Added	984
		12.5.2	Changed	984
		12.5.3	Fixed	984
	12.6	Version	10 1 6 (2025-03-04)	985
	12:0	12.6.1	Added	085
		12.0.1	Changed	005
		12.0.2	Changeu	905
		12.6.3		985
	12.7	Version	10.1.5 (2025-02-10)	985
		12.7.1	Added	985
		12.7.2	Changed	985
	12.8	Version	10.1.4 (2025-01-31)	985
		12.8.1	Added	985
		12.8.2	Changed	085
		12.0.2	Eived	005
	12.0	12.0.3	FIXEU	905
	12.9	Version	$10.1.3 (2024-12-13) \dots \dots$	986
		12.9.1	Added	986
		12.9.2	Fixed	986
	12.10	Version	10.1.2 (2024-11-08)	986
		12.10.1	Added	986
		12 10 2	Changed	986
		12.10.2	Fived	006
	10.11	12.10.3	F1AUU	700
	12.11	version	10.1.1 (2024-09-20)	98/
		12.11.1	Added	987
		12.11.2	Changed	987
		12.11.3	Fixed	987

12.12 Version 10.1.0 (2024-08-15)
12.12.1 Added
12.12.2 Changed
12.12.3 Fixed
12.13 Version 10.0.15 (2024-06-21) 985
12 13 1 Changed 98
12 14 Version 10 0 14 (2024-06-18) 98
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12.15 Version 10.0.13 (2024.05.28)
12.15 Version 10.0.15 (2024-05-26)
12.15.1 Added
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12.10 Version 10.0.12 (2024-03-21)
12.10.1 Added
12.16.2 Changed
12.10.3 Fixed
12.1/ Version 10.0.11 (2024-04-12)
12.17.1 Added
12.17.2 Changed
12.17.3 Fixed
12.18 Version 10.0.10 (2024-03-07)
12.18.1 Changed
12.18.2 Added
12.19 Version 10.0.9 (2024-02-27)
12.19.1 Added
12.19.2 Changed
12.19.3 Fixed
12.20 Version 10.0.8 (2024-02-12)
12.20.1 Changed
12.21 Version 10.0.7 (2024-02-02)
12.21.1 Added
12.21.2 Changed
12.21.3 Fixed
12.22 Version 10.0.6 (2024-01-26)
12.22.1 Fixed
12.23 Version 10.0.5 (2023-12-21)
12.23.1 Changed
12.24 Version 10.0.4 (2023-12-18)
12.24.1 Added
12.24.2 Changed
12.24.3 Fixed
12.25 Version 10.0.3 (2023-12-01)
12.25.1 Added
12.25.2 Changed
12.25.3 Fixed
12.26 Version 10.0.2 (2023-10-26)
12.27 Version 10.0.1 (2023-10-23)
12.27.1 Added
12.27.2 Changed
12.27.3 Fixed
12.28 Version 10.0.0 (2023-06-23)
12.28.1 Changed
12.28.2 Fixed
12.29 Version 9.2.10 (2023-06-01)
12.29.1 Changed
12.29.2 Fixed
12.30 Version 9.2.9 (2023-05-24)
12.30.1 Added
12.30.2 Changed

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12.37.3 Fixed	995
12.57.2 Changed	993
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12.36.2 Changed	995
12.30 Version 7.2.3 (2023-01-10)	995
12.35.5 1100	005
12.35.2 Changed	005
12.35.1 Thatea	994
12.35 (Clober 2.2-02-02-03)	994
12.37.5 Γ 100 Γ 12.37.5 Γ 1	004
12.34.2 Fixed	00/
12.37.1 Autou	00/
12.34 Version 9.2.5 (2025-02-10)	994 00/
12.55.5 Fixed	994
12.33.2 Changed	993
12.33.1 Added	993
12.33 Version 9.2.6 (2023-03-10)	993
12.32.3 Fixed	993
12.32.2 Changed	993
12.32.1 Added	993
12.32 Version 9.2.7 (2023-04-03)	993
12.31.2 Fixed	993
12.31.1 Added	993
12.31 Version 9.2.8 (2023-04-25)	993
12.30.3 Fixed	992

Bibliography

Index

1003

Reference and user manual for MCRA 10 (version 10.2.0).

Part I

User Guide

INTRODUCTION TO MCRA

Humans are exposed to a mixture of multiple chemicals via food intake, inhalation and dermal contact. The risk to health that may result from this depends on the effects of different chemicals in the mixture and how they combine.

MCRA is the model and data toolbox developed in the EuroMix project. It implements methods for exposure, hazard and risk assessment, following guidelines from a.o. the Joint Research Centre (JRC) of the European Commission and the European Food Safety Authority (EFSA). The toolbox should provide computational tools for future risk management decisions on the safety of chemicals in mixtures to be taken by the European Commission and the Codex Alimentarius.

MCRA is a collection of data and models. The system consists of modules that are arranged in eight categories according to a *modular design*. See *Modules overview*.

Each module represents a certain type of data, which can be computed from data provided by other (sub)modules, or the data may be obtained from a dataset selected from the *data repository*. Likewise, each module may be of interest by its own merit, or may just be required as a sub-part of larger calculations. The modular design of MCRA reveals a network of data and models, and shows how data of types and from various sources can be combined in overarching modules. The most overarching module is *Risks or health impact estimates*. The toolbox allows the user to start in any of the modules in the modular design for performing calculations.

For each module, an *action* can be created to configure and run the module. For data modules, such as the *concentrations module*, such an action comprises specifying the dataset, specifying the scope (i.e., *foods* of interest, *substances* of interest, etc.), and perhaps specifying specific selections or model settings for data manipulations (e.g., *imputation of water concentrations* in the concentrations module). For calculation modules, when calculating the data of the module based on other data, configuration of an action comprises specification of the model settings and selection of the calculation inputs, which is data provided by other (sub-)modules. While running an action in MCRA, the module produces output of its associated data type (which can be used as input for other modules), and a report will be generated of the selected data, the selection and model settings, and the module and all intermediate (i.e., sub-modules) results.

1.1 User interface

This section describes the user interface of the MCRA Web Application.

The following sections give an overview of the general functionality of MCRA, broken down into subjects.

1.1.1 Welcome page

The MCRA welcome page contains general information on MCRA, links to the documentation and login and registration functionality.

In the top right corner the 'person' dropdown menu is expanded and contains the 'Log in' and 'Register' menu items. At the bottom of the page the current version of MCRA is shown.

Exposure, Hazard & Risk Assessment	
	🔔 Log in
	+ Register
Welcome to MCRA	
Chemical exposure, hazard and risk assessment	
On a daily basis, people are exposed to multiple chemicals via food intake, inhalation and derm health resulting from this exposure depends on the effects of the different chemicals in the mix MCRA stands for Monte Carlo Risk Assessment . It is a web-based platform containing various assess these health risks for specific populations in various scenarios.	al contact. The risk to human ture and how they combine. models that users can use to
In MCRA, more than 50 modules are available to address all major areas of risk assessment, inchazard characterisation, exposure assessment and risk assessment. MCRA contains models for regulatory methodologies of the European Commission and the European Food Safety Authority includes novel scientific models that could improve or refine future risk assessment.	cluding hazard identification, ollowing the guidelines and y (EFSA). Besides this, it also
MCRA was and is being developed in multiple projects, including national funded projects, parts (2015-2025) and EU projects ACROPOLIS (2010-2013), EuroMix (2015-2019), FNS Cloud (2019)	nership between EFSA, RIVM
2029).	-2023), and PARC (2022-
2029). MCRA documentation Publications and reports using MCRA	-2023), and FARG (2022-
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2029). MCRA documentation Publications and reports using MCRA MCRA account Use of the MCRA web-platform requires an active account. An account can be requested by filli Register for an account Do you already have an account? Log in here. Contact: MCRA Support, National Institute for Public Health and the Environment (RIVM). MCRA is developed by Wageningen University & Research, Biometris for RIVM and EFSA (2007 - 2023)	ng in the registration form.



1.1.2 Registering

If you don't have an MCRA account, the registration procedure is as follows.

In the MCRA welcome screen, click on the 'Register for an account' button, which opens the following screen.

The screen consists of a number of sections which contains several required and optional items to complete before the register button is activated at the bottom of the input form (you need to scroll down on the page to see all options).

lease complete the following sections to register for your MCRA account.	
/lake sure you fill the required fields and click the Register button at the bottom of the lage, after which you will receive an email to confirm your registration.	
'our details	
lease provide some details about you, your affiliation and the email address we can u o send a verification code for your account.	ise
Full name *	
Affiliation *	
Please select your affiliation type *	*
Email *	
four account	
lease choose your preferred user account name and password.	
User name *	
Password *	
Conference on a second at	
commin passworg *	
Show password - Show password 'our password should consist of minimal 8 lower and upper case characters and at le- ine number and one of the following special characters: 1, @, #, \$, %, ^, & or *.	ast
Show password - Show password 'our password should consist of minimal 8 lower and upper case characters and at le ine number and one of the following special characters: 1, @, #, \$, %, ^, & or *. Ise of MCRA	ast
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The following fields are required:

- Full name: your full name
- Affiliation: please provide the organisation you are affiliated with
- Affiliation type: select an option from the drop-down list to specify the type of your organisation
- Email: please provide a valid email address, if you use your email address as your user name, please repeat it here
- User name: choose a user name to use to log in to MCRA, you may use your email address also. The user name cannot contain spaces only upper and lower case letters, numbers, or the characters @ (at), - (dash), . (dot) or _ (underscore) are allowed with a length between 8 and 50 characters.
- Password: provide a secure password, it should consist of minimal 8 lower and upper case characters and at least one number and one of the following special characters: !, @, #, \$, %, ^, & or *.

You can type a short message on how you wish to use MCRA in the text field under 'Use of MCRA'

Optionally you can specify whether you want to use MCRA specifically for the EU-PARC project.

You are required to accept MCRA's Terms of Use to be able to register.

For security reasons, please type the characters that show up in the image at the bottom of the page, which consists of all lowercase letters and digits.

Clicking the 'Register' button submits the filled form to MCRA. You will receive an email at the email address you provided, containing a link to confirm your registration. This will verify your email account and confirm your registration request. The screen will show a verification message, as shown below.

Your registration request still needs to be approved by RIVM. You will receive another email when your account has been approved, after which you can log in to MCRA with your new account.

Registration e-mail verification

Thank you, MCRA Demonstration Account, for completing your registration

Your request to access this version of MCRA is now pending approval. You will be notified by e-mail when access to this version of MCRA has been granted and you can log in.

Note: It may take some time for us to approve your account, so we ask for your patience. If you haven't received an account approval e-mail in due time, please check your 'spam' folder. Otherwise send an e-mail to MCRA support with your details to approve your account.

Go to start page



Figure 1.3: MCRA registration request has been verified

Log in					
	Username *				
-	mcra-demo				
01	Password *				
	Login				
	Forgot password	Create an account			

Figure 1.4: The MCRA login dialog

Two-factor authentication					
MCRA requires the use of two-factor authenticatior your user name and password. To set up two-factor	n to comply to the latest security standards. T r authentication for your MCRA account, plea	his will require entering a 6-digit code in addition to se follow the next few steps.			
Step 1					
Two factor authentication requires the use of an app on your mobile device. The following apps are recommended for this, but any <u>time-based</u> <u>one-time password</u> generator will do.	Microsoft Authenticator	Google Authenticator			
Step 2					
Please click the button to the right to request a personal QR code which allows you to add your MCRA account to the Authenticator app.	Request Authenticator QR code				
Step 3					
Enter the 6-digit code provided by your authenticator app in the input box to the right and click 'Validate':	• Type the 6-digit code here *				
	Validate				

Figure 1.5: Set up two-factor authentication

Authenticator apps

Two factor authentication requires the use of an app on your (mobile) device. The following authenticator apps are commonly used to log in to systems which require two-factor authentication.

Use one of the links below to navigate to the website where you can install the app of your choice. Note that there are many other apps offering time-based one-time password authentication, simply search for two-factor (2FA) apps in your favourite search engine.

Google Authenticator



Microsoft Authenticator



Windows app: 2Fast Authenticator



Request an authenticator QR code

When you request the authenticator QR code, the following screen appears:

Here you need to type your password and a verification code. Click the 'Send' button to finish the request. You will receive an email containing your personal QR code

The image above shows an example of a QR code email. The Authenticator app allows you to add an account and scan the QR code. Alternatively, a readable code is also given as a way to add the account to your authenticator app.

After the authenticator has been set up correctly, you can return to the MCRA login screen and log in using your user name and password, and continue to enter the 6-digit code you get from your authenticator app to access MCRA.

If you should somehow lose your QR code, you can always request to send it to your email address using the 'I lost my authenticator code' button on the two-factor authentication page.

1.1.4 Main user interface

The MCRA main page contains general information on MCRA, links to the data, workspaces and documentation.

1.2 Data and calculation model

1.2.1 Modular design

The modular design distinguishes between three types of modules: primary entity modules, data modules, and calculation modules. For an overview see *Modules*.

- The primary entity modules are data modules determining the scope of the assessments in MCRA. That is, in each assessment, the scope specifies the *foods*, *substances*, *effects*, *populations*, *responses*, and/or *test systems* that are of interest.
- The data modules give summaries of the available data which depend on (some of) the primary entities. For example *consumptions* data.

Two Factor Authentication QR Code				
Please type your password and the text in the image to e-mail with your two-factor authenticator QR code.	receive an			
user name mcra-demo				
Please type your password *				
Show password				
hoj ⁿ b v c i	n			
Type the text in the image above * hbjnbvcm				
C New image	Send			

Figure 1.6: Request your personal two-factor QR code

From: No-Reply mcra-test <noreply.mcra-test@wur.nl> Sent: donderdag 2 november 2023 10:46 To: Subject: MCRA - Authenticator QR Code

Your MCRA two-factor authenticator QR code

You requested the Authenticator QR code for account mcra-demo (MCRA Demonstration Account)

Using your Authenticator app, please scan the following QR code to add 'MCRA':



Alternatively, you may enter the following code into your Authenticator app:

DEMODEMODEMO123456789DEMODEMODEMO

Figure 1.7: QR code email example

	MCRA Exposure, Hazard & Risk Assessment
V	Velcome to MCRA
С	hemical exposure, hazard and risk assessment
O h N a	In a daily basis, people are exposed to multiple chemicals via food intake, inhalation and dermal contact. The risk to human ealth resulting from this exposure depends on the effects of the different chemicals in the mixture and how they combine. ICRA stands for Monte Carlo Risk Assessment . It is a web-based platform containing various models that users can use to ssess these health risks for specific populations in various scenarios.
lr h re ir	MCRA, more than 50 modules are available to address all major areas of risk assessment, including hazard identification, azard characterisation, exposure assessment and risk assessment. MCRA contains models following the guidelines and egulatory methodologies of the European Commission and the European Food Safety Authority (EFSA). Besides this, it also includes novel scientific models that could improve or refine future risk assessment.
N (2 2	1CRA was and is being developed in multiple projects, including national funded projects, partnership between EFSA, RIVM 2015-2025) and EU projects ACROPOLIS (2010-2013), EuroMix (2015-2019), FNS Cloud (2019-2023), and PARC (2022-029).
	MCRA documentation Publications and reports using MCRA
ι	
Y E ta	ou can specify your models, such as a dietary exposure assessment, within actions that are organised in workspaces . ach action is of a module type and contains selected data and settings. After specifying data and settings, the modelling ask can be started. The output report contains concise sections of main results and detailed drilldown information.
Y E ta T d	Source for the second s
Y E ta T d	You can specify your models, such as a dietary exposure assessment, within actions that are organised in workspaces. ach action is of a module type and contains selected data and settings. After specifying data and settings, the modelling ask can be started. The output report contains concise sections of main results and detailed drilldown information. he data used in the actions is organised in the data repository. Users have their own private data repository for uploading ata. In addition, shared repository folders are available used for sharing data among user groups. or more information on using MCRA consult the documentation pages.
Y E ta d F	Tou can specify your models, such as a dietary exposure assessment, within actions that are organised in workspaces. ach action is of a module type and contains selected data and settings. After specifying data and settings, the modelling ask can be started. The output report contains concise sections of main results and detailed drilldown information. the data used in the actions is organised in the data repository. Users have their own private data repository for uploading ata. In addition, shared repository folders are available used for sharing data among user groups. or more information on using MCRA consult the documentation pages.
Y E ta f F C	DSING WCKA You can specify your models, such as a dietary exposure assessment, within actions that are organised in workspaces. ach action is of a module type and contains selected data and settings. After specifying data and settings, the modelling ask can be started. The output report contains concise sections of main results and detailed drilldown information. he data used in the actions is organised in the data repository. Users have their own private data repository for uploading ata. In addition, shared repository folders are available used for sharing data among user groups. or more information on using MCRA consult the documentation pages. Image: Contains Consult the documentation pages. Image: Contact: MCRA Support, National Institute for Public Health and the Environment (RIVM).
Y E ta d F C	Sing WCKA Fou can specify your models, such as a dietary exposure assessment, within actions that are organised in workspaces. fach action is of a module type and contains selected data and settings. After specifying data and settings, the modelling ask can be started. The output report contains concise sections of main results and detailed drilldown information. he data used in the actions is organised in the data repository. Users have their own private data repository for uploading ata. In addition, shared repository folders are available used for sharing data among user groups. or more information on using MCRA consult the documentation pages. Image: The Workspace models with the for Public Health and the Environment (RIVM). ICRA is developed by Wageningen University & Research, Biometris for RIVM and EFSA (2007 - 2023).

Figure 1.8: MCRA main user interface

• The calculation modules perform calculations on input data to produce data on another type, as specified by the module name. E.g. the *dietary-exposures* calculation module calculates dietary exposures from consumption and occurrence data. Some calculation modules can also act as a data module, in which case the data are directly specified rather than calculated. Examples are, the *relative potency factors* module: relative potency factors can be supplied as such (*Data*) or computed based on *hazard characterisations* (*Compute*); the *single value consumptions* module: Large Portions can be supplied as such (*Data*) or computed based on consumption distribution data of a population (*Compute*).

Risks / Relative potency factors	► €	
Relative potency factors		0
🗮 Use data 🏾 🏟 Compute		

Figure 1.9: Relative potency factors supplied as data or computed based on hazard characterisations.

1.2.2 Nominal run and uncertainty analysis

Within MCRA two types of simulation runs are distinguished: the nominal run and the uncertainty analysis loop.

The nominal run represents a single simulation which is aimed to compute the most likely, unbiased estimates for the specified model. E.g., when a *dietary exposure assessment* is requested, in the nominal run a single exposure distribution is estimated using nominal values for all data and parameters.

In the *uncertainty analysis* loop, each simulation run is repeated a large number of times. Each run starts with a different scenario using data obtained with bootstrapping, parametric resampling and/or re-calculation of uncertain values. As a result, a large number of uncertain dietary exposure distributions is estimated which are used to estimate uncertainty limits (p5, p95).

Running a nominal run first has the advantage that the user may evaluate these modelling results before doing the final analysis. The model specification of rather complex simulation models and the corresponding output results are evaluated to detect any errors or misspecifications. Possible errors in the data and/or model settings are identified and corrected. In the final exposure assessment, the uncertainty analysis is included and the uncertainty of estimates is assessed.



Figure 1.10: Uncertainty analysis settings.

1.2.3 Variability diagnostics

In MCRA, the nominal run might be followed by an uncertainty analysis to assess the uncertainty limits (e.g. 2.5 and 97.5%) of the nominal percentiles (e.g. p50, p95, p99, p99.9, p99.99) of the exposure or risk distribution. For these percentiles, the nominal run of an acute assessment consists of preferably 100.000 iterations, the uncertainty analysis preferably of 100 runs with 10.000 iterations each. Note that for the percentile p99.99 the minimal number of iterations should be 10.000. Likewise, to estimate uncertainty limits of 2.5 and 97.5%, the minimal number of bootstrap runs should be 100.

In general, the number of iterations and bootstrap runs will be restricted due to limited computational resources or simulation runs that are time consuming. MCRA offers a diagnostic tool to visualize whether the estimated percentiles and uncertainty limits are stable or vary due to small simulation runs with limited number of iterations and uncertainty runs.

The diagnostic tool focus on the stability of the percentiles or, re-frasing, quantify 1) the amount of Monte Carlo variability and 2) the amount of variability due to resampling e.g. consumption and monitoring data or others sources of uncertainty. By quantifying both quantities, the influence of both sources of variability on the estimated value of the percentiles is assessed.

The diagnostics are displayed in a number of graphs (as many as the number of specified percentiles). For each percentile, the graph is used to draw inference about the optimal number of MC-iterations, the number of uncertainty runs and the number of iterations in each uncertainty run.

In the section below it is assumed that the nominal run consists of 100.000 iterations and uncertainty (95% confidence) is assessed with 100 uncertainty runs of 10.000 iterations each.

To make inference, the set of nominal (Monte Carlo) values is split in 2 samples of 50.000 iterations each, 4 samples of 25.000 each, 8 samples of 12.500 each etc. By doing so, we get n partitions of samples and in each partition we have 2^n samples of size $100.000/2^n$. In each partition, the percentiles of the available samples are estimated and the standard deviation of the percentiles. So in partition n = 1, the estimate of the standard deviation is based on 2 percentiles derived from samples of size 50.000; in partition n = 2, the estimate of the standard deviation is based on 4 percentiles derived from samples of size 25.000, etc. The estimated standard deviations are plotted against the number of MC-iterations per sample of each partition. It is expected that the standard deviation decreases as a function of sample size, so for larger sample sizes MC-variability decreases. For each standard deviation the 90% confidence limits are calculated.

A similar procedure is applied to the 100 uncertainty runs (of size 10.000). In each uncertainty run the percentiles are estimated. Then, in partition n = 1, percentiles are estimated on the first 10.000/2 = 5000 iterations of each sample; in partition n = 2 percentiles are estimated on the first 10.000/4 = 2500 iterations of each sample, etc. Then standard deviations of the percentiles of the partitions of 100×10.000 , 100×5000 , 100×2500 , etc are estimated and plotted against the number of iterations in each partition. It is expected that the standard deviation decreases as a function of sample size.

In Figure 1.12, the estimates and uncertainty limits are displayed. In Figure 1.11, the diagnostics are dlotted

- Red dots: standard deviation (sd) of percentile estimates between subsets of simulations. Error bars indicate parametric 90% confidence interval for the sd.
- Blue dots: standard deviation of percentile estimates between uncertainty iterations of subsets of simulations. The red square indicates the specified number of iterations (10000) of an uncertainty run.
- Each blue dot represents the standard deviation of 100 percentiles

For percentages, p50, p90, p95 and p99 the curves of the nominal estimates are smooth. For the extreme percentiles, p99.9 and p99.99, patterns do not monotonically decrease and vary, indicating that percentile estimates are variable. Also the 90% confidence interval around the sd's are much larger than for the smaller percentiles. An general conclusion is that percentiles and uncertainty limits for p50, p99, p95 and p99 are stable. The percentile of the p99.99 is unstable meaning that running the simulation with a different initialisation seed estimates will vary. The percentile p99.9 is an intermediate case.

The boxplots for uncertainty show the p25 and p75 as edges of the box, and p2.5 and p97.5 as edges of the whiskers. The reference value is indicated with the dashed black line, the median with the solid black line within the box. Outliers are displayed as dots outside the wiskers.

1.2.4 Retain & Refine and tiered approaches

A basic idea of Retain & Refine is that entities (e.g., substances) can be handled in different ways (more or less refined) while still being considered together in the same risk assessment (retain). We refer to such different approaches as tiers.

In the modular design, a tier is defined as a specific set of settings for a module or a group of modules. Tiers differ in many aspects, and there is no single dimension to rank tiers as low vs. high. In risk assessment, typical tiers contrast deterministic to probabilistic approaches, conservative to realistic approaches, approaches using restricted data to



Figure 1.11: Variability diagnostics for percentiles p50, p90, p95, p99, p99.9 and p99.99.

Percentage	Exposure (µg/kg bw/day)	Median (p50)	Uncertainty lower bound (p2.5).	Uncertainty upper bound (p97.5).
50.00	0.2278	0.2026	0.1837	0.2395
90.00	0.6236	0.56	0.4896	0.653
95.00	0.82	0.7339	0.6417	0.8664
99.00	1.339	1.217	1.035	1.438
99.90	2.439	2.128	1.693	2.743
99.99	3.752	3.13	2.411	4.777

Figure 1.12: Exposure estimates for percentiles p50, p90, p95, p99, p99.9 and p99.99 with 95% confidence limits



Figure 1.13: Boxplots for summary statistics for percentiles p50, p90, p95, p99, p99.9 and p99.99 with 95% confidence limits

approaches using more extensive data, and approaches using different degrees of model complexity. For each of the modules of MCRA, as many tiers are implemented as considered useful for the practice of risk assessment.

Each calculation in the modular design may involve multiple, nested, calculations of sub-modules. A *risk* (or health impact) assessment builds on an *exposure assessment* and a *hazard assessment*, the exposure assessment builds on a *dietary* and a *non-dietary exposure* assessment, the dietary exposure assessment builds on a *consumption assessment* and an *occurrence assessment*, etc. Tiers can be defined at each node of the assessment network.

Each calculator has as a main output entities that can be specified to have different tiers (tiered entities). For example, in a *hazard assessment*, some substances may be assessed using a tier 'Hazard Dose from dose-response data', other substances may be assessed using a tier 'TTCx100' or 'sample from general NOAEL distribution x100' (which only requires knowledge of the Cramer class of the substance). As another example, in dietary exposure assessment some food-substance combinations may be recognised as risk drivers for which a more complex approach (e.g. probabilistic modelling) is required, whereas a simpler approach (e.g. *deterministic modelling*) may be sufficient for all other food-substance combinations. So in this case the tiered entity is 'food-substance'. A typical risk assessment will start at a tier that is simple to perform for all tiered entities (potential risk drivers). Note that, based on data availability and ease of application, the initial assessment can already include more complex elements, such as probabilistic modelling. If the initial calculations produce risk estimates that do not exclude concern, refinement of the modelling for the perceived risk drivers is useful for checking whether this concern is real.

1.2.5 Uncertainty

Uncertainties arise in different forms in many of the models and data of MCRA.

- uncertainty in the data values (e.g., uncertain NOAELs, uncertain RPFs, or uncertain processing factors),
- uncertainty due to limited data (e.g., a limited number of food samples),
- uncertainty due to a lack of data (e.g., missing concentration data for some foods/substances or missing processing factors),
- uncertainty of the models, (e.g., due to a lack of detail).

MCRA offers the following options to handle uncertainty:

- for many types of data, the possibility to provide data including quantifications of uncertainty,
- imputation methods for filling in missing data in various types of models, and
- a generic uncertainty analysis method that providing uncertainty estimates of the modelling results for many of the modules, which are based on bootstrapping, parametric resampling, and/or re-calculation on all sub-modules for which this is possible.

Uncertainty due to limited sampled data

For some type of data, e.g., processing factors, it is possible to not only provide nominal estimates of the data values, but also to provide quantified estimates of the uncertainties of these values. Occasionally, quantifications of the uncertainties of these estimates are not available. MCRA provides the possibility to work with both quantified and unquantified uncertainties: include uncertainties in a quantitative uncertainty analysis when available, or, when not available, use nominal estimates, followed by an offline qualitative uncertainty analysis.

Uncertainties of data values are available in different forms. For some data values, uncertainty may be quantified by means of parametric distribution parameters (e.g., *processing factor uncertainties*, or kinetic model instance parameter uncertainties). Alternatively, uncertainty values may be provided in the form of an empirical set of uncertainty values (e.g., *relative potency factor uncertainties*, or *points of departure uncertainties*).

For each data sub-module that has quantified uncertainties, it is optional to include the source of uncertainty in the uncertainty analysis of the main module. Then, when specified, data values are resampled in each *uncertainty analysis cycle*.

The basic *acute exposure* distribution is estimated in a Monte Carlo simulation by combining dietary consumption records (person-days) with sampled residue values. The resulting distribution represents a combination of variability in consumption within the population and between residues in a food lot. Percentiles may be used for further quantification e.g. the median or 99th percentile. Due to the limited size of the underlying data, these outcomes

are uncertain. Confidence (or uncertainty) intervals reflect the uncertainty of these estimates, where MCRA uses bootstrap methodology and/or, depending on the available data, parametric methods to estimate the uncertainty.

Empirical method, resampling

The empirical bootstrap is an approach to estimate the accuracy of an outcome. In its most simple, non-parametric form, the bootstrap algorithm resamples a dataset of n observations to obtain a *bootstrap sample* or *resampled set* of again n observations (sampling with replacement, that is: each observation has a probability of 1/n to be selected at any position in the new resampled set). By repeating this process *B* times, one can obtain *B* resampled sets, which may be considered as alternative data sets that might have been obtained during sampling from the population of interest. Any statistic that can be calculated from the original dataset (e.g. the median, the standard deviation, the 99th percentile, etc.) can also be calculated from each of the *B* resampled sets. This generates a *uncertainty distribution* for the statistic under consideration. The uncertainty distribution characterises the uncertainty of the inference due to the sampling uncertainty of the original dataset: it shows which statistics could have been obtained if random sampling from the population would have generated another sample than the one actually observed (Efron (1979) and Efron and Tibshirani (1993)).

Parametric methods

Instead of bootstrapping the observed data, inference about parameters is based on parametric methods. For processing, where factors are specified through a nominal and/or upper value this is the natural choice. For concentration data, where the lognormal model is used to represent less conservative scenario's (EFSA (2012)), the *parametric bootstrap* may be an alternative, especially when data are limited and the empirical bootstrap fails.

According to Cochran's theorem, sample variance $\hat{\sigma}_y^2$ follows a scaled chi-square distribution. In the parametric bootstrap for the *lognormal* distribution, the sample variance $\hat{\sigma}_y^2$ is replaced by a random draw from a chi-square distribution with $n_1 - 1$ degrees of freedom; the sample mean $\hat{\mu}_y$ is replaced by a random draw from a normal distribution with parameters $\hat{\mu}_y$ and $\hat{\sigma}_y^{*2}/n_1$, giving a new set of parameters $\hat{\mu}_y$ and $\hat{\sigma}_y^{*2}$. This is repeated *B* times.

For the *truncated lognormal* and *censored lognormal*, large sample maximum likelihood theory is used to derive new parameters $\hat{\mu}_y$ and $\hat{\sigma}_y^{*2}$. This is repeated B times.

The binomial fraction of the censored values for the *mixture lognormal* and *mixture truncated* distribution is sampled using the beta distribution with uniform priors a = b = 1 (with the *beta* distribution as the empirical Bayes estimator for the binomial distribution). This is repeated B times.

Uncertainty due to missing data

In some cases, data are only available for specific (primary) entities and missing for others. E.g., points of departure (such as NOAELs or BMDs) may only be available for some of the substances of interest.

Uncertainty due to modelling approach

Model uncertainty or uncertainty of model outcomes arise by applying different modelling approaches or applying alternative model assumptions.

i Note		
TODO		

1.3 Data repository

The data used for the *modelling actions* of MCRA is organised in the data repository. All users have their own (personal) repository folder in which they can *upload* their own data files and organise these in folders and sub-folders to their own preference. In addition, there are shared repository folders that are specifically created for sharing data sources among user groups. Users may be granted access to one or more shared repositories: shared, maintained, and used by multiple users. Shared repositories and their contents are free to use by granted users in their own calculations.

Each data source in the data repository contains data of one or more data groups, indicating the module(s) for which the data set can be used as data. Each primary entity module and data module of the *modular design* is linked to a data group. If a data source is recognized as a data source belonging to a module, then it can be used as a data source of this module in a modelling action. The data groups of a data source are automatically recognized when the data is uploaded, based on whether the data matches with the specified data format(s) of the data group.

Figure 1.14 shows the data repository browser. The repository browsers allows users to browse through the data repository, upload and organise their own datasets and share these with other users. The central panel of the repository browser shows the data sources and sub-folders of the currently opened folder/repository. The top bar of the repository browser shows the path of the currently opened repository, buttons to collapse/expand the repository folder tree-view sidebar on the left \equiv and the info-sidebar on the right \bullet , and a button to open the action menu \vdots . The tree-view sidebar shows the hierarchical structure of the repositories and sub-repositories to which the user has access. The info panel shows the types of data available in the data source or folder. If the selected item is a data source, then the info panel shows the types of data available in the data source and the different data source versions of the data source. If the selected item is a folder, then the info panel shows info about the owner of the repository, the *access level* of the user, and info about the other users and user groups that have access to this repository.

Users with read-write access (or higher) may upload new data source files by pressing the add button + on the bottom right and selecting the *upload new file(s)* item. A new sub-repository can be created by pressing the same add button and selecting the *create new folder* item. A third option is to create an external Proast link, which can be seen as a data source repository folder in which the data sources link to datasets (outputs) available on Proast web.

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Folders	Name 👻	Version	Date	Uploader		RfD-HenG2-DGA-Mivtures-2 vlev
kruisselbrink <	↑ (EuroMix)					BIKTIEPOZ-KOA-MIXtures-2.XISX
Acropolis <	BfR-HepaRG-AdipoRed-Mixtures.xlsx	2	25-06-2019 10:20	kruisselbrink	:	Data groups
EuroMix Combined datasets	BfR-HepaRG-AdipoRed-Single.xlsx	2	25-06-2019 10:20	kruisselbrink	:	Dose response data
Concentrations	BfR-HepG2-RGA-Mixtures-2.xlsx	3	25-06-2019 10:19	kruisselbrink	÷	Versions
Consumptions	HepaRG-AdipoRed-one-expfive-subst-for training no summary.xlsx	1	18-03-2019 16:23	kruisselbrink	:	BfR-HepG2-RGA-Mixtures-2.xl v1 (28/02/2019 04:02 kruisselbrink)
Dose-response data Effects and AOP networks	HepaRG-AdipoRed-one-exp-five-subst-for training.xlsx	1	18-03-2019 16:23	kruisselbrink	:	BfR-HepG2-RGA-Mixtures-2.xl v2 (25/06/2019 10:06 <u>kruisselbrink</u>)
Foods and food translations	HepaRG-AdipoRed-one-exptwo-subst-for training no summary.xlsx	: 1	18-03-2019 16:23	kruisselbrink	:	BfR-HepG2-RGA-Mixtures-2.xl v3 (25/06/2019 10:06 <u>kruisselbrink</u>)
Hazard data	HepaRG-AdipoRed-one-exptwo-subst-for training.xlsx	1	18-03-2019 16:23	kruisselbrink	:	
In-silico data	RIVM-EST-CardioDiff-Mixtures.xlsx	2	25-06-2019 10:19	kruisselbrink	:	
Kinetic models	UGent-HepaRG-Mitochondria-Mixtures.xlsx	2	25-06-2019	kruisselbrink	:	
Non-dietary exposures			10.20			
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Test-systems and responses						•

Figure 1.14: The data repository browser.

1.3.1 Creating and uploading data files

Users with an access level of read-write (or higher) are allowed to upload new data source files to a repository folder. This can be done by pressing the add button + on the bottom right and selecting the *upload new file(s)* item.

Accepted upload file types are:

- **Microsoft Excel files (.xlsx):** An Excel file contains one or more sheets, each sheet containing tabular data. The sheet names and the fields of the data tables must comply with the conventions as specified in the data format section of the module(s) for which the data is presented.
- Microsoft Access files (.mdb): An Access file contains one or more tables. The table names and the fields of the data tables must comply with the conventions as specified in the data format section of the module(s) for
which the data is presented.

• Zip archives with CSV files (.zip): A CSV file (.csv) is a comma-separated values file containing data in tabular format. One or more CSV files are archived in a zipped file format (.zip) to facilitate the upload of collections of multiple CSV files. The names of the CSV files in the zip archive must follow the accepted table names of the module(s) for which the data is presented and the tables in the CSV files must follow the data format of that/those module(s). Note, that it is not allowed to upload single CSV files.

1.3.2 Moving repositories and data sources

The data repository browser (Figure 1.14) supports moving files (data sources) and folders (sub-repositories) via drag-and-drop functionality.

Click on a folder or file item in the browser and drag it to another folder while holding the primary mouse button. The mouse icon will change and the item will be visibly moving while dragging. Release the mouse button over the desired destination folder. MCRA will ask for confirmation before the move is executed.

The following rules apply:

- You can move items within the tree-view, list-view and between both, vice versa.
- You cannot move a root repository folder, a root repository folder is not draggable.
- You cannot move a repository to one of it's descendants.
- If you have insufficient privileges for the source or destination folder, MCRA will show an error message.

Note that moving a data source or any of it's parent repositories to a different folder does **not** affect any actions which use this data source (or any of it's versions). The action's data source will reflect the new location correctly after the data source or any of it's parent repositories has been moved.

1.3.3 Repository access levels

Shares and access rights can be granted on the level of repositories and sub-folders. Data sources inherit the access rights of the repository/folder in which these are located. The following access rights are available:

- **visible:** the user can only see that the repository exists, but cannot see its contents, except for sub-folders that may also visible to the user.
- **use:** the user is only allowed to use the data sources in this repository, but is **not** allowed to download the original data of the data sources of the repository.
- **read:** the user can use data sources in this repository **and** is allowed to download the original data files of the data sources of the repository.
- **read/write:** the user can use and download data sources in this repository and is allowed to add/remove files and folders to/from this repository.
- **admin:** the is considered as an administrator of this repository and has full control over it, including the rights to add/remove files and folders to/from this repository and to add/remove user and group shares.
- owner: the user is considered to be the owner of this repository and therefore has full control over it.

Users with administrator or owner rights on a repository/folder are allowed to add/remove user and group access using the *edit shares dialog* (Figure 1.15) that can be opened by pressing the *edit shares* button \leq .

1.3.4 Linking remote data repositories

MCRA also offers to link external data repositories **①**. These are remote websites not part of MCRA, but contain data sources that can be used for calculations. Currently, only one remote source can be linked as external repository in MCRA, the PROASTweb (https://proastweb.rivm.nl/). PROASTweb users may link directly the outputs of their PROAST analyses (i.e., dose response models) as an external repository to MCRA.

Figure 1.16 shows how PROAST outputs of a PROASTweb user are linked to an external repository in MCRA. Data sources of remote repositories have to be explicitly imported in MCRA before they can be used in analyses. Initially,

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Non-dietary exposures			
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Figure 1.15: The edit-shares dialog of the data repository browser: user and group access rights are added and removed by repository owners and administrators.

all data sources in a remote repository have a status of not-imported \triangle . Pressing the import button \bullet , MCRA will attempt to import the data source and once that is finished, the data source is ready to be used in analyses.

A new PROAST remote repository link is created by pressing the add button + on the bottom right and selecting the *Create Proast link* option. A dialog (Figure 1.17) opens asking for the local name of the external repository/folder, the PROASTweb username of the user of which the outputs should be linked, and the PROASTweb access key of the user, which is required as authentication token to access the analyses of the specified user.

1.4 Workspaces and actions

In MCRA, user work is organised in workspaces. A workspace is collection of work items (actions) and data that are logically grouped together. Each workspace has a name and can be given a description and tags. The *workspace browser* page provides users with an overview of all available workspaces and allows for creation of new workspaces. The *workspace overview page* page is shown when opening a workspace and provides an overview of the actions, data and tasks of the workspace.

The modelling tasks of MCRA are specified through actions. Each action is of a certain action type, which is the *main module* for which this action specifies the modelling task. Depending on the type of action, the user can configure the settings and the data sources of the main module and all relevant/linked sub-modules in the *action area* or, if it is a so-called standard action, in the *standard action page*. When all required settings and data sources of an action are configured and the action is in a valid state, then the modelling task described by the action can be started by pressing the run button. The status of this modelling task (which, depending on the complexity, may take some time to run) is shown in the results page and when the task has completed, output is available in the form of reports or in the form of data that can be used as input in other actions. Note that users are free to change the settings of an action at any time. Actions can therefore have multiple outputs from multiple tasks created with different settings.

A special kind of action is a so-called standard action. In a full action, the user has to link up all data and configure all settings to run the action. To get an action up and running can be quite cumbersome and the result is an output report which can be quite detailed. Although output sections follow the modular design, specific sections are not instantly available. To encourage users to explore MCRA, *standard actions* are implemented to facilitate an easy introduction to the use of MCRA. The actions can be specified with only a few settings that are important within the scope of the standard action, and the output report contains a limited selection of main sections only.

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Imports		🛆 DAS11						6
Issues	<	DAS11-NoCovar						6
Misc		Foetal	1	04-07-2019 11:46				:
Proast Kruisselbrink		Furan	1	01-07-2019 13:55				:
Projects	<	Furan-NoCovar	1	01-07-2019 12: <mark>36</mark>				:
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Figure 1.16: The remote (PROASTweb) repository in the data repository browser.

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Figure 1.17: The dialog for creating a new PROASTweb remote repository link.

A new action can be added to a workspace by clicking the + button at the bottom right of the *workspace overview* page. This will present an action menu with three options (see Figure 1.18):

- Create a new action: will open a wizard to create a (full) *action* by selecting the desired action type, specifying name, description and tags, and, depending on the action type, some main action settings.
- Create a new standard action: will open a dialog to create a *standard action* by selecting one of the available standard actions.
- **Import an action from zip file:** will open a local file browser that lets the user *import* a specifically formatted zip-file containing an action definition.



Figure 1.18: Three options to add an action to the workspace: create new action, create new standard action, and import action from zip file.

1.4.1 Workspace browser

Figure 1.19 shows the workspace browser. Users scroll through their workspaces and select the workspace which they want to work with. Detailed information about the selected item in the browser is shown in the info panel, which can be expanded/collapsed using the info button \bullet on the right of the toolbar. The *filter text box* \triangleleft is used to quickly find/filter workspaces by name or tag. A workspace is opened by clicking on the workspace name or selecting the *open workspace* \bullet option of the *action menu* \vdots of the workspace. Opening a workspace will redirect you to the *workspace overview page*.

A new workspace is added by pressing the add button + on the bottom right of the screen. Delete workspaces by opening the *action menu* \ddagger of the workspace item in the browser and selecting the delete \blacksquare option.

1.4.2 Workspace overview page

Figure 1.20 shows the workspace overview page. This page provides an overview of the actions, data, tasks, and results of a workspace, shown as four tabs at the top of the page. The actions tab shows all actions of the workspace, and from this tab, actions are opened. The data tab shows all data sources used in this workspace. I.e., all data sources that are used by the actions of the workspace. The results tab shows all tasks and results of simulation jobs that have been submitted by the actions of the workspace. The properties tab shows the general information of the workspace (i.e., name, descriptions, and tags) and edit functionality.

In the actions tab, all actions of the workspace are listed. The list of actions can be filtered by action type or by filter text using the controls on the toolbar. An action is opened by clicking on the action name or by selecting the *open action* option of the action menu \bullet of the selected action item. Opening a workspace will redirect you to the *action area*. A new action is added to the workspace by pressing the *add button* + at the bottom right of the page.

1.4.3 Action area

After opening an action, the user is directed to the main panel of the action. Each action has its own specific panel. In the main action page and sub-action pages, an action is configured, simulation jobs started, and output results are evaluated. The panel in Figure 1.21 shows the following sections:

- Scope: Links to the scope-panels in which the scope entities of the action are set (e.g., foods or substances).
- **Inputs:** Links are shown for panels in which the calculation inputs or selection inputs are set (e.g., concentration models that are inputs for computing dietary exposures).

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Workspaces		Order	r by	▼ Q euromix		×	:	
Name 👻	Created	Last modified	Tags					
EuroMix - CAG memberships calculations	01-10-2018 15:51	08-07-2019 14:42	EuroMix				:	
EuroMix - Calculations case study R&R	15-12-2018 10:31	08-07-2019 14:37	EuroMix				:	
EuroMix - Dose response models	14-03-2019 09:11	08-07-2019 14:40	EuroMix				:	
EuroMix - Effect representations	01-05-2018 12:11	08-07-2019 14:40	EuroMix				:	
EuroMix - Examples hazard characterisations	23-08-2018 10:01	08-07-2019 16:07	EuroMix				:	
EuroMix - Examples PROAST	03-04-2018 10:41	08-07-2019 16:41	EuroMix	PROAST			:	
EuroMix - Exposure mixtures calculations	23-04-2019 15:21	08-07-2019 16:42	EuroMix	Mixtures			:	
EuroMix - Hazard characterisation calculations	18-09-2018 14:51	08-07-2019 14:36	EuroMix				:	
EuroMix - Hazard characterisation tests	13-07-2018 14:41	08-07-2019 14:41	EuroMix				:	
EuroMix - Human monitoring example	11-11-2018 14:41	08-07-2019 15:54	EuroMix				:	
EuroMix - Ivive	28-04-2019 11:41	08-07-2019 16:14	EuroMix	IVIVE Risk			:	
EuroMix - Kinetic model calculations	07-01-2019 09:41	08-07-2019 16:40	EuroMix	PBPK Kinetic-mode	els		:	
EuroMix - RPF calculation scenarios	17-07-2018 15:51	08-07-2019 16:43	EuroMix	RPF		+		
EuroMix - Target exposure assessments	30-04-2019 16:31	08-07-2019 15:54	EuroMix				: .	Ŧ



MCRA 9 - EuroMix toolbox / = EuroMix toolbox / = EuroMix toolbox	🛓 kruisselbrink 🖪 🚍 🏭	0			
Actions 🗮 Data	II. Results	Properties			
Workspace actions			(44 selected)	✓ Q Type filter text here	0
Name 👻	Туре	Created	Last modified	Tags	
Aggregate exposure assessment	Exposures	07-05-2018 12:45	22-06-2018 13:57	aggregate target-exposures	:
Dietary exposure assessment	Dietary exposures	07-05-2018 16:56	08-05-2018 08:58	dietary	:
Example hazard characterisation calculation	Hazard characterisation	ons 07-05-2018 14:08	08-06-2018 14:43	target euromix example	:
Example target exposures calculation	Exposures	07-05-2018 17:06	08-06-2018 16:56	target-exposures	:
Relative potency factors	Relative potency facto	rs 07-05-2018 16:04	07-05-2018 16:41	rpf	:
Relative potency factors from data	Relative potency facto	ors 07-05-2018 16:45	07-05-2018 16:53	rpf	:
Risk assessment example	Risks	08-05-2018 09:19	08-05-2018 09:44	Risk	:



+

- **Data source:** If the action is a data action, then a form is shown in which the data source should be specified (e.g., selection of the concentration data source in a concentrations action).
- Settings: A form is shown in which the calculation and/or selection settings of the action are set/changed (e.g., specify the exposure type, chronic/acute, of an exposure assessment).

All modules of MCRA have equally structured panels. In each panel, data sources and settings for the action are specified and the scope and input sub-module links that are relevant are shown. This presentation reflects the modular design and allows the user to select the data and settings required for running the action. In the summary panel \blacksquare the main settings and data of the action are summarized. The output settings panel is used to specify general output settings. In the uncertainty settings panel \vec{z} the number of uncertainty runs and uncertainty sources is specified. In the results panel $\boldsymbol{\oplus}$ running tasks and output results of the actions are shown. An alternative form of navigating from action to sub-action is provided by the navigation menu in the left sidebar that can be expanded/collapsed by clicking the menu button on the top left in the Action bar. In this menu, all required modules for the action are shown in one list, allowing a linear way of navigation.

An action is valid and ready to run when all scopes and inputs are valid and all required data and settings are configured. For each sub-action, the check symbol \checkmark indicates that it has been configured correctly and is ready to run. In case a sub-action has a warning symbol \blacktriangle , some user action is required. When the main action is ready to run, a simulation job is started by clicking the run button \blacktriangleright in the green (sub)action bar on the top right. Optionally, sub-actions can be started by clicking the run button \triangleright in the green (sub)action bar on the top right. Clicking the run button will send the simulation task of this (sub)action to the job-scheduler, and the progress of the task is shown in the results panel \bigoplus . After completing the task, output is available in the form of a screen report, download as pdf, or as an html report with tables (csv) and charts (svg) in a downloadable zip file.



Figure 1.21: The main page of an action.

Scoping: entity selection

Each action starts with the selection of the relevant primary entities. In this context, entity selection or scoping plays an important role. Scoping of the action is defining the members for its primary entities, and, occasionally, also for other entities.

As an example, Figure 1.22 shows the substances module panel. At the top, the data source file with substances is selected containing the primary entity data of substance codes. In the selection card, a selection is made of the entities

in the dataset that are relevant for the current action (3 in scope). Note that if no explicit selection is made, the scope is set to all entities by default. In the settings form, additional (selection) settings are shown, e.g., selection of the index substance (relevant for a cumulative assessment). In this way, the scope of the action is specified by selection of the primary entities.

The panels for the data modules have a similar structure and selection is essentially the same. The only difference is that data actions always have a scope. I.e., data modules always relate to one or more primary entities.

MCRA 9 - EuroMix to Exposure, Hazard & Risk Assessm	oolbox /	The Acties / Hazard characteris	lbrink		≡ ≋	•	
Hazard characterisa Hazard characterisations action	Hazard characterisations from dose response models Hazard characterisations action ■						
Summary	^	Hazard characterisations / Substances		476ec	•	e	
Effects	~						
Responses	~	Substances data source					
Substances	~	✓ EuroMix Substances Inventory (v6).zip			~ /	۶	
• Test systems	~	Substances: 3 in scope (clear filter) 1626 only in table (add to scope)			~ =		
Hazard characterisations	~			_	-		
Active substances	~	Substances selection					
Dose response models	~	Substances: 3 in scope					
AOP networks	~						
Dose response data	~	Substance settings		B Sav	e Chang	les	
Effect representations	~	Index substance Imazalil (aka enilconazole) (RF-0246-001-PPP)			-	•	
Inter-species conversions (def	faults)						
Intra species factors (defaults	;)						
4	+						

Figure 1.22: The substances module panel as an example of a primary entity module panel.

Implicit versus explicit scoping

MCRA distinguishes between implicit and explicit selection of entities (scoping). By default, the selection is defined implicitly as 'all entities' found in all data are linked to the action. For instance, the substance scope will contain all substance codes found. That is, not only substances as specified in the substance data source, but also all other substances found in data sources that link to substances like concentration sample data or points of departure data. These are implicit selections. Explicit selections are made in the specific module panel of this data type (e.g., by selecting the substances in the substances panel). Once made explicit, selections are no longer automatically expanded when new data sources are linked to the action.

For example, the substances scope shown in Figure 1.22 is defined explicitly, having three substances in the scope, and excluding 1626 substances also present provided through substances data source and/or other linked data sources like concentration samples. By pressing the *clear filter* button, the explicit scope is cleared and is made implicit again. Then, the scope contains all substances found as primary entities and found in all linked data sources, in total 1629 (1626 + 3) substances.

Comparing new data to set scopes

After linking a data source to an action, MCRA performs a check whether the new data links well to the current scope (selected entities) of the action and reports the results. For instance, after linking new substance concentration data to an action which already has an implicit or explicit substance scope, it should be checked whether the substance codes used in the concentration data match with the current substances in scope. Note that this check is also performed after linking a primary entity substances data source to an action which already has a set of substances in scope, i.c. substances already specified in other selected data sources.

After linking a data table from a new data source to an action which already has a defined scope for one of the entities in the table, there are three possible states for entity codes:

- codes included in both the scope and the data source
- codes included in the scope, but not present in the data source
- codes included in the data source, but not present in the scope

The first case represents a successful link, no further action is required. For the second and third type of mismatch, it depends on the type of data link whether this is considered a serious problem (red flag \blacktriangle) or merely a point of attention (green flag \bigstar). For instance, in the case of concentration data, for some substances no concentrations are available, and therefore MCRA allows missing concentration data for part of the substances in the scope: a green warning symbol is shown. The concentration data source may equally well contain codes that are not in the scope (e.g., concentrations for substances that are not specified in the primary entity data for substances). It may be desirable to extend the scope with these substances found in the concentration data. Also this situation is flagged with a green warning symbol.

Figure 1.23 shows an example of a point of departure action. The substances scope has already been defined by other data in the action (in this case points of departure data), and subsequently a substances data source is selected. Here, there are 140 substances in the current scope (explicitly defined). However, 132 of these 140 substances are not present in the substances data source (*not in table*). Hence, we are missing the definitions of these substances. This is considered a critical linking issue that should be solved by updating the substances data source to include these substances, therefore a red warning symbol is shown. On the other hand, the substances data source also contains 3 substances that are not part of the current scope (*only in table*). This is a non-critical error, normally leading to a green warning symbol, but in this case, it is overruled by the red warning symbol.



Figure 1.23: Checking substances data in a substances data source against an already set substances scope.

Another example is shown in Figure 1.24. The primary entities effects and substances are selected and in the scope. Then, a points of departure data source is selected containing effect and substance codes. For effects, no linking errors are observed, hence the new data source matches perfectly with the effects already in scope. For substances, we see that there are 7 substances that are in the points of departure data source but not in the substances scope (*new*) and for 3 substances in the scope no points of departure are available (*not in table*). The former is fine, but it might be needed to extend the scope with these 7 substances (*add to scope*). The latter, in general, is not a problem but just a point of consideration. These substances might be removed from the scope (*remove from scope*) or not.

Points of departure be168c0	ib 🕨 🕒
Scope Effects (1 in scope) Substances (140 in scope)	▲ ▲
 ✓ CAG_steatose_PESTICIDES_april 2017.mdb 	~ <i>r</i>
✓ Hazard doses:	~
Effects: no linking errors	Ŧ
Substances: 7 new (add to scope) 3 not in table (remove from scope)	▲ =

Figure 1.24: Checking substances data in a POD data source against an already set substances scope.

1.4.4 Standard action area

Standard actions facilitates MCRA users in organizing the data and configuring all setting to run an action. Some standard actions are for demonstration purposes only, other standard actions are realistic examples of risk assessment or dietary exposure assessment using real data.

Creating a standard action

A new standard action is added to the workspace by pressing the *add button* + at the bottom right of the workspace page, see *workspace overview page*. Select the \bowtie *Create standard action* option.



Figure 1.25: Add standard action.

Then a pop-up appears, see Figure 1.26 and after clicking one of the available panes the standard action is created. Currently, a limited number of standard actions is available for authorized users. Please contact the MCRA system administrator at https://rivm.nl for more information. New standard actions will be developed in the near future.

lect standard action type	0
Chronic cumulative exposure assessment PA This standard action can be used to calculate a chronic cumulative expos	sure for 18 PA. Assuming equipotency, or using proposed RPFs.
Chronic cumulative exposure assessment PFAS	
'his standard action can be used to calculate a chronic cumulative expos	sure for four PFAS. Assuming equipotency, and using proposed RPFs.
Demo acute cumulative risk assessment	
n this demo with fictitious data, acute cumulative risk assessments can l nd Tier 2). Here, also the effect of applying processing factors can be as	pe performed following various calculation methods (EFSA 2012 Optimistic and Pessimistic, EC 2018 Tier 1 sessed.
EU acute cumulative exposure assessment (2018) Ti	er I and Tier II
his standard action will enable you to reproduce the exposure assessme	ent of acute cumulative effects of pesticide residues in food affecting the nervous system. These are
etrospective exposure assessments of the cumulative exposure for the r onsumption data are used. The results, data used and methodology are	ervous system using monitoring data from 2014, 2015 and 2016. In this standard action Dutch monitoring and reported in a scientific report following published on the EFSA website in September 2019. The methodology
ulfils the requirements set by the European Commission.	
EU chronic cumulative exposure assessment (2018)	Tier I and Tier II
his standard action will enable you to reproduce the exposure assessme xposure assessments of the cumulative exposure for the thyroid using r ised. The results, data used and methodology are reported in a scientific iet by the European Commission.	ent of chronic cumulative effects of pesticide residues in food affecting the thyroid. These are retrospective nonitoring data from 2014, 2015 and 2016. In this standard action Dutch monitoring and consumption data are report following published on the EFSA website in September 2019. The methodology fulfils the requirements
IPGF Acute Cumulative Risk Assessment	

Figure 1.26: Create standard action.

Standard action reports

Although a standard action produces a short output report, see Figure 1.27, by clicking the *Show detailed report* the full report becomes available. By clicking *Show short report*, the short output report is returned.

Converting a standard action to a full action

A standard action is easily converted to a full action by opening the *action menu* \vdots in the white bar of the standard workspace in your browser and selecting the \square *Convert to full action* or \square *Clone to full action* option. The first option replaces the standard action by a full action, the second option makes a clone to a full action and the standard action is still available.

1.4.5 Action zip files

MCRA also provides the functionality to import and export actions (with or without data) as specially formatted zip files (.zip) containing 'ready to run' actions. This special file archive contains two XML files (.xml) named _*ActionSettings.xml* and _*ActionData.xml* that respectively describe the action's settings and linked data sources. In addition, this zip archive may include the action data itself, either in their original form, located in a sub-folder (data) or in the form of (.csv) files.

These action zip files can be imported into and workspace by pressing the *add button* + at the bottom right of the workspace page, see *workspace overview page* and then selecting the *Import action from zip file* option. Once uploaded, the action doesn't require further user interaction and is ready to run.

An action can be exported as a zip archive by by clicking the *action menu* : located in the white bar on top of the panel. Currently, this allows export in three different formats (see Figure 1.29):

- A link Edit metadata to clone the current action to a new action (change action type) and to add a description.
- The **download action** (**no data**) option will create a zip archive containing only the two xml files defining the action, but will not include the data.

MCRA Exposure,	9.1 (Localhos Hazard & Risk Assessme	t) / 📰 Sta ent workspa	andard Act	tions			•			?
IPGF Acu Standard action	te Cumulative F - IPGF Acute Cumulative	Risk Asses Risk Assessme	ssment				¢	¢	Þ	:
Results / IPO	GF Acute Cumulativ	e Risk Asse	ssment					Show deta	ailed re	port
Resul	ts Se	ettings								
 ✓ Ha Reference Hazard ch Mean exp Mean haz Image: Constraint of the second se	azard index dist aracterisation value: osure: 0.0167 (µg/kg/ ard index: 0.0167 stazard index: 0.0167 stazard index: 0.0144 0.02772 0.03326 0.05679 0.05679	tribution p 1 (μg/kg/day) /day). Hazard Index 0.0144 0.02772 0.03326 0.05679	bercentiles	3						
99.90 ✓ Ha Hazard in Safety cha	0.1343 Azard index by s dices for 45 substanc art: bar shows variabilit CU C C N Chloorfenvi Chl	0.1343 0.1343 SUDSTANCE es based on r ty of HI (range MULATIVE - Carbofuran - Methiocarb - nfos (som) - triazofos - Cxamyl - Ethephon -	e (overviev multiple effects p5.0 - p95.0) in	V) s. n the popu	lation.	100				

Figure 1.27: Short output report.

	Edit metadata
•	Download action (no data)
•	Download action + data (data as zipped cs
►	Download action + data (original data files)
	Convert to full action
	Clone to full action
C	Refresh

Figure 1.28: Convert to full action.

- The **download action + data (data as zipped cvs)** option will create a zip archive containing the action definition xml files and also the action data in the form of csv files of the internally used data table formats.
- The **download action + data (data as zipped cvs)** option will create a zip archive containing the action definition xml files and also the original action data files (i.e., the files that were originally uploaded in the repository).
- The **download action diagram** (.svg) option will create a plot of the modular design of the current action (scalable vector graphics, SVG). Open the SVG file and it will launch in your browser. Most browser support SVG files. The nodes in the diagram contain hyperlinks to the documentation.

1.5 MCRA web and core

The Monte Carlo Risk Assessment (MCRA) platform, developed by Wageningen University & Research (WUR, Biometris) for the Dutch National Institute for Public Health and the Environment (RIVM), is a web-based system for risk assessment of chemicals, which brings together statistical models, shared data and data uploaded by the users.

EFSA and RIVM previously agreed on the use of this MCRA system for the cumulative risk assessment of pesticides. This agreement was formalised in two framework partnership agreements for the periods 2015-2016 and 2017-2020. As part of these partnership agreements, the MCRA system was continuously improved in terms of capacity and functionality, in accordance with requirements defined by EFSA. In response to needs on transparency and data accessibility, the MCRA system was further developed into a transparent, collaborative, EU harmonised, interoperable, open-source accessible platform.

1.5.1 MCRA core

Source codes of the statistical models are open source (i.e., freely accessible and re-usable) and made more flexible to allow for cooperation and co-creation. The source code of MCRA core is available at a repository located at https://github.com/rivm-syso/mcra-core/. RIVM is the owner of the MCRA core, with co-ownerships of specific parts by WUR Biometris, FERA, INERIS and possibly others in the future. RIVM will manage MCRA core, such that it is available as open source.

= Val-Exposures-Chrc	ê e	≣ :	¢ = 0 () (
Internal exposures / Overview			Edit metadata
	_	<u>+</u>	Download action (no data)
General		<u>+</u>	Download action + data (data as zipped cs
Name	Val-Exposu	<u>+</u>	Download action + data (original data files)
Description Tags	no descrip no tags	<u>+</u>	Download action diagram (.svg)
Data sources		1	Apply settings from settings xml
Effects	AOPN-Effe	1	Apply data source configuration from xml
Foods	NL-VCP-RF	C	Refresh

Figure 1.29: Export an action zip archive and other down/up-loads

1.5.2 MCRA web

The MCRA web portal provides tailored access to the MCRA core actions. In the web portal, each module of MCRA can be accessed as a starting point to perform the corresponding action. RIVM is the owner and manager of the MCRA web platform. The source code of the MCRA web platform is maintained at a dedicated RIVM github site. Access (both for reading and contributing) is restricted to the MCRA development and operations team. It is not anticipated that co-developers will contribute to the code of the web platform.

1.5.3 Running MCRA core using the command line interface

The MCRA core library is supplied with a *command line interface* (CLI) utility to run MCRA actions using input files and producing output files.

1.6 Results panel

The output of all runs is shown in the results panel. By clicking the hyperlink of an output section the output is opened in the browser. Most output tables have three icons. Click the left icon to download the table in csv format, click the middle icon to sort table columns or multiple columns (sort column, press/hold shift and sort next column) or to filter (search box). Press the question mark for additional information about the table headers.

All plots and charts can be saved by right clicking the picture.

An easy way to compare multiple outputs is to select a few outputs as shown in Figure 1.30, then click om the three dots and click Compare selected.

A new panel pops up and each output section can be reached by clicking the output name, e.g., *PARC Exposure mixtures training 0.8.* Expand the tree on the left side and select the sections you want to compare. In Figure 1.31, *Concentrations by substance* is selected in the Human monotoring analysis sub-action. Navigate to the output by clicking name of the output sections e.g., *PARC Exposure mixtures training 0.8.*

■ PARC Exposure mixtures training test Exposure mixtures action					¢	1.	¢	►	:
Results									
Results					Q Type	filter text	: here		:
Output		Status	Message	Date			Compar	e selecte	d
PARC Exposure mixtures training 0.1	1	Ran to completion		30-11	-2022 11		Delete s	elected	_
PARC Exposure mixtures training 0.8	1	Ran to completion		30-11	-2022 11	C	Refresh		
PARC Exposure mixtures training 3 components	1	Ran to completion		30-11	-2022 12	:00 00	0:00:06		:
Y PARC Exposure mixtures training kmeans	1	Ran to completion		30-11	-2022 12	:49 00	0:00:11		:
					_				

Figure 1.30: Compare selected outputs



Figure 1.31: Output comparison panel

CHAPTER

TWO

EXAMPLES

1 Note

This section is under construction.

Guidance notes used in PARC training sessions:

- Guidance how to use PARC HBM data in MCRA
- Guidance how to perform mixture analysis on PARC HBM data in MCRA
- Example HBM data

Training materials used in EuroMix training sessions:

- EuroMix dietary exposure
- RPF-exercise 1-for training-draft

There are a few exercises prepared that you could follow to get started.

2.1 Cumulative dietary exposure assessment

2.1.1 Introduction

The goal of this exercise is to perform a probabilistic cumulative dietary exposure assessment, illustrating all data needed. In Example 1 we will upload and use nine different files containing the data. In Example 2 we will upload and use a single data file for the same purpose. In the example the exposure will be characterised by upper tail percentiles, and the risk driving substances and foods can be examined. In Example 3 an uncertainty analysis is added.

2.1.2 Preparation

In the workspace browser (icon), create a new workspace *Examples*, using the + button in the bottom right corner.

2.1.3 Example 1

Calculate a cumulative chronic dietary exposure for Dutch young adults in 2003 regarding a group of eight triazole substances according to the basic optimistic model of the EFSA 2012 guidance document. Use liver steatosis as a focal effect and Cyproconazole as an index substance. The data files are already available in the data folder *Documentation-Examples / Exercise Dietary Exposure Assessment*.

Detailed steps are as follows.

- In the *Examples* workspace, create a new action using the + button in the bottom right corner.
 - Select action type Dietary exposures
 - Name it, e.g. Triazoles exposures

- (Optional) You can also add tags (e.g. triazoles, NL, steatosis) as labels that can be used later to find similar actions
- (Optional) You can add a description for further information
- Click Next
- Specify Dietary exposures settings
 - Tier: EFSA 2012 Optimistic
 - Risk type Chronic
 - Click Create

You are now directed to the main page of the new action. You can always return to this main page by clicking Action settings \clubsuit or the action type name (*Dietary exposures*) in the green bar.

The main page contains at least three blocks of information: Scope, Inputs and Settings. We will now first link all nine data files needed for this cumulative assessment. For most settings we will use default values in accordance with the chosen tier (*EFSA 2012 Optimistic*).

Scope of the assessment:

- Click Effects (path in the green bar changes Total Dietary exposures / Effects)
 - At *Effects data source*, click 🖍 and browse to the file *Effect Steatosis.xlsx*, then click *Select*
 - At Effect Settings for focal effect select Steatosis-liver and click D Save Changes
 - In the green navigation bar, click *Dietary exposures* to go up one level.
- Click Foods (path: Dietary exposures / Foods)
 - At *Foods data source*, click 🖍 and browse to the file *Foods.xlsx*, then click *Select*
 - In the green navigation bar, click *Dietary exposures* to go up one level
- Click Populations (optional) (path: Dietary exposures / Populations)
 - At Populations data source, click 🖍 and browse to the file Populations.xlsx, then click Select
 - This file contains two populations, only one is allowed. Click \checkmark under Populations selection, this opens a pop-up window. Deselect *NL_2006*, then click *Save*. The red warning signs \blacktriangle should now be gone. (Note: green warning signs \bigstar point at details and can usually be ignored)
 - In the green navigation bar, click *Dietary exposures* to go up one level.
- Click Substances (path: Dietary exposures / Substances)
 - At Substances data source, click 🖍 and browse to the file Substances Triazoles.xlsx, then click Select
 - At Substance settings for Index substance select Cyproconazole and click 🖬 Save Changes
 - In the green navigation bar, click Dietary exposures to go up one level

Next we choose the other input data:

- Click Consumptions by modelled foodd (path: Dietary exposures / Consumptions by modelled food)
 - Click Consumptions (path: Dietary exposures / Consumptions by modelled food / Consumptions)
 - At Consumptions data source, click 🖍 and browse to the file FoodConsumptions.xlsx and Select
 - At *Consumptions data selection*, with 🖍 open the food consumption surveys selection.
 - The file contains two surveys, but only one is allowed. Click ✓ under Consumptions data selection, this opens a pop-up window. Deselect *VCP-kids*, then click *Save* (the red warning ▲ should now be gone)
 - In the green navigation bar, click Consumptions by modelled food to go up one level
 - Click Food conversions (path: Dietary exposures / Consumptions by modelled food / Food conversions)

- Click Modelled foods (path: Dietary exposures / Consumptions by modelled food / Food conversions / Modelled foods)
 - Click Concentrations (path: Dietary exposures / Consumptions by modelled food / Food conversions / Modelled foods / Concentrations)
 - At *Concentrations data source*, click \checkmark and browse to the file *ConcentrationData.xlsx*, then click *Select*
 - In the green navigation bar, click Food conversions to go up two levels
- Click Food recipes (path: Dietary exposures / Consumptions by modelled food / Food conversions / Food recipes)
 - At *Food recipes data source*, click \checkmark and browse to the file *FoodTranslations.xlsx*. then click *Select*
 - In the green navigation bar, click *Dietary exposures* to go up three levels
- Click Concentration models (path: Dietary exposures / Concentration models)
 - Click Relative potency factors (path: Dietary exposures / Concentration models / Relative potency factors)
 - At *Relative potency data source*, click / and browse to the file *RPFs.xlsx*, then click *Select*
 - In the green navigation bar, click *Dietary exposures* to go up two levels
- Click Processing factors (path: Dietary exposures / Processing factors)
 - At Processing factors data source, click 🖍 and browse to the file ProcessingFactors.xlsx, then click Select
 - In the green navigation bar, click Dietary exposures to go up one level
- Click Active substances (optional) (path: Dietary exposures / Active substances)
 - In this example we have a fixed list of relative potency factors for the eight substances, and don't need point of departure (POD) data to decide which substances are active with respect to the health effect and therefore belong to the cumulative assessment group. Deselect the setting "Derive memberships from POD presence", then click Save Changes
 - In the green navigation bar, click *Dietary exposures* to go up one level

Now run the model, either by clicking the \blacktriangleright run icon in the grey bar, or by clicking the \blacktriangleright run icon in the green bar (Note: \blacktriangleright in the green bar can also be used to run subactions on their own).

The \blacktriangleright icon is replaced by the text "Running". When the run has finished, the interface automatically changes to the Results screen. You can also click the Results icon \clubsuit to go there.

As an exercise, try find the following results:

- 1. The 99th percentile of cumulative exposure
- 2. The substance(s) with highest contribution to the total exposure
- 3. The food(s)-as-measured with the highest contribution to the upper tail of the exposure distribution

Answers:

- In the grey bar, browse to the results panel by clicking the ① icon and click on the latest output (path: *Results / Dietary exposures*)
 - In the *Dietary exposures* tab, browse in the tree (unfold by clicking > where necessary) to > *Dietary exposures* > *Distribution (OIM)* > *Percentiles*
 - In the table it states that the 99% exposure percentile is at an exposure of $0.02127 \,\mu$ g/kg bw/day.
 - In the *Dietary exposures* tab, browse in the tree (unfold by clicking > where necessary) to > *Dietary exposures* > *Details* > *Exposures by substance* > *Total distribution*
 - From the pie chart it is clear that Tebuconazole contributes the most to the total exposure distribution with 32.7%. In the table below the graph more details can be found.

- In the *Dietary exposures* tab, browse in the tree (unfold by clicking > where necessary) to > *Dietary exposures* > *Details* > *Exposures by food and substance* > *Risk drivers upper tail*
 - From the pie chart it is clear that Flusilazole in grapefruit contributes the most (16.7%) to the upper tail exposure distribution

2.1.4 Example 2

We will create a new action to demonstrate uploading all the data at once. All data is now contained within one file, *MCRA-Documentation Example Dietary exposures.xlsx*.

Detailed steps are as follows.

- In the *Examples* workspace, create a new action (using +)
 - Select action type Dietary exposures
 - Name it, e.g. Triazoles exposures from one data file
 - Click Next
- Specify Dietary exposures settings
 - Tier: EFSA 2012 Optimistic
 - Risk type *Chronic*
 - Click Create
- Then go to the actions settings 🌣 of this action (path: *Dietary exposures*)
 - Click Effects (path: Dietary exposures / Effects)
 - At Effects data source, click
 And browse to the file MCRA-Documentation Example Dietary exposures.
 At Effects data source, click
 At eff

You still need to specify the focal effect (under *Effects*), index substance (under *Substances*), and food surveys (under *Consumptions by modelled food / Consumptions*). You also need to deselect the "Derive memberships from POD presence" setting under *Active substances*. Navigate to the subaction where these changes have to be made using the green bar.

You now have achieved the same as in Example 1, only with the upload of one single file. You can now run the model, and inspect the results, which should be the same as for Example 1.

2.1.5 Example 3

Repeat the run of the previous task, but in addition to the nominal run, perform an uncertainty analysis as well.

- Click on the \neq icon (in the grey bar) to open the uncertainty settings panel
 - At Uncertainty settings, check ✓ Perform uncertainty analysis
 - For Monte Carlo iterations per uncertainty run choose 100, and press Save Changes
- Now run the model, by pressing the 🕨 run icon in the grey bar. Note that the run will take much more time.

Compare with the previous results, to find:

- 1. 95% uncertainty bounds for the 99% exposure percentile
- 2. 95% uncertainty bounds for the highest contribution from a substance to the total exposure distribution
- 3. 95% uncertainty bounds for the highest contribution from a food to the total exposure distribution

2.2 TDS-based exposure and risk assessment

Total Diet Studies (TDS) monitor chemical levels in representative consumed foods as eaten (e.g. bread, pizza), so after processing steps (Lee et al. (2015)). TDSs contrast to the common chemical monitoring studies on raw primary commodities (e.g. wheat, tomato). In some TDSs, samples are analysed directly, in other TDSs samples of similar food products are pooled before chemical analysis. Pooling of samples means that average concentrations can be well estimated, but that less or no infromation is available about variability. Therefore, in the context of risk assessment, TDS data are used for the assessment of chronic risks from long-term exposure.

This section contains three demonstrators (*MCRA standard actions*) on how to use TDS data for exposure and risk assessment. The demonstrators were developed in the FNS-Cloud project. Two examples illustrate simple TDS data on consumed foods. The first demonstrator shows the exposure to methyl-mercury and the risk for German children. The second demonstrator shows the exposure to nickel and the risk for several Belgian age groups. DON for Dutch children. The third demonstrator contains the previous two as specific cases, but also includes a case where foods are pooled before analysis, for the exposure to the mycotoxin DON and the risk for Dutch children. In this last case additional data are included to describe the TDS sample compositions in terms of the modelled foods.

After trying out the standard actions, prospective users can convert or clone these standard actions to full actions for use with other data or using other exposure or risk models.

2.2.1 Standard actions

The following standard actions are available as TDS exposure and risk assessment demonstrators:

- TDS-based long-term exposure and risk assessment of methylmercury for German children
- Long-term dietary exposure and risk of nickel for the Belgian population
- TDS-based long term dietary exposure and risk assessment

2.3 Aggregate exposure assessment

2.3.1 Introduction

The goal of this exercise is to assess aggregate exposure assessment.

2.3.2 Preparation

If you haven't done so, in the workspace browser (use the \square icon), create a new workspace named *Examples*, using the +.

The data files used in the example(s) in this section, are located in the data folder *Documentation-Examples / Exercise* Aggregate Exposure Assessment.

2.3.3 Example 1

- In the *Examples* workspace, create a new action (using +)
 - Then select ✓ Show all action types, select Exposures
 - Name it exposures
 - At Exposure settings choose:
 - As Risk type Chronic
 - Check ✓ Include dietary and non-dietary routes of exposure
 - Press Create
- Then go to the Actions settings 🍄 of this action (path: *Exposures*)
 - At Scope, click Effects (path: Exposures / Effects)
 - At *Effects data source* with 🖍 browse to the file *Effect Steatosis.xlsx* and *Select*

- At *Effect settings* for *Focal effect* select *Steatosis-liver* and press **D** *Save Changes*
- In the green navigation bar, click Exposures to go up one level
- At Scope, click Foods (path: Exposures / Foods)
 - At *Foods data source* with *I* browse to the file *Foods.xlsx* and *Select*
 - In the green navigation bar, click *Exposures* to go up one level
- At Scope, click Substances (path: Exposures / Substances)
 - At Substances data source with 🖍 browse to the file Substances.xlsx and Select
 - At Substance settings
 - for Index substance select Cyproconazole and press Save Changes
 - In the green navigation bar, click Exposures to go up one level
- At Inputs, click Dietary exposures (path: Exposures / Dietary exposures)
 - At Inputs, click Consumptions by modelled food (path: Exposures / Dietary exposures / Consumptions by modelled food)
 - At Inputs, click Consumptions (path: Exposures / Dietary exposures / Consumptions by modelled food / Consumptions)
 - At Consumptions data source with 🖍 browse to the file Consumptions.xlsx and Select
 - At *Consumptions data selection* with \checkmark open the food consumption surveys selection.
 - The file contains two surveys, but only one is allowed. So deselect everything by clicking ✓ on the first line, next to the word *Code*
 - Now select *DNFCS_2003* and press *Save* (the red warning **A** should now be gone)
 - In the green navigation bar, click *Consumptions by modelled food* to go up one level
 - At Inputs, click Food conversions (path: Exposures / Dietary exposures / Consumptions by modelled food / Food conversions)
 - At Inputs, click Modelled foods (path: Exposures / Dietary exposures / Consumptions by modelled food / Food conversions / Modelled foods)
 - At Inputs, click Concentrations (path: Exposures / Dietary exposures / Consumptions by modelled food / Food conversions / Modelled foods / Concentrations)
 - At Concentration data source with 🖍 browse to the file ConcentrationData.xlsx and Select
 - In the green navigation bar, click Food conversions to go up two levels
 - At Inputs, click Food recipes (path: Exposures / Dietary exposures / Consumptions by modelled food / Food conversions / Food recipes)
 - At Food recipes data source, with 🖍 browse to the file FoodRecipes.xlsx and Select
 - In the green navigation bar, click *Dietary exposures* to go up three levels
 - At Inputs, click Concentration models (path: Exposures / Dietary exposures / Concentration models)
 - At Inputs, click Relative potency factors (path: Exposures / Dietary exposures / Concentration models / Relative potency factors)
 - At *Relative potency factors data source* with browse to the file *RelativePotencyFactors.xlsx* and *Select*
 - In the green navigation bar, click Dietary exposures to go up two levels
 - At Inputs, click Processing factors (path: Exposures / Dietary exposures / Processing factors)
 - At Processing factors data source with 🖍 browse to the file ProcessingFactors.xlsx and Select
 - In the green navigation bar, click *Dietary exposures* to go up one level

- At Inputs, click Active substances (optional) (path: Exposures / Dietary exposures / Active substances)
 - At Inputs, click Points of departure (path: Exposures / Dietary exposures / Active substances / Points of departure)
 - At Points of departure data source with 🖍 browse to the file HazardDoses Triazoles.xlsx
 - In the green navigation bar, click Dietary exposures to go up two levels
- At Dietary exposure settings, for Dietary exposure calculation tier select EFSA 2012 Optimistic, and press Save Changes
- In the green navigation bar, click Exposures to go up one level
- At Inputs, click Non-dietary exposures (path: Exposures / Non-dietary exposures)
 - At Non-dietary exposures data source with ✔ browse to the file NonDietaryExposures.xlsx and Select
- Now run the model, by pressing the **>** run icon in the grey bar.

Try to find the following results:

- 1. Exposure percentiles daily intakes with uncertainty bounds
- 2. Substance with highest contribution to the total exposure distribution
- 3. The modelled food measured with the highest contribution to the upper tail of the exposure distribution

2.3.4 Example 2

In this example we will elaborate on the previous one with kinetic models.

- Go to the Actions settings 🌣 of this action (path: *Exposures*)
 - At Inputs, click Kinetic models (default) (path: Exposures / Kinetic models)
 - At Kinetic models data source with browse to the file UserGroupDemo-KineticModelsArtificial.xlsx and Select
 - At Kinetic model settings for Kinetic model select Cosmos Version 5
- Now run the model, by pressing the ▶ run icon in the green bar.

2.4 Hazard characterisations from PoDs

2.4.1 Introduction

The goal of this exercise is to try to establish hazard characterisations from PoDs (NOAELs).

2.4.2 Preparation

If you haven't done so, in the workspace browser (use the \square icon), create a new workspace named *Examples*, using the +.

The data files used in the example(s) in this section, are located in the data folder *Documentation-Examples / Exercise Hazard characterisations*.

2.4.3 Example 1

In this example, Imazalil target dose from NOAEL will be calculated.

- In the *Examples* workspace, create a new action (using +)
 - Then select ✓ Show all action types, and select Hazard characterisations
 - Name it TargetDoseImazalil
 - · Use as Hazard characterisation settings

- Risk type: Chronic
- Target level: External
- Press Create
- Then go to the Actions settings 🌣 of this action.
 - At Scope, click Effects (path: Hazard characterisations / Effects)
 - At Effects data source with ✓ browse to the file Effects and AOP Network Steatosis.xlsx and Select
 - At Effects selection with ▲
 - Deselect everything by clicking \checkmark on the first line, next to the word *Code*
 - On the second page, select only *Steatosis-liver*, and **D** Save
 - At *Effect Settings* for *focal effect* select *Steatosis-liver* and press **D** Save Changes.
 - In the green navigation bar, click Hazard characterisations to go up one level
 - At Scope, click Substances (path: Hazard characterisations / Substances)
 - At Substances data source with 🖍 browse to the file TargetDosescalculation-Substances.xlsx and Select
 - At Substances selection with
 - Deselect everything, by clicking the \checkmark on the first line, next to the word *Code*
 - Select only *Imazalil*, and **B** Save
 - In the green navigation bar, click Hazard characterisations to go up one level
 - At Inputs, click Points of departure (path: Hazard characterisations / Points of departure)
 - At *Points of departure data source* with \checkmark browse to the file *TargetDosesCalculation-HazardDoses.xlsx* and *Select*
 - In the green navigation bar, click Hazard characterisations to go up one level
 - At Hazard characterisations settings, for Expression type select NOAEL (convert all hazard characterisations as NOAELs)
 - At Hazard characterisations settings, Select \checkmark Use inter-species conversions
 - At *Hazard characterisations settings*, Select \checkmark Use intra-species factors, and press 🖻 Save Changes
- Now run the model, by pressing the **>** run icon in the grey bar.

Try to find the following results:

- 1. The NOAEL for Imazalil used as point of departure.
- 2. The target hazard dose based on the default assessment factors 1/10 and 1/10 for inter-species and withinspecies conversion.

Answers:

- In the grey bar, browse to the results panel by clicking the \bigcirc icon and click on the latest output (path: *Results / TargetDoseImazalil*)
 - In the *Hazard characterisations* tab, browse in the tree (unfold by clicking > where necessary) to > *Available hazard characterisations*
 - The NOAEL for Imazalil is 40 μ g/kg bw/day.

2.5 Health impact estimates

2.5.1 Introduction

The goal of this exercise is to assess a health impact estimate.

2.5.2 Preparation

If you haven't done so, in the workspace browser (use the \square icon), create a new workspace named *Examples*, using the +.

The data files used in the example(s) in this section, are located in the data folder *Documentation-Examples / Exercise Health Impact*.

2.5.3 Example 1

- In the *Examples* workspace, create a new action (using +)
 - Then select ✓ Show all action types, select Risks
 - Name it Risks
 - Press Create
- Then go to the Actions settings 🌣 of this action (path: *Risks*)
 - At Scope, click Effects (path: Risks / Effects)
 - At Effects data source with ✓ browse to the file Effects and AOP Network Steatosis.xlsx and Select
 - At * Effect settings*, for *focal effect* select *Steatosis-liver* and press **D** *Save Changes*
 - In the green navigation bar, click Risks to go up one level
 - At Scope, click Foods (path: Risks / Foods)
 - At *Foods data source* with *I* browse to the file *Foods.xlsx* and *Select*
 - In the green navigation bar, click Risks to go up one level
 - At Scope, click Substances (path: Risks / Substances)
 - At Substances data source with 🖍 browse to the file Substances.xlsx and Select
 - At Substance settings, for index substance select Cyproconazole, and press B Save Changes
 - In the green navigation bar, click *Risks* to go up one level
 - At Inputs, click Exposures (path: Risks / Exposures)
 - At Inputs, click Dietary exposures (path: Risks /Exposures / Dietary exposures)
 - At Inputs, click Consumptions by food measured (path: Risks /Exposures / Dietary exposures / Consumptions by modelled food)
 - At Inputs, click Consumptions (path: Risks /Exposures / Dietary exposures / Consumptions by modelled food / Consumptions)
 - At Consumptions data source with 🖍 browse to the file Consumptions.xlsx and Select
 - At *Consumptions data selection* with 🖍 open the food consumption surveys selection.
 - The file contains two surveys, but only one is allowed. So deselect everything by clicking \checkmark on the first line, next to the word *Code*
 - Now select *DNFCS_2003* and press *Save* (the red warning **A** should now be gone)
 - At Consumptions settings for Food survey select DNFCS_2003 and press Save Changes
 - In the green navigation bar, click Consumptions by modelled food to go up one level

- At Inputs, click Food conversions (path: Risks /Exposures / Dietary exposures / Consumptions by modelled food / Food conversions)
 - At Inputs click Food as modelled (path: Risks /Exposures / Dietary exposures / Consumptions by modelled food / Food conversions / Food as modelled)
 - At Inputs, click Concentrations (path: Risks /Exposures / Dietary exposures / Consumptions by modelled food / Food conversions / Food as modelled / Concentrations)
 - At *Concentrations data source* with
 ✓ browse to the file *UserGroupDemo-ConcentrationData.xlsx* and *Select*
 - In the green navigation bar, click Food conversions to go up two levels
 - At Inputs, click Food recipes (path: Risks /Exposures / Dietary exposures / Consumptions by modelled food / Food conversions / Food recipes)
 - At *Food recipes data source* with \checkmark browse to the file *UserGroupDemo-FoodRecipes.xlsx* and *Select*
 - In the green navigation bar, click *Dietary exposures* to go up three levels
- At Inputs, click Processing factors (path: Risks /Exposures / Dietary exposures / Processing factors)
 - At *Processing factors data source* with \checkmark browse to the file *UserGroupDemo-ProcessingFactors.xlsx* and *Select*
 - In the green navigation bar, click *Risks* to go up three levels
- At Inputs, click Hazard characterisations (path: Risks / Hazard characterisations)
 - At Inputs, click Active substances (path: Risks / Hazard characterisations / Active substances)
 - At Inputs, click Points of departure (path: Risks / Hazard characterisations / Active substances / Points of departure)
 - At *Points of departure data source* with \checkmark browse to the file *UserGroupDemo-HazardDoses.xlsx* and *Select*
 - In the green navigation bar, click Active substances to go up one level
 - At 'Active substances' click 🌣 Compute

2.6 Assessment group membership probabilities

2.6.1 Introduction

The goal of this exercise is to assess group membership probabilities.

2.6.2 Preparation

If you haven't done so, in the workspace browser (use the \square icon), create a new workspace named *Examples*, using the +.

The data files used in the example(s) in this section, are located in the data folder *Documentation-Examples / Exercise Dietary Exposure Assessment*.

2.6.3 Example 1

- In the *Examples* workspace, create a new action (using +)
 - Then select Dietary exposures
 - Name it Dietary exposures

- Use as Dietary exposures settings
 - Tier: EFSA Guidance Optimistic
 - Risk type Chronic
 - Select ✓ *Cumulative*
- Press Create
- Then go to the actions settings 🌣 of this action (path: *Dietary exposures*)
 - At Scope, click Foods (path: Dietary exposures / Foods)
 - At Foods data source with 🖍 browse to the file UserGroupDemo-Foods.xlsx and Select
 - In the green navigation bar, click Dietary exposures to go up one level
 - At Scope, click Substances (path: Dietary exposures / Substances)
 - At Substances data source with ✔ browse to the file UserGroupDemo-Substances.xlsx and Select
 - At Substance settings for Index substance select Cyproconazole and press D Save Changes
 - In the green navigation bar, click *Dietary exposures* to go up one level
 - At Scope, click Effects (path: Dietary exposures / Effects)
 - At Effects data source with 🖍 browse to the file Effect Steatosis.xlsx and Select
 - At Effect Settings for focal effect select Steatosis-liver and press D Save Changes
 - In the green navigation bar, click *Dietary exposures* to go up one level.
 - At Inputs, click Consumptions by modelled food (path: Dietary exposures / Consumptions by modelled food)
 - At Inputs, click Consumptions (path: Dietary exposures / Consumptions by modelled food / Consumptions)
 - At *Consumptions data source* with ▲ browse to the file *UserGroupDemo-Consumptions.xlsx* and *Select*
 - At Consumption settings for Food survey select DNFCS_2003 and press Save Changes
 - In the green navigation bar, click Consumptions by modelled food to go up one level
 - At Inputs, click Food conversions (path: Dietary exposures / Consumptions by modelled food / Food conversions)
 - At Inputs, click Modelled foods (path: Dietary exposures / Consumptions by modelled food / Food conversions / Modelled foods)
 - At Inputs, click Concentrations (path: Dietary exposures / Consumptions by modelled food / Food conversions / Modelled foods / Concentrations)
 - At *Concentrations data source* with *I* browse to the file *UserGroupDemo-ConcentrationData.xlsx* and *Select*
 - In the green navigation bar, click Food conversions to go up two levels
 - At Inputs, click Food recipes (path: Dietary exposures / Consumptions by modelled food / Food Food recipes)
 - At *Food recipes data source*, with \checkmark browse to the file *UserGroupDemo-FoodRecipes.xlsx*
 - In the green navigation bar, click *Dietary exposures* to go up three levels
 - At Inputs, click Concentration models (path: Dietary exposures / Concentration models)
 - At Inputs, click Relative potency factors (path: Dietary exposures / Concentration models / Relative potency factors)

- At *Relative potency data source* with *r* browse to the file *UserGroupDemo-RelativePotencyFactors.xlsx* and *Select*
- In the green navigation bar, click Dietary exposures to go up two levels
- At Inputs, click Processing factors (path: Dietary exposures / Processing factors)
 - At *Processing factors data source* with ✓ browse to the file *UserGroupDemo-ProcessingFactors.xlsx* and *Select*
 - In the green navigation bar, click Dietary exposures to go up one level
- At Inputs, click Active substances (optional) (path: Dietary exposures / Active substances)
 - At Inputs, click Points of departure (path: Dietary exposures / Active substances / Points of departure)
 - At Points of departure data source, with 🖍 browse to the file HazardDoses Triazoles.xlsx
 - In the green navigation bar, click *Dietary exposures* to go up two levels
- Now run the model, by pressing the **>** run icon in the grey bar.

Try to find the following results:

- 1. Exposure percentiles daily intakes
- 2. Substance with highest contribution to the total exposure distribution
- 3. The food-as-measured with the highest contribution to the upper tail of the exposure distribution

2.6.4 Example 2

Repeat the run of the previous task, but instead of the nominal run, now do an uncertainty analysis loop.

- Click on the [∓] icon (in the grey bar) to open the uncertainty settings panel, and check ✓ *Perform uncertainty analysis*
 - For Monte Carlo iterations per uncertainty run choose 100, and press 🖬 Save Changes
- Now run the model, by pressing the **b** run icon in the grey bar.

Compare with the previous results, to find:

- 1. Exposure percentiles daily intakes with uncertainty bounds
- 2. Substance with highest contribution to the total exposure distribution
- 3. The food-as-measured with the highest contribution to the upper tail of the exposure distribution

Part II

Reference Manual

CHAPTER THREE

MODULES

MCRA is a modular system. The diagram in Figure 3.1 shows the modules and their relations. Each module is associated with its own type of data, and is linked to one or more other modules. Note that not all details can be fully shown in the scheme, for details consult the table below, which specifies all relations between the modules in MCRA.



Figure 3.1: Modular design of MCRA.

3.1 Primary entity modules

The MCRA modular system is based on six primary entities, defining who (*Populations*) is to be protected against what impact (*Effects*) caused by what agent (*Substances*) originating from where (*Foods*), with an indication how the effects are quantified (*Responses* in *Test systems*).

3.1.1 Effects

Effects are biological or toxicological consequences for human health, that may result from chemical exposure and are the focus of hazard or risk assessment.

Output of this module is used by: Concentration models High exposure food-substance combinations Dietary exposures Exposure mixtures QSAR membership models Molecular docking models Active substances Relative potency factors Hazard characterisations Points of departure Effect representations Inter-species conversions Intra species factors AOP networks Exposure response functions Burden of disease Risks Single value risks

Effects from data

Effects calculation

Option *Multiple effects analysis* selects multiple selects. Effects are selected using **Effects selection**. Press **Clear filter** and/or **change selection** and select multiple effects in the scroll down box. If both this option and *Include related effects of AOP network* is unchecked, it is obligatory to select one and only one effect. If *Include related effects of AOP network* is checked, a **Focal effect** is specified and all related effects in the *AOP network* are selected as well.

Effects data formats

Effects are primary entities of the data model. Health effects are defined as (critical) changes relative to a treatment or exposure.

Download empty dataset template: Zipped CSV Excel

Effects

Effects are uniquely identified by a code (idEffect). Optionally, a name and description can be added. Health effects are commonly distinguished in two types, acute and chronic. Further properties may be specified, e.g. in relation to decision schemes such as the use of thresholds of toxicological concern (TTCs).

Name	Туре	Description	Aliases	Required
idEffect	AlphaNumeric (50)	Unique identification code of the effect.	idEffect, EffectId, Code- FocalEffect, Id, Code, KeyEvent, idKeyEvent	Yes
CodeSystem	AlphaNumeric (100)	Identifier of the coding system of the effect code.	CodeSystem	No
Name	AlphaNumeric (100)	Name of the effect.	Name	No
Description	AlphaNumeric (200)	Additional description or label of the effect.	Description	No
Biological- Organisation	Biological- OrganisationType	Biological organisation of the effect: Molecular, Cellular, Tissue, Organ, Individual, Population. This is in line with AOP wiki terminology and can be used for grouping.	Biological- Organisation	No
KeyEvent- Process	AlphaNumeric (100)	Description of AOP Key event component process. E.g., receptor signalling.	Process	No
KeyEvent- Object	AlphaNumeric (100)	Description of AOP Key event component object. E.g., PPAR-alpha.	Object	No
KeyEvent- Action	AlphaNumeric (100)	Description of AOP Key event component action. E.g., decreased.	Action	No
KeyEventOrgan	AlphaNumeric (100)	Description of AOP Key event organ. E.g., liver.	Organ	No
KeyEventCell	AlphaNumeric (100)	Description of AOP Key event organ. E.g., hepatocyte.	Cell	No
AOPwikiKE	AlphaNumeric (200)	Key event ID number in AOP wiki https://aopwiki.org/events Several ID possible Some effects might not be in the wiki, and this field will be empty.	AOPWikiIds, AOPwikiKE	No
Reference	AlphaNumeric (200)	External reference(s) to sources containing more information about the AOP key event. E.g., the AOP wiki, and the associated AOP wiki Ids.	References	No

Table 3.1: Table definition for Effects.

Accepted table names: Effects, Effect, KeyEvents, KeyEvent.

Effects settings

Selection settings

	e	
Name	Туре	Description
Multiple effects analysis	Boolean	Specifies whether the analysis should consider multiple effects. Otherwise, a single focal effect should be selected.
Focal effect	AlphaNumeric	The main (health) effect of interest.
Include related effects of AOP network	Boolean	Include all related key events of the AOP network.

Table 3.2: Selection settings for module Effects.

Effects as data

Effect definitions are provided as lists/catalogues of effect definitions.

- Effects data formats
- Effects from data
- Effects calculation

3.1.2 Foods

Foods are uniquely defined sources of dietary exposure to chemical substances. Foods may refer to 1) foods as eaten, foods as coded in food consumption data (e.g. pizza); 2) modelled foods, foods as coded in concentration data (e.g. wheat, tomato); 3) any other type of food (e.g. ingredients like flour, tomato sauce).

Output of this module is used by: Consumptions Single value consumptions Market shares Food recipes Concentrations Concentration distributions Single value concentrations Processing factors Unit variability factors Occurrence patterns Occurrence frequencies Substance authorisations Deterministic substance conversion factors Concentration limits Concentration models Modelled foods Focal food concentrations Total diet study sample compositions Food extrapolations Food conversions Consumptions by modelled food High exposure food-substance combinations Dietary exposures Single value dietary exposures Internal exposures

Foods from data

Foods data formats

Foods are of interest in (dietary) consumption assessments and the sources of exposure within exposure assessments. The foods table is the main table of the food definitions. Relevant food related data, such as processing types, additional properties (e.g., unit weight and brand loyalty), facets, and hierarchies, can be described in the food properties, food hierarchies, and faces and facet descriptors tables.

Download empty dataset template: Zipped CSV Excel

Foods

Each food is identified by a unique code (idFood) in a code system of choice, a name, and a description. Food codes can have a hierarchical structure (as in the FoodEx1 coding systems), using '.' or '\$' as separator between adjacent hierarchical levels, e.g. 'A.05' is fruits and fruit products, 'A.05.01' is citrus fruits, and 'A.05.01.001' is grapefruit (citrus paradisi). Additional forms of foods, such as foods in processed form, can be specified via food facets according to the FoodEx2 system of EFSA.

Name	Туре	Description	Aliases	Required
idFood	AlphaNumeric (50)	The unique identification code of the food.	idFood, Code, FoodId, FoodCode, Food, Id	Yes
Name	AlphaNumeric (100)	The name of the food.	Name, FoodName	No
Description	AlphaNumeric (200)	Food description.	Description	No

Table 3.3: Table definition for Foods.

Accepted table names: Foods, Food.

Food properties

Additional food properties. This table is deprecated. See table FoodUnitWeights which partly replaces this table for data on food unit weights.

Table 3.4: Table definition	for Food	properties.
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Name	Туре	Description	Aliases	Required
idFood	AlphaNumeric (50)	The code of the food to which the property is attached. The provided food code should match with a code of the foods table.	idFood, FoodId, Food, FoodCode, Code	Yes

Accepted table names: FoodProperties, FoodProperty.

Food unit weights

Food unit weights as specified for a food, and possibly a location.

Name	Туре	Description	Aliases	Required
idFood	AlphaNumeric (50)	The unique identification code of the food.	idFood, Code, FoodId, FoodCode, Food, Id	Yes
Location	AlphaNumeric (50)	The location for which this food unit weight is defined. If not specified, then the value is considered a default unit weight that can be used when there is no location specific unit weight.	Location	No
ValueType	UnitWeightValue- Type	The value type of the unit weight value (i.e., raw agricultural commodity or edible portion). Controlled terminology.	ValueType, UnitWeight- ValueType	No
Qualifier	ValueQualifier	Qualifier of the unit weight value, e.g. equal-to (=) or smaller-than (<). If omitted, = is assumed.	Qualifier, QualifierType	Yes
Value	Numeric	Unit weight value in grams.	Value, Unit- WeightValue, UnitWeight	Yes
Reference	AlphaNumeric (200)	External reference(s) to source of the unit weight value.	Reference, References	No

Accepted table names: FoodUnitWeights, UnitWeights.

Food hierarchies

Food items are commonly categorised in hierarchies, e.g. oranges and mandarins are citrus fruits. For example FoodEx is a food description and food classification (FDFC) system consisting of a large number of individual food items aggregated into food groups and broader food categories in a hierarchical structure of parent-child relationships.

Table 3.6: Tab	ble definition	for Food	hierarchies.
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Name	Туре	Description	Aliases	Required
idFood	AlphaNumeric (50)	Food node.	idFood, FoodId, Food, Code	Yes
idParent	AlphaNumeric (50)	Parent node of the food.	idParent, ParentId, Parent, ParentCode	Yes

Accepted table names: FoodHierarchies, FoodHierarchy, FoodsHierarchy.

Facets

Food codes can be linked to facets, as e.g. in FoodEx2. FoodEx2 is a comprehensive food classification and description system aimed at covering the need to describe food in data collections across different food safety domains.

See the EFSA catalogue browser for facets currently used e.g. the facet to describe processing technology is: F28 process.

Name	Туре	Description	Aliases	Required
idFacet	AlphaNumeric (10)	The food code of the food to which the facet is attached.	idFacet, Code, Id, FacetCode, FacetId	Yes
Name	AlphaNumeric (200)	Facet name	Name, FacetName	No
Description	AlphaNumeric (200)	Additional description of the facet.	Description	No

Table 3.7: Table definition for Facets.

Accepted table names: Facets, Facet, FoodFacets, FoodFacet.

Facet descriptors

Facet descriptors are elements of additional information on a facet such as processing.

Name	Туре	Description	Aliases	Required
idFacet- Descriptor	AlphaNumeric (10)	The identification code of the facet descriptor.	idFacet- Descriptor, Code, Id, FacetCode, FacetId	Yes
Name	AlphaNumeric (200)	The name of the facet descriptor.	Name, Facet- DescriptorName	No
Description	AlphaNumeric (200)	Additional description of the facet descriptor.	Description	No

Table 3.8: Table definition for Facet descriptors.

Accepted table names: FacetDescriptors, FacetDescriptor, FoodFacetDescriptors, FoodFacetDescriptor.

Processing types

Name	Туре	Description	Aliases	Required
idProcessing- Type	AlphaNumeric (50)	The unique identification code of the processing type.	idProcessing- Type, ProcessingType- Id, ProcType, Id	Yes
Name	AlphaNumeric (100)	The processing name.	ProcName, Name	No
Description	AlphaNumeric (200)	The processing type description.	Description	No
Distribution- Type	Processing- DistributionType	The distribution type. Simulated processing factors are restricted to the interval (0,1) using a logistic-normal distribution (default) or simulated processing factors are restricted to positive values using a log-normal distribution.	Distribution- Type, DistType	No
Bulking- Blending	Boolean	For types of processing applied on large batches, e.g., juicing, sauce/puree. Default is no bulking blending.	Bulking- Blending, BulkBlending, IsBulkBlending	No

Table 3.9: Table definition for Processing types.

Accepted table names: ProcessingTypes, ProcessingType.

Food consumption quantifications

Food consumption quantifications record information about food consumption quantities that are associated with unit-consumptions of foods.

Name	Туре	Description	Aliases	Required
idFood	AlphaNumeric (50)	The food code of the quantification.	idFood, FoodId, Food	Yes
idUnit	AlphaNumeric (50)	The code of the unit of consumption. E.g spoon, plate, cup. Units may depend on food.	idUnit, UnitId, Unit	Yes
UnitWeight	Numeric	The unit weight/portion size of the food, specified in grams.	UnitWeight	Yes
UnitWeight- Uncertainty	Numeric	The uncertainty in unit weight/portion size (%).	UnitWeight- Uncertainty, UnitWeight%	No
Amount- Uncertainty	Numeric	The uncertainty in amount consumed (%). The label 'general' specifies a default value for the uncertainty when specific information for combinations of food and unit in food consumptions table is not available.	Amount- Uncertainty, Amount%	No

Table 3.10: Table definition for Food consumption quantifications.
Accepted table names: FoodConsumptionQuantifications, FoodConsumptionQuantification.

Foods as data

Food definitions are provided as lists/catalogues of food definitions, optionally with encompassing processing type definitions, facet definitions, hierarchy definitions, and additional food property information.

- Foods data formats
- Foods from data

Food coding systems

MCRA is intended to retain complete transparence of the results of risk assessment in terms of the foods that were actually consumed (foods-as-eaten). In many cases measurements of substances have not been made on the **food-as-eaten**, e.g. pizza, but on a raw agricultural commodity (RAC), e.g. tomato, onion etc. The food on which the concentration measurements have been made is termed the **modelled food**. MCRA implements a *recursive search algorithm* to link foods-as-eaten to modelled foods. This means that there can be intermediate steps, e.g. if unpeeled *apple* and *grapes* are the modelled foods, the food-as-eaten *apple pie* contains *peeled apple* and *raisins, peeled apple* is linked to unpeeled *apple*, and *raisins* are dried *grapes*. *Peeled* and *dried* are the *processing types*.

Food classification: FoodEx1

Food code definition

In MCRA, a food code is a string consisting of symbols. Some special symbols (., \$, -, #) are reserved for special use (see below), and can not be used freely in own codes.

Codes can be hierarchical. Any code can be followed by \$ or . plus a subtype code. This can be repeated any number of times, e.g. A\$B\$C\$D, or A.B.C.D.

Codes can specify the food processing type (e.g. peeling). Any code can be followed by a hyphen ('-') plus a processing type code (e.g. FP0226-2). More than one level of processing code is allowed (e.g. FP0226-2-13). Subtype codes should precede processing codes (e.g. NL005\$123\$456-2).

Food codes in consumption surveys

Any coding system for foods-as-eaten can be used in MCRA. For example, in Europe EFSA develops a Food Classification and Description System for exposure assessment named FoodEx 2 (EFSA (2011a), EFSA (2011b)), featuring a hierarchical system of a core list of foods, an extended list, and domain-specific hierarchies.

Food codes in concentration data

Any coding system for modelled foods can be used in MCRA.

Food processing

Concentrations of substances in foods may change when foods are processed. Examples of *processing types* are peeling (e.g. of apples), cooking (e.g. of spinach), drying (e.g. of grapes), juicing (e.g. of oranges). In MCRA a processing factor can be specified for any food. Processing factors specify the ratio of concentrations in the processed and unprocessed food. The food code of the processed food (e.g. FP0226-2) will be converted to the food code of the unprocessed food will then be multiplied by the processing factor. Special attention is needed when food processing also includes changes of the weight of the food. Traditionally, processing factors combine the effects of chemical alteration and weight change, so the weight change should not be double-counted. The *processing correction factor* is introduced to correct processing factors that combine both effects, e.g. when 100g *raisins* (dried grapes) are translated to 300g *grape* (modelled food) and the processing factor for drying combines both effects, the processing correction factor is 3.

Recipes and food translation

Recipes specify the composition of composite foods, e.g. *pizza*, in terms of relevant ingredients, e.g. 100g pizza contains 10g *tomato*, 5g *cheese* and 50g *flour*. Recipes are also used to specify weight changes, e.g. to obtain 100g *raisins* (dried grapes) 300g of the modelled food *grape* is needed, see also *processing correction*.

A special use of recipes and food translation is found in *Total Diet Studies*. Here, the composition of a Total Diet Study food is specified, e.g. TDS-food *FruitMix* is composed of *apple*, *orange* and *pear* with a default translation proportion of 100%. So in MCRA, the food-as-eaten *apple* is converted to *FruitMix* (100%) and *FruitMix* is considered as the modelled food (TDS-food). A conversion from *apple-pie* (food-as-eaten) to *FruitMix* (modelled food) is based on a recipe for apple-pie and a TDS composition for FruitMix.

Another use of converting foods (as-eaten or as an intermediate step), is through the specification of so-called food extrapolations (read across translations), e.g. for *pineapple* no measurements are found but by specifying that *pineapple* is converted to *FruitMix* (with a default proportion of 100%), the TDS sample concentration value of *FruitMix* will be used for *pineapple* (as-eaten or as ingredient).

Market shares and brand loyalty

Sometimes measurements of substances in food are available at a more detailed food coding level than consumption data. For example, measurements may have been made for specific brands of a food whereas the consumption survey did not record the brand. MCRA allows to specify market share data for subtypes of a food (e.g. A\$1, A\$2, A\$3 are three brands of food A), and to calculate acute exposure based on such *market shares*.

Supertypes

Sometimes measurements of substances on food are available at a less detailed food coding level than consumption data. MCRA allows to use the concentration data of a supertype for all underlying food codes. However, this is not the default, and an explicit permission should be given to allow this feature.

Maximum Residue Levels

Maximum residue levels are the upper legal levels of a concentration for substance residues in a food, e.g. pesticide, or feed based on good agricultural practices and to ensure the lowest possible consumer exposure.

MCRA food code conversion algorithm

The conversion algorithm links food as eaten codes to modelled food codes using a 7-step procedure.

Food classification: FoodEx2

'The collection and evaluation of data on levels of chemical occurrence or presence of biological agents in food and feed are important tasks of EFSA. By combining the data with information on food consumption allows for detailed intake and exposure estimates crucial to any food and feed safety risk assessment or nutrient adequacy analysis. The EU Member States provide an increasing volume of data to EFSA and other European bodies. To provide a common link to all the diverse food and feed databases, a system for the unique and universal identification and characterisation of food and feed items is essential. EFSA has developed a preliminary standardised food classification and description system called FoodEx2 (version 2 of the EFSA Food Classification and Description System [FCDC] for exposure assessment). The system consists of descriptions of a large number of individual food items aggregated into food groups and broader food categories in a hierarchical parent-child relationship. Central to the system is a common 'core list' of food items or generic food descriptions that represent the minimum level of detail needed for intake or exposure assessments. More detailed terms may exist in addition to the core list and these are identified as the 'extended list'. A parent-child relationship exists between a core list food item and its related extended list food items. The terms of the core and extended list may be aggregated in different ways according to the needs of the different food safety domains. In the present version four hierarchies are proposed: three domain-specific and a general purpose one. Facets are used to add further detail to the information provided by the food list term. Facets are collections of additional terms describing properties and aspects of foods from various perspectives'. For more information visit: http://www.efsa.europa.eu/en/datex/datexfoodclass.htm.

For MCRA, having a different set of food codes is in itself not a problem. That is, for MCRA, it does not matter how foods are coded, as long as they can be linked to consumptions and concentrations within an exposure assessment. What makes FoodEx2 different from other food coding systems is that it provides additional food hierarchies, food facets, and a combined food/facet coding system. Below follows a brief summary of these main features of the FoodEx 2 coding system from the perspective of exposure assessment using MCRA.

Foods and food hierarchies

FoodEx 2 contains different food hierarchy definitions and allows for creation of custom food hierarchy definitions. These hierarchies could, for exposure assessment, allow to assess intake or consumption data based on the groups defined by these hierarchies.

Code	Level	Name	ParentCode	Scopenotes
A000J	1	Grains and grain-based products	ROOT	The category covers all
A000K	2	Cereals and similar	A000J	
A0001	3	Cereal and cereal-like grains	A000K	
A000M	4	Amaranth grain	A000L	
A000N	5	Buckwheat grain	A000L	
A000P	6	Barley grain	A000L	

Table 3.11: Food hierarchy export from FOODEX 2.0 Browser version 0.1.3

Facets and facet descriptors

FoodEx 2 allows to provide supplementary details on specific aspects of foods by means of so-called facets and facet descriptors. Facets are collections of terms defining specific characteristics of food from particular points of view and facet descriptors describe specific characteristics foods. For example, *processing technology* is a facet, and *baking* is a facet descriptor belonging to this facet. Currently, 26 facets are defined, containing in total 2172 descriptors see EFSA (2011b). Facets are also defined in a hierarchical system. For instance, *cooking in fat (A07GR)* and *baking (A07GX)* are sub-items of the descriptor *cooking and similar thermal preparation processes (A0BA1)*. Facets are coded as small strings that consist of a facet code and a facet descriptor code separated by a '.'-character. For example, the facet code *F28.A07GX* holds

- 1. the facet code F28, which is the facet code for process technology, and
- 2. A07GX , which is the descriptor code for baking.

Table 3.12: Part of the FoodEx 2 facet descriptor codes of the source facet (F01).

Code	Level	Name	ParentCode	Scopenotes
A04SF	1	Animals	ROOT	
A056H	2	Mammals (food source animal)	A04SF	
A056Z	3	Farmed / non-game mammals (food source animal)	A056H	
A057A	4	African buffalo (food source animal)	A056Z	
A057B	4	American buffalo (food source animal)	A056Z	
A057C	4	Buffalo (food source animal)	A056Z	
A057D	4	Cape buffalo (food source animal)	A056Z	
A057E	4	Cattle (food source animal)	A056Z	

Implicit facets

Implicit facets are facets of a product that are already implied by the food product itself. Consider, for example, *potato boiled (A011P)*, where *boiling (A011P)* is an implicit facet, because boiling is already implied by the product. According to EFSA (EFSA (2011a)) 'inclusion of implicit facets in the string recorded for each food database record is not encouraged' and it is suggested to identify and record the implicit facet descriptors in a separate table.

Foods as facets

Foods and facet descriptors share the same unique alphanumerical coding system; in some cases, like *characterising ingredient or sweetening agent* food list elements may be used as facet descriptors.

The FoodEx 2 coding system

In the coding system, facets can be added to the primary food codes to provide supplementary detailed information of particular data records. The structure of the FoodEx 2 codes is:

idFood#idFacet.idFacetDescriptor\$idFacet.idFacetDescriptor\$....

The code starts with the primary FoodEx2 food code. Then, when there are supplementary facets, the food code is followed by a '#'-character and the facets string. The facets string is constructed as a concatenation of the individual facets strings, separated by means of the '\$' character. As an example, consider the string *A011P#F28.A07GL\$F28.A07KQ* which is composed of:

- Food: A011P Potato boiled
- Facet 1: F28.A07GL Process technology Boiling
- Facet 2: F28.A07KQ Process technology Freezing

FoodEx2

For MCRA, FoodEx 2 introduces the following points of attention:

- Reading and dealing with FoodEx 2 coded data sets
- Reading and dealing with food facets
- Reading and exploiting food hierarchy data

Reading and dealing with FoodEx 2 codes

All data entities that contain foods data are potentially affected by the introduction of FoodEx 2. In MCRA, the following data tables are adapted to allow for input of full FoodEx 2 food codes:

- Foods
- Consumptions
- Concentrations

For these tables, the food code is allowed to be the complete FoodEx 2 food code and automatically recognized as such. As an example, Table 3.13 shows how the FoodEx 2 coded consumptions should be provided to the system. On important note: the maximum field length of the food code is 50. This means that there is a maximum of five facets that can be specified for a food.

Individual	DayOfSurvey	Food	Amount	FoodSurvey
14233701	1	A011R# F28.A07GX	153.43	FS01
18843004	1	A011R# F28.A07GX	125.23	FS01
34025701	1	A011R# F28.A07GX	153.60	FS01
14720005	2	A011R# F28.A07GX	105.00	FS01
49174010	1	A011R# F28.A07GX	140.00	FS01
62794010	1	A011R# F28.A07GX	67.00	FS01
61392002	1	A011P# F28.A07GL\$F28.A07KQ	104.72	FS01
61281231	1	A011P# F28.A07GL\$F28.A07KQ	109.72	FS01

Table 3.13: Integrated coding of the facets in the consumed foods field of
food consumptions. Implementation.

Reading and dealing with facets data

Within MCRA, the following facets related aspects are accounted for:

- Reading facets data
- · Dealing with facets
- · Facets in concentration data
- Facets in food conversion
- Using facets as processing factors
- Using hierarchy data in the output

Reading facets data

To incorporate input of facets data in MCRA, two tables Facets and FacetDescriptors are introduced as optional tables of the Foods data group. See table *Facets* and *Facet descriptors*.

Within MCRA, the facets of FoodEx 2 coded foods, consumptions, and concentrations are automatically linked to the provided facets and facet descriptors. Also, the facet descriptor names are added automatically to the foods containing these facets.

Dealing with facets

The introduction of food facets allows for much more detailed specifications of consumption and concentration data. However, it introduces the problem of deciding on which level of detail the exposure assessment should be performed. That is, should concentration models be generated on the level of foods-without-facets or on the level of foods-with-facets? E.g., should the concentrations of *clementine peeled (A01CE#F28.A07LC)* and *clementine unprocessed (A01CE#F28.A07CS)* be modelled separately or should one model be constructed for *clementine (A01CE)*? Treating all clementine's as equal may yield over-simplified conversions, whereas treating all separately may lead to many concentration models based on only few measurements. In MCRA, no implicit grouping of concentrations of equal foods with different facets is applied. If concentrations are provided for both *clementine peeled (A01CE#F28.A07LC)* and *clementine unprocessed (A01CE#F28.A07LC)* and *clementine unprocessed (A01CE#F28.A07LC)* and *clementine unprocessed (A01CE#F28.A0C0S)*, then these are modelled separately. Another question is whether the order of the facets is relevant or not. E.g., is *A0BYV#F02.A06GF\$F03.A06HY* the same as *A0BYV#F03.A06HY\$F02.A06GF*? Regarding this matter, MCRA considers the facet order to be important. I.e., *A0BYV#F02.A06GF\$F03.A06HY* is not the same as *A0BYV#F02.A06GF*.

Facets in food conversion

For conversion of foods-as-eaten to modelled foods, MCRA considers foods with different facet strings as different foods. I.e., there is no implicit conversion of foods-with-facets to foods-without-facets and also the order of the facets is important. However, as it is realistic to convert food-with-facets to the base food without facets, an additional (explicit) conversion step remove-all-facets is added that converts foods with facets to the base foods. I.e., the action is "remove all". There is no conversion step for "stripping off one facet at a time". The reason for this is that there

is no good way of deciding which facet to strip off first. This new conversion step is somewhat equivalent to the already existing default processing conversion step (step 6), and is therefore implemented as step 6b of the conversion algorithm. Particular rules followed by this step:

• Conversion of food-with-facets to food-without-facets.

Using facets that reveal processing data

Facets containing processing information, such as *part-consumed-analysed* (F20) and *processing technology* (F28) could be integrated with processing data. As an example, consider *clementine peeled* (A01CE#F28.A07LC). This could be linked to *clementine* (A01CE), with processing type *removal of external layer* (A07LC). Linking to processing data could be achieved by entering processing data using the facet codes. As an alternative to the current processing factor tables, a facet-based processing factors table is defined for processing facets. That is, the codes for food processed and unprocessed are implicitly defined for FoodEx 2.

Table 3.14: Example of a MCRA processing factors table using FoodEx 2 foods and facets codes.

FacetCode	Substance	FoodCode	ProcNom	ProcUpp	Proc- NomUnc- Upp	Proc- UppUnc- Upp
A07LC	SubstanceX	A01CE	0.5	0.6	0.05	0.06
F28.A07GV	SubstanceX	A0BY	0.2	0.1	0.03	0.04

Note that in the example, the facet code could be specified as the full facet code, or just the code of the facet descriptor. As a more elaborate example consider

French fries from cut potato (A0BYV#F02.A06GF\$F03.A06HY\$F04.A00ZT\$F28.A07GR)

For this food code, the substring of the processing facet is extracted from the list of facets.

- A0BYV#F02.A06GF\$F03.A06HY\$F28.A07GR\$F04.A00ZT with processing facet link A07GR
- *A0BYV#F02.A06GF\$F03.A06HY\$F04.A00ZT*

In MCRA, a table *Processing factors* is introduced that allows for specification of processing factors by means of facets. This table has the following structure:

Column name	Key	Required	Туре	Size	Description
idProcessingType	Yes	Yes	String	5	The facet code of this processing factor definition. May be specified as full facet code, i.e., facet code plus facet descriptor code, or as the facet descriptor code.
idFood	Yes	Yes	String	200	The food code
idCompound	Yes	No	String	50	The substance for which this processing factor is defined.
Nominal	No	Yes	Double		Nominal value (best estimate of 50th percentile) of processing factor (defines median processing factor)
Upper	No	Yes	Double		Upper value (estimate of 95th percentile or "worst case" estimate) of processing factor due to variability
NominalUncertaintyUpper	No	Yes	Double		Upper 95th percentile of nominal value (Nominal) due to uncertainty. A standard deviation for uncertainty of the nominal value (Nominal) is derived using the nominal value (Nominal) and upper 95th percentile (NominalUncertaintyUpper)
UpperUncertaintyUpper	No	Yes	Double		Upper 95th percentile of upper value (Upper) due to uncertainty. From the nominal value (Nominal), upper value (Upper) and the specified uncertainties of these values (NominalUncertaintyUpper and UpperUncertaintyUpper, respectively) the degrees of freedom of a chi-square distribution describing the uncertainty of the standard

Table 3.15: Table FacetDescriptors of the Food data group.

The integration with the food conversion algorithm is as follows: Conversion step 2 (*processing*) is extended with a step 2c (*processing facet*) that attempts to match facets of a food code to processing data provided in the processing facets table. The following important rules are followed:

- Processing factors can be defined for base-food-code/facet-code combinations and translate as food-withprocessing-facet to food-without-processing-facet.
- If multiple processing facets are present in the food-as-eaten code, then the last processing facet is used first for conversion.
- Facet processing factors can be specified using the full facet code (i.e., facet-code plus facet-descriptor-code) or just the facet descriptor code. If both are specified for the same food, the full facet code is used.
- Facet processing factors can be defined substance-specific, and non-substance-specific. Processing factors that are defined substance-specific always precede non-substance specific processing factors.
- Processing factors defined by a food-processed/food-unprocessed combination precede processing factors defined through facets.

Weight reduction factors for processing factors defined for facets should be included in table *Recipes* should match exactly.

Food hierarchies

Reading and dealing with food hierarchy data

Within MCRA, the following hierarchy related aspects are accounted for:

- Reading food hierarchy data
- Using hierarchical data for conversion of foods
- Using hierarchy data in the output

Reading food hierarchy data

A new data group named *Foods data formats* is added. In this group, a new table *Food hierarchies* is used for input of food hierarchies. This table contains food hierarchy node-definition records that reflect a hierarchical structure. For foods that are not in this list as idFood, it is implicitly assumed that these foods are root items.

Note: It is common practice to describe hierarchies using tree structures. Here, the elements of the tree are named *nodes*, the lines connecting the nodes are named *branches*, and nodes without children are *leaf nodes/end-nodes*. This terminology is also used throughout the remainder of this document.

Using food hierarchies for food conversion

The introduction of the hierarchy structure allows for integration with step 4 and step 5 of the food conversion algorithm; the *subtype* and *supertype* linking steps. That is, when no concentration data is found for a certain product, the concentration data of a (according to the hierarchy) related product could be used. In MCRA, the *supertype* conversion step also contains a *hierarchy-supertype* step based on the food hierarchy.

Supertype link (step 5):

- a) Supertype: Try to find supertypes base on '\$'-coded strings, e.g., 'xxx\$yyy' is converted to 'xxx'
- b) **Hierarchy-supertype**: try to find the supertype of the current food based on the food hierarchy (i.e., convert the current food to its parent).

Note 1: the supertype conversion step is optional and should be specified in the conversion settings panel.

Note 2: the *hierarchy-supertype* step only applies for foods-without-facets. The reason for this is that for the conversion, the base type of a food-with-facets can be considered as a better conversion candidate than the parent food with the same facets.

Using hierarchy data in the output

Food hierarchy information could be used in presentation of various tables of the output of MCRA. That is, in the tables in which foods data is presented, these records could be grouped based on the hierarchy and/or a tree-like display can be built for the presentation of this data. Tables that are candidate for being extended are, for example, the input data tables foods-as-eaten/modelled foods and the exposure by food-as-eaten/modelled food output tables.

Summarizing over the food hierarchy is many cases not a straightforward task. Consider, for instance, the statistic *number of consumption days* given the artificial hierarchy of *Citrus Fruits* containing two child-nodes *Mandarin* and *King Mandarin*: the number of consumption of *Citrus Fruits* is not "just" the sum of the consumption day of *Mandarin* and *King Mandarin*. A difficulty for summarizing based on a hierarchy arises when a node contains both data and child-nodes with data. E.g., concentrations are defined on the level of *Citrus Fruits* and on the level of *Mandarin*. In this case, the hierarchy view should ideally summarize for both *Citrus Fruits* as data record and *Citrus Fruits* as summary node. An additional complication is the status of facet-coded foods within the hierarchy. In a hierarchical view, foods-with-facets should ideally be added to their base-foods for visualization.

In MCRA, an alternative view (treetable) is added that can display hierarchical data. This alternative view is used to present a hierarchical view based on the foods hierarchy for the consumption input summary tables food as eaten and modelled food. The data summary methods for these tables are updated such that the data is also summarized per hierarchy-node.

If a node contains both data and a child record, then this node is split-up in two nodes: a summary node that summarizes the data of the node and all of its child nodes, and a data record with the string "(unspecified)" added as a child

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Food name	Food code	Mean consumption (g)	Mean consumption days (g)	Consumption days	n Percentage consumption days	Total weights consumption days	Percentage total weights consumption days
 Fruit and fruit products 	A01BS	167	200	5	83.3 %	5.0	83.3 %
Fresh fruit	A04RK	167	200	5	83.3 %	5.0	83.3 %
 Starchy roots or tubers and products thereof, sugar plants 	A00ZR	100	600	1	16.7 %	1.0	16.7 %
Starchy root and tuber products	A011B	66.7	400	1	16.7 %	1.0	16.7 %
 Processed root and tuber products 	A04MJ	66.7	400	1	16.7 %	1.0	16.7 %
Potato boiled	A011P	66.7	400	1	16.7 %	1.0	16.7 %
 Potato boiled Tuber (as part-nature) 	A011P#F02.A067V	16.7	100	1	16.7 %	1.0	16.7 %
 Potato boiled Tuber (as part-nature), Potatoes, Boiling 	A011P#F02.A067V\$F27.A00ZT\$F28.A07GL	16.7	100	1	16.7 %	1.0	16.7 %
 Potato boiled Tuber (as part-nature), Potatoes, Boiling 	A011P#F02.A067V\$F28.A07GL\$F27.A00ZT	16.7	100	1	16.7 %	1.0	16.7 %
 Potato boiled Tuber (as part-nature), Potatoes, Boiling, Baking 	A011P#F02.A067V\$F27.A00ZT\$F28.A07GL\$F28.A07GX	16.7	100	1	16.7 %	1.0	16.7 %
 Starchy roots and tubers 	A00ZS	33.3	200	1	16.7 %	1.0	16.7 %
 Tubers 	A04MC	33.3	200	1	16.7 %	1.0	16.7 %
 Potatoes 	A00ZT	33.3	200	1	16.7 %	1.0	16.7 %
 Potatoes Potatoes (food source plant), Tuber (as part-nature) 	A00ZT#F01.A05KG\$F02.A067V	16.7	100	1	16.7 %	1.0	16.7 %
 Potatoes Potatoes (food source plant), Tuber (as part-nature), Baking 	A00ZT#F01.A05KG\$F02.A067V\$F28.A07GX	16.7	100	1	16.7 %	1.0	16.7 %

Figure 3.2: Hierarchy view for the foods as eaten input summary table.

of this summary node. See Figure 3.2 for an example (*Citrus Fruits versus Citrus Fruits (unspecified)*). In MCRA, foods-with-facets are added as child nodes of the foods-without-facets.

Food unit weights

Food unit weights specify the standard weights of food units. E.g., the standard weight of an apple. This unit weight may be specified as the weight of the whole food (raw agricultural commodity/RAC) or the weight of the edible portion (EP), e.g., without peel. Unit weights are specified in the table *Food unit weights* and used in combination with *unit variability factors* to account for unit-to-unit variation in concentrations between single units of the same food in *single value dietary exposures assessments* and (*individual*) *dietary exposures assessments*.

Food unit weights can be location specific or specified as overall (default) unit weights. For some models, e.g., the *IESTI model*, location specific unit weights are preferred over overall unit weights. The overall unit weights are then used when no location specific uses are available. For other methods, only overall unit weights are used. If, for a food, an overall unit weight is not available, but there are location specific unit weights available, then the overall unit weight is computed as the average weight of the location specific unit weights (similar to EFSA PRIMo revision 3 EFSA (2018)).

Note

Note that in earlier versions of the software, food unit weights were specified in the *table for Food properties*. Although this is still possible, the recommended way of specifying unit weights is in the *table for Food unit weights*. If, for a food, unit weights are specified in both tables, then the unit weights specified in table *Food unit weights* have priority. The unit weights specified in table *Food properties* are then only used as fallbacks for the overall unit weight when no overall unit weight is specified in the *table for Food unit weights*.

3.1.3 Consumer products

Consumer products are uniquely defined sources of (personal care) products.

Output of this module is used by: Consumer product concentrations Internal exposures Consumer product use frequencies Consumer product exposure determinants Consumer product exposures

Consumer products from data

Consumer products data formats

Consumer products are primary entities of the data model.

Download empty dataset template: Zipped CSV Excel

ConsumerProducts

Each consumer product is identified by a unique code (id), a name, description and/or product group (parent). Consumer products codes are not harmonised.

Name	Туре	Description	Aliases	Required
Id	AlphaNumeric (50)	The unique identification code of the consumer product.	idConsumer- Product, Consumer- ProdufctId, Id	Yes
Name	AlphaNumeric (100)	The name of the consumer product.	Name, Consumer- ProductName	No
Description	AlphaNumeric (200)	Description of the consumer product.	Description	No
idGroup	AlphaNumeric (50)	Identifier of the parent product (i.e., the product group of which this product is a part).	idGroup, GroupId	No

Table 3.16:	Table definition for ConsumerProducts.	
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Accepted table names: ConsumerProducts, ConsumerProduct.

Consumer products as data

Consumer products definitions are provided as lists/catalogues of food definitions or hierarchy definitions.

- Consumer products data formats
- Consumer products from data

3.1.4 Non-dietary exposure sources

Non-dietary exposure sources are the sources containing chemical substances to which individuals in a population are exposed via any of three non-dietary routes: dermal, inhalation or oral, per day.

Non-dietary exposure sources from data

Non-dietary exposure sources data formats

Non-dietary exposure sources are defined in the non-dietary sources table.

Download empty dataset template: Zipped CSV Excel

Non-dietary exposure sources

Each non-dietary exposure source is identified by a unique code in a code system of choice, a name, and a description.

Name	Туре	Description	Aliases	Required
idNonDietary- ExposureSource	AlphaNumeric (50)	The unique identification code of the non-dietary exposure source.	idNonDietary- ExposureSource, idSource, Code, Id	Yes
Name	AlphaNumeric (100)	The name of the non-dietary exposure source.	Name	No
Description	AlphaNumeric (200)	Description of the non-dietary exposure source.	Description	No

Accepted table names: NonDietaryExposureSources.

Non-dietary exposure sources as data

Non-dietary exposure sources are lists/catalogues of definitions of non-dietary sources through which people can be exposed to chemical substances (e.g., personal care products and plant-protection products).

- Non-dietary exposure sources data formats
- Non-dietary exposure sources from data

3.1.5 Populations

Populations are groups of human individuals that are the scope of exposure or risk assessments. Optional descriptors of populations are location (e.g. a country), time period (with a start and end date), age range (with a minimum and maximum age) and gender. Example: the French population in 2005-2007 (= time period) of women (= gender) of child-bearing age 18-45 yr (= age range).

Output of this module is used by: Consumptions Single value consumptions Concentrations Consumptions by modelled food Dietary exposures Single value dietary exposures Single value non-dietary exposures Non-dietary exposures Internal exposures HBM concentrations HBM analysis Biological matrix concentration comparisons Dust exposures Soil exposures Air exposures Individuals Consumer product use frequencies Consumer product exposures Hazard characterisations Burden of disease Risks Single value risks Environmental burden of disease

Populations from data

Populations calculation

In an exposure or risk assessment, the population of interest is implicitly or explicitly defined. An implicit definition is made by selecting one of the available food consumption surveys in the **Compute** option. For example, by selecting the food survey 'NL-Toddlers' with corresponding consumption data, the Dutch population of toddlers is implicitly considered as the population of interest.

After selecting a survey, the population is further defined by checking *Define population based on specified individual properties* in the *Population definition from dietary surveys panel*. Include implicitly defined individual(day) properties in the population definition. E.g., the Dutch population of toddlers is restricted to females by including the property *gender* with level *Female'* in the population definition. The implicitly defined population becomes the Dutch female toddlers.

Although implicit definition of the population of interest works fine, there is a need to make more explicit that the focus of an assessment is on the population. The user should be aware that assessments are about assessing the exposure or risk in a specific population of interest. This becomes even more urgent when dietary exposures are combined

with non-dietary exposures. Without explicitly specifying the population of interest, the exposure of Dutch toddlers may unintentionally be combined with non-dietary exposures of e.g. adult operators. The explicit definition of a population is made by selecting a population datasource in the **Use data** option. By specifying population properties like age, gender or any other property in the data, the population may be further restricted.

For the **Compute** option:

• Setting *Population definition from dietary surveys* Check *Define populations based on specified individual properties* for including individual properties.

When the population of interest is explicitly defined, (Use data option), the population definitions are based on the data in the *IndividualProperties table* referring to populations in the *Populations table*..

Population definition from dietary surveys

This panel is only available when the population of interest is implicitly defined, use the *Compute* option. Currently, population definitions are based on dietary surveys. It is foreseen that the population definition will be extended as soon as Human Biomonitoring (HBM) data and/or non-dietary survey data become available.

Specify the nominal population bodyweight, needed for single value calculations, e.g. 70 kg (default).

After checking *Define populations based on specified individual properties* in Figure 3.3, include individual properties for the population definition.

Select the requested levels for each property.

See also the *consumption panel documentation* for further information on population definition based on a selection of individual properties.

For individual day properties, an extra option is available to include the records without date data. Currently, the default value is to include records without dates.

Populations data selection

This option is only available when the population of interest is explicitly defined, use the *Use data* option and select a data source. Select one of the available populations in the *Populations selection* pane.

Selecting two or more populations, initiates a loop over multiple populations. Check the *Loop over multiple populations* checkbox. See also Figure 3.8

Populations data formats

Populations are primary entities of the data model.

Download empty dataset template: Zipped CSV Excel

Populations

Populations identify human groups in e.g. dietary, nondietary and human monitoring surveys. Optionally, a name and description can be added. Specify a standard bodyweight (optional) as a descriptor of the average or nominal bodyweight in the population. Use table PopulationIndividualPropertyValues to specify descriptors/properties that characterise the population. These population individual property values will be used to restrict the population to e.g. a certain time period (through specifying a start and end date, both dates are inclusive), age (through specifying a minimum and maximum age, both limits are inclusive) or gender (male or female). To facilitate the user, dynamic properties may be added to the table. Three kind of additional properties are available: alphanumeric properties (property name and level), numeric properties (a range through specifying a minimum and maximum using suffixes Min and Max, both bounds are inclusive) and datetime properties (a range through specifying a start and end date using prefixes Start and End, both dates are inclusive). Dynamic or additional properties are ignored when table PopulationIndividualPropertyValues is present in the upload. In table IndividualProperties each property used in the Populations table is described.

Consumptions / Populations	08d837d3 🕨 🕒
Populations	0
Use data 🌣 Compute	
Population settings	Save Changes
Nominal population bodyweight (kg) 70	0
Population definition from dietary surveys	Save Changes
Define populations based on specified individual properties	6
Include breastfeeding in population definition	
Include gender in population definition	
Include haircolor in population definition	
Include region in population definition	
Include age in population definition	
Include education in population definition	
Include height in population definition	
Include month in population definition	0

Figure 3.3: Define populations by including individual properties.

Define populations based on specified individual properties						
Include breastfeeding in population definition						
 Include gender in population definition Selected values * Female (398) 		-				
Include haircolor in population definition						
 Include region in population definition Selected values * East (156), North (80), South (165) 						
 Include age in population definition Lower bound (min. 0) 0 	Upper bound (max. 4) 5					

Figure 3.4: Check *gender*, region and age and select the requested levels Female, East, North and South and age min = 0, age max = 5, resp.

Dietary exposures / Populations	0716d970	×	¢
Populations			?
Use data 🌣 Compute			
Populations data source			
> Populations-DE.zip v2			
Populations selection			
Populations: selected 1 of 17 (clear filter) (change selection)		-	

Figure 3.5: Population data selection (Use Data).

×

Populations selection

Sho	w selected	Q Filter	
1 entrie	s selected		
	Code	Name	
\checkmark	DE-Children-2001-2002-East	German children 2001/2002 region East	
	DE-Children-2001-2002	German children 2001/2002	
	DE-Children-2001-2002-Conventional	German children 2001/2002 conventional	
	DE-Children-2001-2002-East-Summer	German children 2001/2002 region East summer season	
	DE-Children-2001-2002-East-Winter	German children 2001/2002 region East winter season	
	DE-Children-2001-2002-North	German children 2001/2002 region North	
	DE-Children-2001-2002-North-Summer	German children 2001/2002 region North summer season	
		Cancel Sav	ve

Figure 3.6: Available population data: change selection.



Figure 3.7: Loop over multiple populations (Use Data).

				•••	
Ð	ρ	2		IT	•
	•	-	•		•

ults						
Output		Status	Message	Date	Running time	
Dietary exposures	/	Running	1 running, 3 waiting for activation		-	(
German children 2001/2002 region East (DE-Children-2001-2002-East)	1	Running	Summarizing food conversion		-	1
German children 2001/2002 (DE-Children- 2001-2002)		Waiting for activation	Only one active job per user is allowed. Waiting for other job(s) to complete		-	•
German children 2001/2002 region East summer season (DE-Children-2001-2002- East-Summer)	/	Waiting for activation	Only one active job per user is allowed. Waiting for other job(s) to complete		-	•
German children 2001/2002 region East winter season (DE-Children-2001-2002- East-Winter)	/	Waiting for activation	Only one active job per user is allowed. Waiting for other job(s) to complete		-	•

Figure 3.8: Results pane with loop over multiple populations (Use Data).

Name	Туре	Description	Aliases	Required
idPopulation	AlphaNumeric (50)	Unique identification code of the population.	IdPopulation, PopulationId, Code, Id	Yes
Name	AlphaNumeric (100)	The name of the population.	Name, PopulationName	No
Description	AlphaNumeric (200)	Description of of the population.	Description	No
Location	AlphaNumeric (50)	Location.	Location, Country	No
StartDate	DateTime	Starting date of the specific time window marking this population.	StartDate	No
EndDate	DateTime	End date of the specific time window marking this population.	EndDate	No
NominalBody- Weight	Numeric	Nominal body weight (in kg) of the individuals of this population.	NominalBody- Weight, BodyWeight	No
Size	Numeric	The number of individuals of this population.	Size, PopulationSize	No
Additional individual properties, type = Alpha- Numerical		AlphaNumerical population properties specifying a level or levels (comma separated), [property name]. E.g. for individual property [Region] specify a region like [North] or [South]. Note that table IndividualProperties should contain the property Pagion		No
2		contain the property Region with PropertyLevel = Individual and Type = Categorical. For other type of properties use Type =	Chapte	er 3. Modules

Table 3.18:	Table	definition	for	Populations
-------------	-------	------------	-----	-------------

Accepted table names: Populations, Population.

Individual properties

This table is used to describe the properties used in the Populations or PopulationIndividualPropertyValues table characterising the population (table Populations) and/or the properties used in the Individuals table characterising an individual. Properties like Age, Gender, Region are describing an individual (PropertyLevel = Individual). Properties like Period (for populations) or Month (sampling date for an individual day) are describing an individual day (PropertyLevel = IndividualDay).

Name	Туре	Description	Aliases	Required
idIndividual- Property	AlphaNumeric (50)	The code of the property.	idIndividual- Property, Individual- PropertyId, Individual- Property	Yes
Name	AlphaNumeric (100)	The name of the property.	Name	No
PropertyLevel	<i>PropertyLevelType</i>	The level of the property. This type follows a controlled terminology, with possible values: Individual or IndividualDay.	PropertyLevel, LevelProperty	No
Description	AlphaNumeric (200)	Description of the property.	Description	No
Туре	IndividualProperty- Type	This field specifies the type of the values of this individual property. This type follows a controlled terminology, with possible values: Boolean, Categorical (default), Numeric, Nonnegative, Integer, NonnegativeInteger, Month, Datetime, Gender.	Туре	No

Table 3.19: Table definition	for Individual j	properties.
------------------------------	------------------	-------------

Accepted table names: IndividualProperties, IndividualProperty.

Population individual property values

This table describes population individual properties, such as Age, Gender, Period, Region or Breastfeeding. Population individual property values are used to restrict the population to, for example, a range of ages, a gender, a certain time period, a geographical location or women who are breastfeeding. For numerical properties, use Min-Value and MaxValue to specify a range. To specify a time period, use StartDate and EndDate. For Gender, Region and Breastfeeding use Value to specify the gender, categorical level or boolean value, respectively. In the IndividualProperties table, each property used in the PopulationIndividualPropertyValues table is described. NOTE: the individual property ID (idIndividualProperty) must be unique for a population (specified by idPopulation), i.e., the combination idPopulation - idIndividualProperty may occur only once in the table Population individual property values. You may list multiple values in the 'Value' field by separating them with a comma (e.g., 'Summer, Autumn').

Name	Туре	Description	Aliases	Required
idPopulation	AlphaNumeric (50)	The code of the population to which the property is attached. The provided population code should match with a code of the populations table.	idPopulation, PopulationId, Population, PopulationCode, Code	Yes
idIndividual- Property	AlphaNumeric (50)	The name or reference of the individual property.	idIndividual- Property, Individual- PropertyId, Individual- Property	Yes
Value	AlphaNumeric (50)	The value(s) of the property. Multiple values should be comma separated.	Value	No
MinValue	Numeric	Minimum value of the value of the property for a numeric range.	MinValue, ValueMin	No
MaxValue	Numeric	Maximum value of the value of the property for a numeric range.	MaxValue, ValueMax	No
StartDate	DateTime	Start date value of the property for a time period.	StartDate, MinDate	No
EndDate	DateTime	End date value of the property for a time period.	EndDate, MaxDate	No

Table 3.20:	Table	definition	for	Population	individual	property	values.
				1		1 1 2	

Accepted table names: PopulationIndividualPropertyValues, PopulationIndividualPropertyVal, PopulationPropertyValues.

Population characteristics

Characteristics describing specifics of a population, e.g. IQ or average birth weight.

Name	Туре	Description	Aliases	Required
idPopulation	AlphaNumeric (50)	Unique identification code of the population.	IdPopulation, PopulationId	Yes
Characteristic	Population- CharacteristicType	Population characteristic type.	Characteristic, idCharacteristic, Population- Characteristic, IdPopulation- Characteristic	Yes
Unit	AlphaNumeric (50)	Unit of the population characteristic.	Unit, Characteristic- Unit	No
Value	Numeric	Mean of the distribution.	Mean	Yes
Population- Characteristic- Distribution- Type	Population- Characteristic- DistributionType	Distribution type (Normal, Uniform, or LogNormal). If not specified, the characteristic is assumed to be constant.	Distribution- Type, Population- Characteristic- Distribution- Type, Distribution	No
CvVariability	Numeric	Coefficient of variation.	CvVariability, CV	No

Table 5.21. Table definition for 1 opulation characteristics.

Accepted table names: PopulationCharacteristics, PopulationCharacteristic.

Populations settings

Selection settings

Name	Туре	Description
Define populations based on specified individual properties	Boolean	Define a population by selecting specific ranges/values of individual properties. E.g., the female population between ages and 45 is composed of the properties gender (female) and age (between 18 and 45).
Individuals subset definitions	IndividualsSubsetDefinition	Contains a list of subset definitions to filter the population's individuals based on an individual's properties, by property nam and a custom filter query, for example a value range or a list of keywords.
Individual day subset	IndividualDaySubsetDefinition	Individual day subset definition.
Nominal population bodyweight (kg)	Numeric	Nominal population bodyweight in kg (needed for single value calculations).

Table 3.22: Selection settings for module Populations.

Populations as data

Populations are provided as data. When the population of interest is explicitly defined, subset selections are based on the data in the IndividualProperties and PopulationIndividualPropertyValues table.

- Populations data formats
- Populations from data

Calculation of populations

Populations are implicitly defined by the individuals as used in the dietary consumption module and/or the non-dietary module and/or the human based monitoring module.

• Populations calculation

3.1.6 Responses

Responses are measurable entities in test systems. Responses are used to represent effects (see effect representations) and their measured values are collected in dose response data.

This module has as primary entities: Test systems

Output of this module is used by: Dose response models Dose response data Effect representations

Responses from data

Responses data formats

A response is a measurable endpoint on in a test system. E.g., in a rat test system a response may be the percentage of fatty hepatocytes observed after 90 days. Responses are defined in the responses table.

Download empty dataset template: Zipped CSV Excel

Responses

Each response is identified by a unique code (idResponse) in a code system of choice, a name, and a description. Also, each response should be linked to a test system (idTestSystem) on which the response is measured. Responses can be of various types (ResponseType), e.g., ContinuousMultiplicative (= non-negative real values using a ratio scale), ContinuousAdditive (= real values using an interval scale), Ordinal, Quantal, or Binary. For continuous variables, the response unit (ResponseUnit) is also relevant. Additionally, also a reference to the test method guideline, e.g., standardised assay kit may also be specified (GuidelineMethod).

Name	Туре	Description	Aliases	Required
idResponse	AlphaNumeric (50)	Unique identification code of the response. In the EuroMix data collection, a EuroMix coding system has been set up in which the id of the test system prefixes the id of the response. E.g., 'HepaRG-PCR-PPARA', 'RatWEC-PCR-CYP26a1' and 'MouseDevelopmental- FacialPrimordia-malformed- E9'.	idResponse, ResponseId, Response, Id	Yes
CodeSystem	AlphaNumeric (100)	Identifier of the coding system of the response code.	CodeSystem	No
Name	AlphaNumeric (100)	Name of the response.	Name	No
Description	AlphaNumeric (200)	Additional description or label of the response.	Description	No
idTestSystem	AlphaNumeric (50)	Unique identification code of the test system.	idTestSystem, idSystem, SystemId, TestSystem	Yes
Guideline- Method	AlphaNumeric (200)	Reference to the test method guideline, e.g., standardised assay kit.	Guideline- Method	No
ResponseType	<i>ResponseType</i>	The data type of the response measurements (e.g., continuous multiplicative, continuous additive, binary, quantal, count, ordinal, categorical). Controlled terminology.	ResponseType	Yes
ResponseUnit	AlphaNumeric (100)	If the response type is Continuous, then this should be the unit of the response, e.g., kg.	ResponseUnit	No

Accepted table names: Responses, Response.

Responses as data

A response is a measurable endpoint defined in a test system. It has a unit and a measurement type (e.g., continuous non-negative, quantal).

- Responses data formats
- Responses from data

3.1.7 Substances

Substances are chemical entities that can refer to: 1) active substances such as investigated in toxicology; 2) measured substances such as defined in specific analytical methods. MCRA assessments can have one or more substances as the scope. When more than one substance is specified, there is an option to perform a cumulative assessment. In that case one of the substances has to be indicated as the index/reference substance, and results will be expressed in equivalents of the index substance.

Output of this module is used by: Concentrations Concentration distributions Single value concentrations Processing factors Unit variability factors Occurrence patterns Occurrence frequencies Substance authorisations Substance approvals Substance conversions Deterministic substance conversion factors Concentration limits Concentration models Modelled foods Focal food concentrations Exposure biomarker conversions Consumer product concentrations Food conversions Consumptions by modelled food High exposure food-substance combinations Dietary exposures Single value dietary exposures Single value non-dietary exposures Non-dietary exposures Internal exposures Exposure mixtures HBM concentrations HBM analysis Biological matrix concentration comparisons Dust concentration distributions Dust exposure determinants Dust exposures Soil concentration distributions Soil exposure food-substance Consumer product exposures Indoor air concentrations Air exposure determinants Air exposures Consumer product exposure determinants Consumer product exposure QSAR membership models Molecular docking models Absorption factors PBK model parametrisations Kinetic conversion factors Active substances Relative potency factors Hazard characterisations Points of departure Dose response models Dose response data Inter-species conversions Intra species factors Exposure response functions Risks Single value risks Environmental burden of disease

Substances from data

Substances data formats

Substances are primary entities of the data model. Substance intakes are of main interest in exposure assessments and the effect of intake on human health is of interest in risk assessments. In the substances table, the substance entities and other relevant substance properties that are relevant for the assessment at hand should be defined.

Download empty dataset template: Zipped CSV Excel

Substances

Each substance should have a unique identification code (idSubstance), and optionally, a name and description may be used for a more detailed description of the entity. Additional properties, such as the molecular mass (MolecularMass) and Cramer class (CramerClass) may also be specified. Example: Captan (idSubstance RF-0061-001-PPP) has MolecularMass 300.5922 and CramerClass 3.

Name	Туре	Description	Aliases	Required
idSubstance	AlphaNumeric (50)	The unique identification code of the substance. This code may be from an existing coding system, such as CAS-codes or Param codes of EFSA, or it may be a used-defined code.	idSubstance, SubstanceId, SubstanceCode, Substance, Code, Id	Yes
Name	AlphaNumeric (100)	The substance name.	Name, SubstanceName, PesticideName	No
Description	AlphaNumeric (200)	Substance description.	Description	No
Concentration- Unit	<i>ConcentrationUnit</i>	Contains a coding to determine the default unit in which concentrations for this substance are expressed.	Concentration- Unit, Unit, Reference- Concentration- Unit	No
CramerClass	Integer	The Cramer class of the substance.	CramerClass	No
MolecularMass	Numeric	The molecular (molar) mass.	MolecularMass, Mass, MolarMass, Molecular- Weight, MolarWeight	No
IsLipidSoluble	Boolean	States whether the substance is soluble in lipid $(0 = no, 1 = yes)$.	IsLipidSoluble, IsSoluble	No

Table 3.24:	Table	definition	for	Substances.
-------------	-------	------------	-----	-------------

Accepted table names: Substances, Substance.

Substances settings

Selection settings

Table 3.25:	Selection	settings	for	module	Substances.
-------------	-----------	----------	-----	--------	-------------

Name	Туре	Description
Compute cumulative exposures	Boolean	Specifies whether the assessment involves multiple substances a results should be cumulated over all substances.
Multiple substances analysis	Boolean	Specifies whether the assessment involves multiple substances.
Index substance	AlphaNumeric	The substance of interest or index substance.

Substances as data

Substances are provided as data (code, name).

- Substances data formats
- Substances from data

3.1.8 Test systems

Test systems are biological or artificial systems used for assessing hazard in relation to chemical exposure from substances in varying doses. Test systems may refer to 1) in-vivo test systems (e.g. a rat 90-day study, a human biomonitoring study); 2) in-vitro test systems (e.g. HepaRG cells).

Output of this module is used by: Responses Dose response models Dose response data

Test systems from data

Test systems data formats

Test systems are the biological systems (e.g., animals) or in-vitro systems on which responses related to health effects can be measured.

Download empty dataset template: Zipped CSV Excel

Test Systems

Each test system should have a unique identification code (idSystem), and (optionally) a name and a description. The test system's type (TestSystemType) indicates the type whether the test system is an in-vivo test system (in which case it is a model for external exposure) or any of a range of other, in-vitro, options (cell-line, etc., which all will be interpreted as models for internal exposure). Additionally, if applicable, the organ (e.g., liver) of the test system and the route of exposure (RouteExposure) for in-vivo test systems (oral, dermal or inhalation) may be specified.

Name	Туре	Description	Aliases	Required
idTestSystem	AlphaNumeric (50)	Unique identification code of the test system.	idTestSystem, idSystem, Id, Code	Yes
CodeSystem	AlphaNumeric (100)	Identifier of the code system of the test systems.	CodeSystem	No
Name	AlphaNumeric (100)	Name of the test system.	Name	No
Description	AlphaNumeric (200)	Additional description or label of the test system.	Description	No
TestSystem- Type	TestSystemType	The type of the test system, i.e., in-vivo, cell-line, primary cells, tissue, organ. Controlled terminology.	TestSystem- Type, SystemType	No
Organ	AlphaNumeric (100)	If applicable, the organ that the cells originate from associated with the in vitro test-system.	Organ	No
Species	AlphaNumeric (100)	If applicable, the species associated with the test-system.	Species	No
Strain	AlphaNumeric (100)	If applicable, the strain of the species associated with the test-system.	Strain	No
RouteExposure	ExposureRoute	If applicable, the route of exposure associated with the in vivo test-system, oral, dermal, inhalation, s.c., i.v. Controlled terminology.	ExposureRoute- Type, ExposureRoute, RouteExposure	No
Guideline- Method	AlphaNumeric (200)	Reference to test guideline.	GuidelineStudy	No
Reference	AlphaNumeric (200)	External reference(s) to other sources containing more information about the test system. E.g., publications, website, documents.	Reference	No

Accepted table names: TestSystems, TestSystem, System.

Test systems as data

Test systems are provided as data.

- Test systems data formats
- Test systems from data

3.2 Consumption modules

Consumption modules specify the *consumptions* or *single value consumptions* of *foods* by surveyed individuals in *populations*. Foods can be related to each other using *food recipes*.

3.2.1 Consumptions

Consumptions data are the amounts of foods consumed on specific days by individuals in a food consumption survey. For acute exposure assessments, the interest is in a population of person-days, so one day per individual may be sufficient. For chronic exposure assessments, the interest is in a population of persons, so preferably two or more days per individual are needed.

This module has as primary entities: Populations Foods

Output of this module is used by: Food conversions Consumptions by modelled food

Consumptions from data

Consumptions calculation

The consumptions module offers a number of options and filters that specify how the consumption data should be included in the assessment.

- After including individual properties for defining a population in the *populations panel* an option becomes available *Match consumption data to population definition options*. Use all populations definitions (default), ignore all populations definitions or use a selection of properties to define the population.
- When individual sampling weights have been recorded in the *data*, the option *ignore sampling weights* specifies whether sampling weights are ignored or not in all calculations.
- It is possible to *restrict to consumptions of specific foods*.
- Depending on the *exposure type* (acute or chronic), the individuals (chronic) or individual-days (acute) can be filtered in several ways. It is possible to restrict the consumers or consumer days to consumers that consume or consumer days with a consumption. Use option *consumers or consumer days only*, and only consumers or consumer days with consumption of *specific (focal) products* are selected.
- For a chronic dietary or risk assessment (OIM, LNN or BBN), the individuals with less than N surveys days may be filtered out. Check setting *Exclude individuals with less than N days*, then./ option *N (number of days in survey)* appears. Specify a number, all individuals with a number of survey days less than N are filtered out.
- In general, one consumption data source is selected. However, the menu allows selection of multiple datasources through the use of the *add another data source* option. Press the add button + on the right side of the panel and select a datasource. Then, in the settings panel a new checkbox appears *Loop over multiple surveys* and after checking it all red triangles in the left panel disappear. See also *multiple surveys*.

In Figure 3.9 the consumption panel is shown. The setting *Match consumption data to population definition* is selected as default. All properties that are included in the *populations definition* are used. See *other options* for more info.

Consumption population definition

In Figure 3.10 all options of setting Match consumption data to population definition options are shown.

- Match consumption data to population definition: all selected properties in the population definition are used (default),
- Ignore population definition (use all individuals in survey): all selected properties in the population definition are ignored,
- Match consumption data to population definition using selected properties only: a selection of individual properties is used to define the population.

In the *populations panel*, the population was defined by including individual properties. These properties are all used (default), ignored or *a subset is selected* for further restricting the population.

In Figure 3.12 the selected individual properties that are available are shown. Check one or more for restricting the population definition to the specified set.

Consumptions	02e5f03d	▶ ₿
Consumptions		0
Consumptions data source		
> ConsumptionsProp_Age_Gender.xlsx	e) /
add another data source		+
Inputs		
Populations		~
Foods (5464 selected)		~
Consumptions data selection		
Food consumption surveys: selected 1 of 2 (clear filter) (change selection)		1
Consumption settings	B Save	Changes
Match consumption data to population definition options		
Match consumption data to population definition		_ U
Restrict population to consumers or consumer days only		6
Consumption subset: restrict to consumptions of specific foods		0
Ignore sampling weights		0

Figure 3.9: Consumption and consumption settings.

Consumption settings	Save Changes
Match consumption data to population definition ontions Match consumption data to population definition	D
Ignore population definition (use all individuals in survey)	Ð
Match consumption data to population definition using selected properties only	

Figure 3.10: Consumption settings panel: Match consumption data to population definition options.

Consumption settings	Save Changes
Match consumption data to population definition options Match consumption data to population definition using selected properties only	- 0
Select one or more individual(day) properties to filter the population	. 0

Figure 3.11: Match consumption data to population definition using selected properties only.

Toggle all/none	
Match individuals by gender	
Match individuals by region	
Match individuals by gender	_



Multiple surveys

In the Consumption panel, consumption data sources are selected. After selecting two or more data sources (*add another data source*), MCRA performs a loop over the selected surveys, see Figure 3.13. Check *Loop over multiple surveys*.

Risks / Consumptions 8b	o0f9cdd	•	¢
Consumptions data source			
✓ [2 data sources]			
> Fpfas_1-79y_20220323.accdb v1		1	
> Hpfas_1-79y_20220323.accdb v1		1	
> [4 merged tables]			urce
add another data source		+	
Inputs			
Populations		~	•
Foods (18648 selected)		~	•
Consumptions data selection			
Food consumption surveys: selected 2 of 2 (clear filter) (change selection)		-	
Consumption settings	B Save C	hanges	
✓ Loop over multiple surveys		0	
Restrict population to consumers or consumer days only		Ð	

Figure 3.13: Loop over multiple surveys.

The advantage of running a multiple survey loop instead of running separate actions for each surveyn is that outputs are directly compared. In the Results panel, Figure 3.14, a toc appears containing links to the output of the risk or dietary exposure assessment (main action) for the selected surveys, but also a link to the combined output panel.

After clicking the link 2022 surveys, the combined output is represented in violin plots and a table with percentiles for the specified percentages. Here a combined risks action is run and the p99 of the risk characterisation ratio (exposure/hazard) distribution is selected. For a dietary exposure action similar results are available.

Results						${\bf Q}_{\rm c}$ Type filter text here	:
] Output		Status	Message	Date	Running time	
> □	2022 surveys MOE	*	Ran to completion		08-03-2023 09:35	00:00:00	:
~ <	2022 surveys	*	Ran to completion		17-03-2023 15:29	00:00:01	:
	fFCS2016_Core	*	Ran to completion		17-03-2023 15:28	00:02:10	:
	hFCS2016_Core	*	Ran to completion		17-03-2023 15:29	00:01:18	:

Figure 3.14: Results panel multiple surveys.

Risks - Compari	son	Output info															
∽ Risks -	Compar	ison															:
p50.00	p90.00	p95.00	p97.50 j	o99.00 C	Combined overv	view hazard ind	ex										
Violin plots of uncertainty di	f the uncertai istribution. Th	nty distributio ne nominal rui	n of the haza is indicated	rd index at th by the black o	e p99.00 percer dot.	ntile of the popul	lation risk di	stribution	s. The vertica	al lines rep	present the me	dian and the	lower p2.5 ai	nd upper p	97.5 bound o	of the	
fFCS2016_Con	re					_											
hFCS2016_Cor	re																
		1		1.1 Hazaro	1.2 I index	1.3	1.	.4									
žl																	
Model	p50.00	p90.00	p95.00	p97.50	p99.00												
fFCS2016_Core	0.174 [0.171, 0.18]	0.478 [0.466, 0.495]	0.644 [0.624, 0.673]	0.812 [0.781, 0.846]	1.09 [1.03, 1.15]												
hFCS2016_Core	0.202 [0.195, 0.207]	0.541 [0.529, 0.554]	0.708 [0.681, 0.728]	0.916 [0.875, 0.992]	1.26 [1.22, 1.35]												

Figure 3.15: Risks comparison multiple surveys.

Results

Consumptions data formats

Consumption data is often collected in 24-hour dietary recall studies and contains the food consumptions and consumption amounts for a number of individuals on a number of days. For each of the individuals, the bodyweight should be specified, and optionally also age, sex, and other properties may be recorded. If applicable, sampling weights may also be specified that can be used to correct the sample of individuals in the survey to a more representative sample of the targeted population. The consumption amounts are usually expressed in grams, but may also be expressed in alternative units of plates, cups, or spoons. Optionally, the uncertainty of food consumption quantifications can be specified, see Souverein et al. (2011).

Consumption surveys are described using three tables: FoodSurveys, Individuals, and Consumptions. Individuals are linked to food surveys using the survey code (idFoodSurvey), and consumptions are linked to individuals using the individual codes (idIndividual). The food codes used to identify the consumed foods should match with the codes provided by the foods entity definitions.

Download empty dataset template: Zipped CSV Excel

Food consumption surveys

The records of the food consumption surveys table contain the ids, names, descriptions, and other relevant metadata of consumption surveys.

Name	Туре	Description	Aliases	Required
idSurvey	AlphaNumeric (50)	Unique identification code of the food consumption survey.	idSurvey, idFoodSurvey, Survey, FoodSurvey, SurveyId, FoodSurveyId, Code, Id	Yes
Name	AlphaNumeric (100)	The name of the food consumption survey.	Name, SurveyName	No
Description	AlphaNumeric (200)	Description of the food consumption survey.	Description	No
Location	AlphaNumeric (50)	The location or country where survey is held. It is recommended to use ISO Alpha-2 country codes.	Location, Country	No
BodyWeight- Unit	Body Weight Unit	The unit of bodyweight of the individuals of the survey: kg (default) or g.	BodyWeight- Unit, UnitBody- Weight, WeightIn	No
Consumption- Unit	<i>ConsumptionUnit</i>	The unit of the use/consumption amounts of the consumptions of the survey: g (default) or kg or CustomUnit (see table food consumption quantifications table).	AmountUnit, UnitAmount, AmountUnit, Consumption- Unit	No
StartDate	DateTime	The start date of the survey.	StartDate	No
EndDate	DateTime	The end date of the survey.	EndDate	No
NumberOf- SurveyDays	Integer	The number of days each individual participated in the survey.	NumberOf- SurveyDays, NDaysInSurvey	Yes
idPopulation	AlphaNumeric (50)	Unique identification code of the population.	IdPopulation, PopulationId	No

Table 3.27:	Table	definition	for Food	consumption surveys.
				1 2

Accepted table names: FoodConsumptionSurveys, ConsumptionSurveys, FoodSurveys, Surveys.

Individuals

The individuals of a survey are recorded in the Individuals table. Add additional properties like Region, Breastfeeding to further describe an individual. In table IndividualProperties, each property in the Individuals table is described (recommended way). Note that only those properties that are available in the Individuals table are used in module Populations, table Populations or PopulationIndividualPropertyValues to subset the individuals. This is only relevant when the UseData option in the population module is used.

Name	Туре	Description	Aliases	Required
idIndividual	AlphaNumeric (50)	Unique identification code of the individual.	idIndividual, IndividualId, Individual, Id	Yes
idFoodSurvey	AlphaNumeric (50)	The identification code / short name of survey.	idSurvey, idFoodSurvey, Survey, FoodSurvey, SurveyId, FoodSurveyId, SurveyCode	Yes
BodyWeight	Numeric	The body weight of the individual.	BodyWeight, Weight	No
Sampling- Weight	Numeric	The sampling weight for an individual (default = 1).	SamplingWeight	No
NumberOf- SurveyDays	Integer	The number of days the individual participated in the survey.	NumberOf- SurveyDays, NumberOfDays- InSurvey, DaysInSurvey, NDaysInSurvey	No
Name	AlphaNumeric (100)	Name or label of the individual.	Name	No
Description	AlphaNumeric (200)	Additional description of the individual.	Description	No
Individual properties		Other individual properties can be added like the fields Age, Gender, Region etc. These properties are automatically parsed as co-factors or co-variables.		No

Table 3.28: Table definition for Individuals.

Accepted table names: Individuals, SurveyIndividuals, ConsumptionSurveyIndividuals, FoodConsumptionSurveyIndividuals.

IndividualDays

The individuals and days in the survey are recorded in the individualDays table.

Name	Туре	Description	Aliases	Required
idIndividual	AlphaNumeric (50)	The identification code of the individual.	idIndividual, IndividualId, Individual, Id	Yes
idDay	AlphaNumeric (50)	Identification code of the day of consumption, sequential number	idDay, DayId, Day, DayOfSurvey	Yes
SamplingDate	DateTime	The date of the consumption.	Date, SamplingDate, SurveyDate	No

Table 3.29: Table definition for IndividualDays.

Accepted table names: IndividualDays, SurveyIndividualDays, ConsumptionSurveyIndividualDays, FoodConsumptionSurveyIndividualDays.

Individual properties

This table is used to describe the properties used in the Populations or PopulationIndividualPropertyValues table characterising the population (table Populations) and/or the properties used in the Individuals table characterising an individual. Properties like Age, Gender, Region are describing an individual (PropertyLevel = Individual). Properties like Period (for populations) or Month (sampling date for an individual day) are describing an individual day (PropertyLevel = IndividualDay).

Name	Туре	Description	Aliases	Required
idIndividual- Property	AlphaNumeric (50)	The code of the property.	idIndividual- Property, Individual- PropertyId, Individual- Property	Yes
Name	AlphaNumeric (100)	The name of the property.	Name	No
PropertyLevel	PropertyLevelType	The level of the property. This type follows a controlled terminology, with possible values: Individual or IndividualDay.	PropertyLevel, LevelProperty	No
Description	AlphaNumeric (200)	Description of the property.	Description	No
Туре	IndividualProperty- Type	This field specifies the type of the values of this individual property. This type follows a controlled terminology, with possible values: Boolean, Categorical (default), Numeric, Nonnegative, Integer, NonnegativeInteger, Month, Datetime, Gender.	Туре	No

Table 3.30:	Table defin	ition for	Individual	properties.
14010 5.50.	rable defin	intion for	marviauai	properties.

Accepted table names: IndividualProperties, IndividualProperty.

Individual property values

Not recommended. This table describes individual property values. Property values are describing an individual for properties like e.g. Region, Breastfeeding. The recommended way is to add these columns as additional columns in the Individuals table. In table IndividualProperties, each property in the IndividualPropertyValues table is described.

Name	Туре	Description	Aliases	Required
idIndividual	AlphaNumeric (50)	The identification number of the Individual.	Id	Yes
PropertyName	AlphaNumeric (50)	The name of the property.	Name	Yes
TextValue	AlphaNumeric (50)	The value of the property as text value.		No
DoubleValue	Numeric	The value of the property as number.		No

Table 3.31: Table definition for Individual property values.

Accepted table names: IndividualPropertyValues, IndividualPropertyValue.

Consumptions

The individual consumptions are recorded in the consumptions table.

Name	Туре	Description	Aliases	Required
idIndividual	AlphaNumeric (50)	The unique identification code of the consumer (individual).	idIndividual, IndividualId, Individual	Yes
idFood	AlphaNumeric (50)	The food code (food as eaten code).	idFood, Food, FoodId, FoodConsumed, FoodAsEaten	Yes
Facets	AlphaNumeric	The codes of the facets/treatments recorded for this consumption. Multiple treatments are separated by a '\$'.	Treatments, Treatment, Facets	No
idUnit	AlphaNumeric (50)	Identification code of the unit in which the food is consumed (e.g. plate, cup, spoon).	idUnit, Unit, UnitId	No
idDay	AlphaNumeric (25)	Identification code of the day of consumption, sequential number	idDay, DayId, Day, DayOfSurvey	Yes
idMeal	AlphaNumeric (25)	Identification code of the meal (eating occasion within a day).	idMeal, MealId, Meal	No
Amount	Numeric	The consumed portion of food in g (default) or kg or quantity of a plate, cup, spoon. Days without consumptions are not recorded.	Amount, Amount- Consumed	Yes

Table 3.32:	Table	definition	for	Consumptions.	
-------------	-------	------------	-----	---------------	--

Accepted table names: FoodConsumptions, FoodConsumption, Consumptions, Consumption.

Consumptions settings

Selection settings

Name	Туре	Description
Selected tier	SettingsTemplateType	Specifies all module settings should be set according to a pre-defined tier or using custom settings.
Exposure type	ExposureType	The type of exposure considered in the assessment; acute (shorterm) or chronic (long-term).
Food survey	AlphaNumeric	The food consumption survey representative for the population interest.
Restrict population to consumers or consumer days only	Boolean	Specifies whether the population should be restricted to the individuals (chronic) or individual days (acute) that have non-ze consumption.
Restrict population to consumers or consumer days with consumptions of specific foods	Boolean	Specifies whether the population should be restricted to the individuals (chronic) or individual days (acute) consuming any the foods of the specified subset.
Selected foods-as-eaten	AlphaNumeric	Set of consumed foods that are of particular interest for restrict the consumers / consumption days.
Consumption subset: restrict to consumptions of specific foods	Boolean	If checked, then the consumptions are restricted to those of the specified food-as-eaten subset.
Selected foods-as-eaten	AlphaNumeric	Set of consumed foods that are of particular interest.
Match consumer selection to population definition options	IndividualSubsetType	Match consumption data to population definition. Use population definitions (default), ignore all population definitions or use a selection of properties.
Select one or more individual(day) properties to filter the population	AlphaNumeric	Select one or more individual(day) properties to filter the individuals(days) in the population.
Ignore sampling weights	Boolean	If checked, individual sampling weights are not used (sampling weight = 1). If unchecked, the specified sampling weights are used.
Exclude individuals with less than N days	Boolean	Filter out all individuals with less than N survey days.
N (number of days in survey)	Numeric	Specify the nominal number of days in the survey to filter out a individuals with less than N survey days.
Cofactor name	AlphaNumeric	Specify the name of the cofactor.
Covariable name	AlphaNumeric	Specify the name of the covariable.
Individual day subset	IndividualDaySubsetDefinition	Individual day subset definition.
Individuals subset definitions	IndividualsSubsetDefinition	Contains a list of subset definitions to filter the population's individuals based on an individual's properties, by property nan and a custom filter query, for example a value range or a list of keywords.

Table 3.33: Selection settings for module Consumptions.

Calculation settings

Table 3.34: Calculation settings for module Consumptions.

Name	Туре	Description
Define populations based on specified individual properties	Boolean	Define a population by selecting specific ranges/values of individual properties. E.g., the female population between ages and 45 is composed of the properties gender (female) and age (between 18 and 45).

Output settings

		—
Name	Туре	Description
Lower percentage for variability (%)	Numeric	The default value of 25% may be overruled.
Upper percentage for variability (%)	Numeric	The default value of 75% may be overruled.

Table 3.35: Output settings for module Consumptions.

Uncertainty settings

	· •	•
Name	Туре	Description
Resample individuals	Boolean	Individual data are resampled from the original database using bootstrap methodology (Efron 1979, Efron & Tibshirani 1993)
Resample portion sizes	Boolean	Specifies whether portion sizes should be resampled based on food consumption quantification data, see (Souverein et al. 201

Table 3.36: Uncertainty settings for module Consumptions.

Consumptions tiers
Overview

N EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFSA 2022 Acute Tier II	EFSA 2022 Chronic Tier II	EFSA 2023 Acute Prospec- tive Tier II	EFS 2023 Chros Pros tive II
Ig true ne sa pl w	true	true	true	true	true	true	true	true	true
E false cl in di vi u al w le th N di	true	false	true	false	true	false	true	false	true
N (r b) o) d: ir su ve	2		2		2		2		2

Table 3.37: Tier overview for module Consumptions.

Retrospective dietary CRA (EC 2018) - Acute / Tier I

Table 3.38: Tier definition for Retrospective dietary CRA (EC 2018) - Acute / Tier I.

Name		Setting
	Ignore sampling weights	true
	Exclude individuals with less than N days	false

Retrospective dietary CRA (EC 2018) - Chronic / Tier I

Table 3.39: Tier definition for Retrospective dietary CRA (EC 2018) - Chronic / Tier I.

Name	Setting
Ignore sampling weights	true
Exclude individuals with less than N days	true
N (number of days in survey)	2

Retrospective dietary CRA (EC 2018) - Acute / Tier II

Table 3.40: Tier definition for Retrospective dietary CRA (EC 2018) - Acute / Tier II.

Name		Setting
	Ignore sampling weights	true
	Exclude individuals with less than N days	false

Retrospective dietary CRA (EC 2018) - Chronic / Tier II

Table 3.41: Tier definition for Retrospective dietary CRA (EC 2018) - Chronic / Tier II.

Name	Setting
Ignore sampling weights	true
Exclude individuals with less than N days	true
N (number of days in survey)	2

Retrospective dietary CRA (EFSA 2022) - Acute / Tier I

Table 3.42: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier I.

Name	Setting
Ignore sampling weights	true
Exclude individuals with less than N days	false

Retrospective dietary CRA (EFSA 2022) - Chronic / Tier I

Table 3.43: Tier definition for Retrospective dietary CRA (EFSA 2022) - Chronic / Tier I.

Name	Setting
Ignore sampling weights	true
Exclude individuals with less than N days	true
N (number of days in survey)	2

Retrospective dietary CRA (EFSA 2022) - Acute / Tier II

Table 3.44: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier II.

Name	Setting
Ignore sampling weights	true
Exclude individuals with less than N days	false

Retrospective dietary CRA (EFSA 2022) - Chronic / Tier II

Table 3.45: Tier definition for Retrospective dietary CRA (EFSA 2022) - Chronic / Tier II.

Name	Setting
Ignore sampling weights	true
Exclude individuals with less than N days	true
N (number of days in survey)	2

Prospective dietary CRA (EFSA 2023) - Acute / Tier II

Table 3.46: Tier definition for Prospective dietary CRA (EFSA 2023) - Acute / Tier II.

Name	Setting
Ignore sampling weights	true
Exclude individuals with less than N days	false

Prospective dietary CRA (EFSA 2023) - Chronic / Tier II

Table 3.47: Tier definition for Prospective dietary CRA (EFSA 2023) - Chronic / Tier II.

Name	Setting
Ignore sampling weights	true
Exclude individuals with less than N days	true
N (number of days in survey)	2

Prospective dietary CRA (EFSA 2024) - Acute / Tier II

Table 3.48: Tier definition for Prospective dietary CRA (EFSA 2024) - Acute / Tier II.

Name	Setting
Ignore sampling weights	true
Exclude individuals with less than N days	false

Prospective dietary CRA (EFSA 2024) - Chronic / Tier II

Table 3.49: Tier definition for Prospective dietary CRA (EFSA 2024) - Chronic / Tier II.

Name	Setting
Ignore sampling weights	true
Exclude individuals with less than N days	true
N (number of days in survey)	2

Consumptions uncertainty

In MCRA, in an *acute exposure* assessments, individual consumption day data are *resampled*, thus preserving the multivariate consumption patterns and associated weights and/or other individual characteristics. In MCRA we resample the set of individuals x number of survey days. We think that this implementation better reflects the notion of acute exposure which is expressed as the normalized intake per day. For *chronic exposure* assessments the resampling algorithm remained unchanged and the set of individuals (with corresponding days) is *resampled*.

Consumptions as data

Consumptions data are the amounts of foods consumed on specific days by individuals in a food consumption survey.

- Consumptions data formats
- Consumptions from data
- Consumptions calculation

Settings used

• Calculation Settings

3.2.2 Dust exposure determinants

Dust exposure determinants specify the factors that determine the exposure of individuals to dust.

This module has as primary entities: Substances

Output of this module is used by: Dust exposures

Dust exposure determinants from data

Dust exposure determinants data formats

Distributions for exposure determinants of substances in dust.

Download empty dataset template: Zipped CSV Excel

Dust ingestions

Variability distributions specifying the intake/ingestions of dust, specified using a distribution type, a mean and a coefficient of variation (CV). Distributions may be specified for different subgroups of age and sex.

Name	Туре	Description	Aliases	Required
idSubgroup	AlphaNumeric (50)	The id of the subgroup.	idSubgroup	Yes
AgeLower	Numeric (50)	Specifies the lower bound of the age interval (in years) of the dust exposure determinant subgroup. Individuals belong to a subgroup when the age of the individual is equal or greater than the specified lower bound and smaller than the specified lower bound of the next subgroup.	AgeLower, LowerAge	No
Sex	GenderType	The sex of the subgroup.	Sex, Gender	No
Value	Numeric	Distribution mean.	Mean	Yes
ExposureUnit	ExternalExposure- Unit	The unit of the dust ingestion. If not specified, then a default unit of g/day is assumed.	Unit, Dust- IngestionUnit	No
Distribution- Type	DustIngestion- DistributionType	Distribution type (Uniform, or LogNormal). If not specified, the dust adherence amount is assumed to be a constant factor.	Distribution- Type, Distribution	No
CvVariability	Numeric	Coefficient of variation of the distribution.		No

Accepted table names: DustIngestions, DustIngestion.

Dust body exposure fractions

Variability distributions specifying the individual fractions of the body exposed to dust. The distributions are specified using a distribution type, a mean and a coefficient of variation (CV). Distributions may be specified for different subgroups of age and sex.

Name	Туре	Description	Aliases	Required
idSubgroup	AlphaNumeric (50)	The id of the subgroup.	idSubgroup	Yes
AgeLower	Numeric (50)	Specifies the lower bound of the age interval (in years) of the dust body exposure fractions subgroup. Individuals belong to a subgroup when the age of the individual is equal or greater than the specified lower bound and smaller than the specified lower bound of the next subgroup.	AgeLower, LowerAge	No
Sex	GenderType	The sex of the subgroup.	Sex, Gender	No
Value	Numeric	Mean.	Value, Mean, Mean	Yes
Distribution- Type	DustBodyExposure- FractionDistribution- Type	Distribution type (Uniform, or LogNormal). If not specified, the dust body exposure fraction is assumed to be constant.	Distribution- Type, Distribution	No
CvVariability	Numeric	Coefficient of variation.	CvVariability, Sd	No

Table 3.51: Table definition for Dust body exposure fractions

Accepted table names: DustBodyExposureFractions, DustBodyExposureFraction.

Dust adherence amounts

Variability distributions specifying the dust adherence amounts. The distributions are specified using a distribution type, a mean and a coefficient of variation (CV). Distributions may be specified for different subgroups of age and sex.

Name	Туре	Description	Aliases	Required
idSubgroup	AlphaNumeric (50)	The id of the subgroup.	idSubgroup	Yes
AgeLower	Numeric (50)	Specifies the lower bound of the age interval (in years) of the dust adherence fractions subgroup. Individuals belong to a subgroup when the age of the individual is equal or greater than the specified lower bound and smaller than the specified lower bound of the next subgroup.	AgeLower, LowerAge	No
Sex	GenderType	The sex of the subgroup.	Sex, Gender	No
Distribution- Type	DustAdherence- AmountDistribution- Type	Distribution type (Uniform, or LogNormal). If not specified, the dust adherence amount is assumed to be a constant factor.	Distribution- Type, Distribution	No
Value	Numeric	Mean adherance amount (in g/m2).	Value, Mean	Yes
CvVariability	Numeric	Coefficient of variation.	CvVariability, CV	No

Table 3.52: Table definition	for Dust adherence amounts.
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 $\label{eq:Accepted table names: DustAdherenceAmounts, DustAdherenceAmount.$

Dust availability fractions

Distributions for fractions of substance in dust available for dermal contact can be dependent on e.g. age.

Name	Туре	Description	Aliases	Required
idSubgroup	AlphaNumeric (50)	The id of the subgroup.	idSubgroup	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
AgeLower	Numeric (50)	Specifies the lower bound of the age interval (in years) of the dust exposure determinant subgroup. Individuals belong to a subgroup when the age of the individual is equal or greater than the specified lower bound and smaller than the specified lower bound of the next subgroup.	AgeLower, LowerAge	No
Sex	GenderType	The sex of the subgroup.	Sex, Gender	No
Value	Numeric	Mean of the distribution.	Mean	Yes
Distribution-	DustAvailability-	Distribution type (Uniform, or	Distribution-	No
Туре	FractionDistribution-	LogNormal). If not specified,	Туре,	
	Туре	the dust availability fraction is assumed to be constant.	Distribution	
CvVariability	Numeric	Coefficient of variation.		No

Table 3.53:	Table	definition	for Dust	availability	fractions.
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Accepted table names: DustAvailabilityFractions, DustAvailabilityFraction.

Dust exposure determinants as data

Dust exposure determinant data can be entered using the internal, relational data format.

- Dust exposure determinants data formats
- Dust exposure determinants from data

3.2.3 Soil exposure determinants

Soil exposure determinants specify the factors that determine the exposure of individuals to soil.

This module has as primary entities: *Substances*

Output of this module is used by: Soil exposures

Soil exposure determinants from data

Soil exposure determinants data formats

Distributions for exposure determinants of substances in soil. Download empty dataset template: Zipped CSV Excel

Soil ingestions

Distributions for soil ingestions are dependent on e.g. age.

Name	Туре	Description	Aliases	Required
idSubgroup	AlphaNumeric (50)	The id of the subgroup.	idSubgroup	Yes
AgeLower	Numeric (50)	Specifies the lower bound of the age interval (in years) of the soil exposure determinant subgroup. Individuals belong to a subgroup when the age of the individual is equal or greater than the specified lower bound and smaller than the specified lower bound of the next subgroup.	AgeLower, LowerAge	No
Sex	GenderType	The sex of the subgroup.	Sex, Gender	No
Value	Numeric	Mean of the distribution.	Mean	Yes
ExposureUnit	ExternalExposure- Unit	The unit of the soil ingestion. If not specified, then a default unit of g/day is assumed.	Unit, Soil- IngestionUnit	No
Distribution- Type	SoilIngestion- DistributionType	Distribution type (Uniform, or LogNormal). If not specified, the soil adherence amount is assumed to be a constant factor.	Distribution- Type, Distribution	No
CvVariability	Numeric	Coefficient of variation.		No

Table 3.54: Table definition for Soil ingestions.	•
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Accepted table names: SoilIngestions, SoilIngestion.

Soil exposure determinants as data

Soil exposure determinant data can be entered using the internal, relational data format.

- Soil exposure determinants data formats
- Soil exposure determinants from data

3.2.4 Air exposure determinants

Air exposure determinant data are distributions of chemical substances occurring in air samples. The distributions can differ per age group.

This module has as primary entities: Substances

Output of this module is used by: Air exposures

Air exposure determinants from data

Air exposure determinants data formats

Distributions for exposure determinants of substances in air.

Download empty dataset template: Zipped CSV Excel

AirIndoorFractions

Point estimates for indoor fractions,,can be dependent on e.g. age.

Name	Туре	Description	Aliases	Required
idSubgroup AgeLower	AlphaNumeric (50) Numeric (50)	The id of the subgroup. Specifies the lower bound of the age interval (in years) of the air exposure determinant subgroup. Individuals belong to a subgroup when the age of the individual is equal or greater than the specified lower bound and smaller than the specified lower bound of the next subgroup.	idSubgroup AgeLower, LowerAge	Yes No
Fraction	Numeric	Fraction indoors, between 0 and 1.	Fraction, FractionIndoors	Yes

Table 3.55:	Table definition	for AirIndoorFractions.
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Accepted table names: AirIndoorFractions, AirIndoorFraction, FractionsAirIndoor.

Air ventilatory flow rates

Distributions for air ventilatory flow rates (default m3/day) can be dependent on e.g. age.

Name	Туре	Description	Aliases	Required
idSubgroup	AlphaNumeric (50)	The id of the subgroup.	idSubgroup	Yes
AgeLower	Numeric (50)	Specifies the lower bound of the age interval (in years) of air ventilatory flow rate subgroup. Individuals belong to a subgroup when the age of the individual is equal or greater than the specified lower bound and smaller than the specified lower bound of the next subgroup.	AgeLower, LowerAge	No
Sex	GenderType	The sex of the subgroup.	Sex, Gender	No
Value	Numeric	Mean of the distribution.	Value, Mean	Yes
Distribution-	VentilatoryFlow-	Distribution type	Distribution-	No
Туре	RateDistributionType	(LogNormal). If not specified, the air ventilatory flow rate is assumed to be constant.	Type, Distribution	
CvVariability	Numeric	Coefficient of variation.	CvVariability, Sd	No

Table 3 56.	Table definition	for Air ventilatory	flow rates
1 auto 5.50.	rable demittion	IOI All volutilatory	now rates.

Accepted table names: AirVentilatoryFlowRates, AirVentilatoryFlowRate.

Air exposure determinants as data

Air exposure determinant data can be entered using the internal, relational data format.

- Air exposure determinants data formats
- Air exposure determinants from data

3.2.5 Consumer product exposure determinants

Consumer product exposure determinant data are distributions of chemical substances occurring in consumer products. The distributions can differ per age group.

This module has as primary entities: Substances Consumer products

Output of this module is used by: Consumer product exposures

Consumer product exposure determinants from data Consumer product exposure determinants data formats

Distributions for exposure determinants of substances in consumer products.

Download empty dataset template: Zipped CSV Excel

Consumer product exposure fractions

Fractions of the substance used with the product to which a consumer is exposed.

Name	Туре	Description	Aliases	Required
idProduct	AlphaNumeric (50)	The consumer product.	idProduct, ProductId	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
ExposureRoute	ExposureRoute	The exposure route(s) (only applicable if exposure level is external) for which this exposure.	ExposureRoute, Route	Yes
Exposure- Fraction	Numeric	Fraction between 0 and 1.	Exposure- Fraction, Fraction	Yes

Γ_{0} $h_{10} 2 57$	Tabla	definition	for	Comment	meno durat	011000011000	fractiona
1 able 5.5/1	rable	demnition	TOF	Consumer	Droduct	exposure	fractions.

Accepted table names: ConsumerProductExposureFractions, CPExposureFractions, ExposureFractions.

Consumer product application amounts

Distributions of consumer product application amounts represent the quantities used or applied during a single use. These distributions can be characterized for the general population or for specific subgroups based on age and/or sex.

Name	Туре	Description	Aliases	Required
idProduct	AlphaNumeric (50)	Identifier/code of the consumer product.	idProduct, ProductId	Yes
AgeLower	Numeric (50)	Specifies the lower bound of the age interval (in years) of the application amount in case the distribution applies to a specific population age subgroup. Individuals belong to a subgroup when the age of the individual is equal or greater than the specified lower bound and smaller than the specified lower age of the next subgroup.	AgeLower, LowerAge	No
Sex	GenderType	Specifies the sex associated with this application amount in case the distribution applies to a specific population sex subgroup.	Sex, Gender	No
Amount	Numeric	Mean of the distribution. Amounts are in g (gram, default unit).	Amount, Mean	Yes
Distribution- Type	ApplicationAmount- DistributionType	Type of the distribution. If not specified, the application amount is assumed to be constant.	Distribution- Type, Distribution	No
CvVariability	Numeric	Coefficient of variation of the distribution.	Cv, CvVariability	No
Unit	ApplicationAmount- Unit	The unit for the consumer product application amount. When not specified, then a default unit of g (gram) is assumed.	Unit, Application- AmountUnit	No

Table 3.5	8. Table	definition	for	Consumer	product	application	amounts
14010 5.5	0. 1 auto	ucilition	101	Consumer	produce	application	amounts.

Accepted table names: ConsumerProductApplicationAmounts, CPApplicationAmounts, ApplicationAmounts.

Consumer product exposure determinants as data

Consumer product exposure determinant data can be entered using the internal, relational data format.

- Consumer product exposure determinants data formats
- Consumer product exposure determinants from data

3.2.6 Consumer product use frequencies

Consumer product use frequencies specify the frequency of use for a consumer product (personal care, do-it-yourself, cleaning product) per day.

This module has as primary entities: Populations Consumer products

Output of this module is used by: Consumer product exposures

Consumer product use frequencies from data

Consumer product use frequencies data formats

Consumer product use frequencies are described by five table.

Download empty dataset template: Zipped CSV Excel

Consumer product surveys

The records of the consumer product surveys table contain the ids, names, descriptions, and other relevant metadata of consumer product surveys.

Name	Туре	Description	Aliases	Required
idSurvey	AlphaNumeric (50)	Unique identification code of the consumer product survey.	idSurvey, Survey, SurveyId, Id	Yes
Name	AlphaNumeric (100)	The name of the consumer product survey.	Name, SurveyName	No
Description	AlphaNumeric (200)	Description of the consumer product survey.	Description	No
Country	AlphaNumeric (50)	The location or country where the survey is held. It is recommended to use ISO Alpha-2 country codes.	Location, Country	No
BodyWeight- Unit	Body Weight Unit	The unit of bodyweight of the individuals of the survey: kg (default) or g.	BodyWeight- Unit, UnitBody- Weight, WeightIn	No
AmountUnit	<i>ConsumptionUnit</i>	The unit of the amounts of consumer product use in the survey: g (default) or kg or CustomUnit (see table food consumption quantifications table).	AmountUnit, UnitAmount	No
StartDate	DateTime	The start date of the survey.	StartDate	No
EndDate	DateTime	The end date of the survey.	EndDate	No
idPopulation	AlphaNumeric (50)	Unique identification code of the population.	IdPopulation, PopulationId	No

Table 3.59: Table definition for Consumer product surveys.

Accepted table names: ConsumerProductSurveys, CPSurveys.

Individuals

The individuals of a survey are recorded in the Individuals table. Add additional properties like Region, Breastfeeding to further describe an individual. In table IndividualProperties, each property in the Individuals table is described (recommended way). Note that only those properties that are available in the Individuals table are used in module Populations, table Populations or PopulationIndividualPropertyValues to subset the individuals. This is only relevant when the UseData option in the population module is used.

Name	Туре	Description	Aliases	Required
idIndividual	AlphaNumeric (50)	Unique identification code of the individual.	idIndividual, IndividualId, Individual, Id	Yes
idFoodSurvey	AlphaNumeric (50)	The identification code / short name of survey.	idSurvey, idFoodSurvey, Survey, FoodSurvey, SurveyId, FoodSurveyId, SurveyCode	Yes
BodyWeight	Numeric	The body weight of the individual.	BodyWeight, Weight	No
Sampling- Weight	Numeric	The sampling weight for an individual (default = 1).	SamplingWeight	No
NumberOf- SurveyDays	Integer	The number of days the individual participated in the survey.	NumberOf- SurveyDays, NumberOfDays- InSurvey, DaysInSurvey, NDaysInSurvey	No
Name	AlphaNumeric (100)	Name or label of the individual.	Name	No
Description	AlphaNumeric (200)	Additional description of the individual.	Description	No
Individual properties		Other individual properties can be added like the fields Age, Gender, Region etc. These properties are automatically parsed as co-factors or co-variables.		No

Table 3.60: Table definition for Individuals.

Accepted table names: Individuals, SurveyIndividuals, ConsumptionSurveyIndividuals, FoodConsumptionSurveyIndividuals.

Individual properties

This table is used to describe the properties used in the Populations or PopulationIndividualPropertyValues table characterising the population (table Populations) and/or the properties used in the Individuals table characterising an individual. Properties like Age, Gender, Region are describing an individual (PropertyLevel = Individual). Properties like Period (for populations) or Month (sampling date for an individual day) are describing an individual day (PropertyLevel = IndividualDay).

Name	Туре	Description	Aliases	Required
idIndividual- Property	AlphaNumeric (50)	The code of the property.	idIndividual- Property, Individual- PropertyId, Individual- Property	Yes
Name	AlphaNumeric (100)	The name of the property.	Name	No
PropertyLevel	<i>PropertyLevelType</i>	The level of the property. This type follows a controlled terminology, with possible values: Individual or IndividualDay.	PropertyLevel, LevelProperty	No
Description	AlphaNumeric (200)	Description of the property.	Description	No
Туре	IndividualProperty- Type	This field specifies the type of the values of this individual property. This type follows a controlled terminology, with possible values: Boolean, Categorical (default), Numeric, Nonnegative, Integer, NonnegativeInteger, Month, Datetime, Gender.	Туре	No

Accepted table names: IndividualProperties, IndividualProperty.

Individual property values

Not recommended. This table describes individual property values. Property values are describing an individual for properties like e.g. Region, Breastfeeding. The recommended way is to add these columns as additional columns in the Individuals table. In table IndividualProperties, each property in the IndividualPropertyValues table is described.

Name	Туре	Description	Aliases	Required
idIndividual	AlphaNumeric (50)	The identification number of the Individual.	Id	Yes
PropertyName	AlphaNumeric (50)	The name of the property.	Name	Yes
TextValue	AlphaNumeric (50)	The value of the property as text value.		No
DoubleValue	Numeric	The value of the property as number.		No

Table 3.62: Table definition for Individual property values.

Accepted table names: IndividualPropertyValues, IndividualPropertyValue.

Individual consumer product use frequencies

Individual consumer product use frequencies are specified as a number of use per day for a consumer product (>= 0).

Name	Туре	Description	Aliases	Required
idIndividual	AlphaNumeric (50)	The unique identification code of the consumer (individual).	idIndividual, IndividualId, Individual	Yes
idProduct	AlphaNumeric (50)	The consumer product id or code.	idProduct, ProductId, Product	Yes
Frequency	Numeric	The frequency of use, a value equal or higher than zero (freq $>= 0$).	Frequency	Yes

Table 3.63: Table definition for Individual consumer product use frequencies.

Accepted table names: ConsumerProductUseFrequencies, CPUseFrequencies, UseFrequencies.

Consumer product use frequencies as data

Consumer product use frequencies specify the frequency of use.

- Consumer product use frequencies data formats
- Consumer product use frequencies from data

3.2.7 Food recipes

Food recipes data specify the composition of specific foods (typically: foods-as-eaten) in terms of other foods (intermediate foods or modelled foods) by specifying proportions in the form of a percentage.

This module has as primary entities: Foods

Output of this module is used by: Food conversions

Food recipes from data

Food recipes data formats

Recipe data to specify the ingredients of foods. Food recipes can be used to describe the ingredients of a composite food (e.g., of apple pie), or to specify the amount of a primary ingredient needed to obtain 100g of the food (e.g., grapes to raisins). Recipe is commonly used recursively (e.g., apple pie contains apple and flour, flour contains wheat).

Download empty dataset template: Zipped CSV Excel

Recipes

Name	Туре	Description	Aliases	Required
idFromFood	AlphaNumeric (50)	The code of the composite food (from-code), i.e., the code of the food for which the ingredient(s) are specified.	idFromFood, FromFoodId, FromFood, FoodFrom, Food	Yes
idToFood	AlphaNumeric (50)	The code of the ingredient food (to-code).	idToFood, ToFoodId, ToFood, FoodTo, Ingredient	Yes
Proportion	Numeric	Proportion of each ingredient in the food (%).	Proportion, Proportion%	Yes
idPopulation	AlphaNumeric (50)	Unique identification code of the population.	IdPopulation, PopulationId	No

Table 3.64: Table definition for Recipes.

Accepted table names: FoodTranslations, FoodTranslation, FoodCompositions, FoodComposition.

Food recipes as data

Food recipes are provided as data in the form of simple composition tables.

- Food recipes data formats
- Food recipes from data

3.2.8 Market shares

Market shares data specify for a given food, percentages of more specific foods (subfoods, e.g. brands) representing their share in a market. Market shares are used when consumption data are available at a more generalised level than concentration data.

This module has as primary entities: Foods

Output of this module is used by: Food conversions

Market shares from data

Market shares data formats

Describes the shares (proportions) in a market.

Download empty dataset template: Zipped CSV Excel

Market shares

Market shares main table.

Name	Туре	Description	Aliases	Required
idFood	AlphaNumeric (50)	The subtype of the food.	idFood, FoodId, Food, FoodType	Yes
Percentage	Numeric	Market share percentage of the subtype.	Percentage, Marketshare- Percentage, MarketShare, MarketShare- Percentage, MarketShare%	Yes
BrandLoyalty	Numeric	A parameter used in brand loyalty modelling, where 0 (default) specifies no brand loyalty (on each eating occasion a random selection of the next lower level in the hierarchy of food codes), and 1 specifies absolute brand loyalty (on subsequent eating occasions the same selection of the next lower level in the hierarchy of food codes).	BrandLoyalty	No

Accepted table names: MarketShares, MarketShare, FoodMarketShares, FoodMarketShare.

Market shares as data

Market shares are provided as data in the form of percentages.

- Market shares data formats
- Market shares from data

Market shares and brand loyalty

Sometimes measurements of substances in food are available at a more detailed food coding level than consumption data. For example, measurements may have been made for specific brands of a food whereas the consumption survey did not record the brand. MCRA allows to specify market share data for subtypes of a food (e.g. A\$1, A\$2, A\$3 are three brands of food A), and to calculate acute exposure based on such market shares.

For chronic assessments **brand loyalty** should be specified according to a simple Dirichlet model (Goodhardt et al. (1984)). Technically, the Dirichlet model for brand choice needs nbrand parameters α_i (which should be positive real numbers). The average brand choice probability for each brand is

 α_i/S

where

$$S = \sum \alpha_i$$

By definition, the market shares m_i should be proportional to the brand choice probabilities, and thus to the parameters α_i . Thus means that S, the sum of the alphas, is the only additional parameter that should be specified, and indeed this is the parameter that determines brand loyalty. S = 0 corresponds to absolute brand loyalty, and brand loyalty decreases with increasing S. We define $L = (1 + S)^{-1}$ as an interpretable brand loyalty parameter, where now L = 0 and L = 1 correspond to the situations of no brand loyalty and absolute brand loyalty, respectively.

The multinomial distribution models the probability of counts and is a generalization of the binomial distribution. The Dirichlet does the opposite, it models for a number of counts the distribution of probabilities, so for numbers $\alpha_1 = x_1, ..., \alpha_k = x_k$ the distribution of $m_1, ..., m_k$ is modelled with m_k the marketshare for brand k.

Given empirical or parametric distributions of consumption and concentration values, the algorithm for chronic exposure assessment now operates as follows:

- 1. Simulate consumptions for n individual(day)s.
- 2. Simulate n selection probabilities from the Dirichlet distribution.
- 3. For each individual, simulate d brand choices from a multinomial distribution using the individual specific selection probabilities from step 2.
- 4. For all individuals and days simulate values from the appropriate concentration distribution.
- 5. Multiply consumption with concentration to obtain exposure.

3.2.9 Single value consumptions

Single value consumption data are the single value amounts (Large Portion, Mean Consumption, p97.5Consumption) of modelled foods (foods-as-measured) consumed in a population.

This module has as primary entities: Populations Foods

Output of this module is used by: Single value dietary exposures

Single value consumptions from data

Single value consumptions calculation

Single value consumptions can be supplied *as data* or computed. When single value consumptions are computed from *consumptions by modelled food*, then the mean, median and large portion (p97.5 percentile) are computed for all modelled food consumption distributions. Besides these statistics, also the mean bodyweight of the population is computed. The following options are relevant in this calculation:

- Set the *risk type* option to *acute* if the consumptions should be based on the individual-day distributions. Otherwise, choose *chronic* to base them on the distributions aggregated by individual.
- Checking the *apply processing factors* option will compute the single value consumptions for the processed foods. When using this option, the output will also show a reverse yield factor, that is the ratio of the quantity of the raw commodity required to to obtain the processed commodity. Note, when no processing factors are available, the single-value consumption amounts of processed foods are expressed in terms of the processed commodities. The yield factor, i.e. the factor for translating the processed amount to the unprocessed amount, is not applied. In the IESTI calculations (and also chronic single-value calculations), calculations are done using the processed amounts, this is on the level of raw processed foods.
- Check the *restrict population to consumers or consumer days only (modelled-food)* option to compute the single value consumption statistics for each food based on the food consumers only. Note that checking this option will also affect the computed bodyweight, which is then computed by food based on the food-consumers only and can be different for each food.
- There is also an option to *ignore sampling weights* in the calculation.
- Check the *standardise consumption with body weight before calculation of single values or afterwards (with mean bodyweight)* option to compute the single value consumptions from the per bodyweight distribution. If unchecked, the per-person distribution will be used for computing the statistics. Note that although the results are reported per-day, the statistics are established by multiplying the statistics obtained from the per bw distribution by the bodyweight.

Single value consumptions data formats

Single value consumptions data provides a single per-individual-day and per-food consumption amount for a population. Also the bodyweight should be specified, and optionally also age, sex, and other properties may be recorded. The consumption amounts are usually expressed in grams, but may also be expressed in alternative units of plates, cups, or spoons. Optionally, the uncertainty of food consumption quantifications can be specified, see Souverein et al. (2011).

Single value consumptions are described using one table: PopulationConsumptionSingleValues.

Download empty dataset template: Zipped CSV Excel

Population consumption single values

Population consumption single values describe population food consumptions in the form of single value statistics.

Name	Туре	Description	Aliases	Required
idPopulation	AlphaNumeric (50)	Unique identification code of the population.	IdPopulation, PopulationId	Yes
idFood	AlphaNumeric (50)	The unique identification code of the consumed food.	idFood, FoodCode, Food	Yes
Value type of the single value consumption amount.	Consumption Value- Type	The value type of this consumption value.	Consumption- Type, ValueType, Consumption- ValueType, Consumption- SingleValue- Type	Yes
Percentile	Numeric	The percentile (if consumption value type is a percentile).	Percentile	No
Consumption- Amount	Numeric The consumed amount.		Amount, Consumption, Consumption- Amount, Amount- Consumed	Yes
Consumption- Unit	ConsumptionIntake- Unit	The unit of the consumption amount.	AmountUnit, UnitAmount, Consumption- Unit	No
Reference	AlphaNumeric (200)	Reference to the source from which this value is obtained.	Reference, References, Source, Sources	No

Table 3.66: Table definition for Population consumption single values.

Accepted table names: ConsumptionSingleValues, SingleValueConsumptions, PopulationConsumptionSingleValues, PopulationConsumptionValues.

Single value consumptions settings

Calculation settings

Name	Туре	Description
Exposure type	ExposureType	The type of exposure considered in the assessment; acute (shor term) or chronic (long-term).
Restrict population to consumers or consumer days only	Boolean	Specifies whether the population should be restricted to the individuals (chronic) or individual days (acute) that have non-ze consumption.
Ignore sampling weights	Boolean	If checked, individual sampling weights are not used (sampling weight = 1). If unchecked, the specified sampling weights are used.
Use standardised consumption distributions before calculation of single values	Boolean	Specifies whether single values are calculated on individual consumptions standardised with body weight and then multiplie by the mean body weight. Otherwise, single values are calculate on the original consumptions (per day). Note that both method lead to different estimates for the single value.
Apply processing factors	Boolean	Specified in table ProcessingFactor. If checked, processing fact are applied. Concentrations in the consumed food may be different from concentrations in the modelled food in monitorir programs (typically raw food) due to processing, such as peelin washing, cooking etc. If unchecked, no processing information used. This is in most (though not all) cases a worst-case assumption
Restrict population to consumers or consumer days only (food-as-measured)	Boolean	Specifies whether the population should be restricted to the individuals (chronic) or individual days (acute) with consumption containing any of the modelled foods.

Table 3.67: Calculation settings for module Single value consumptions.

Single value consumptions as data

Single value consumption data are the single value amounts of modelled foods (foods-as-measured) consumed in a population.

- Single value consumptions data formats
- Single value consumptions from data

Settings used

• Calculation Settings

Calculation of single value consumptions

Single value consumptions are calculated as a percentile (p97.5 or p99) or mean of the modelled food consumption distribution. For an acute single value dietary exposure assessment, this is the individual day consumption distribution, for chronic single value dietary exposure assessment, the individual consumption distribution is used.

• Single value consumptions calculation

Inputs used: Consumptions by modelled food

Settings used

• Calculation Settings

3.3 Occurrence modules

The basic occurrence data are *concentrations* for *substances* in *foods*, sometimes specified separately for a focal food as *focal food concentrations*. In some cases *concentration limits* are used as a stand-in when data are missing.

Concentration data are recalculated (if needed) as *active substance concentrations* in *modelled-foods*. If substance concentrations are not specified directly for the *active substances*, then they are converted using *substance conversions* and/or specified authorised *occurrence patterns*. The composition of mixed samples in total diet studies is described in *total diet study sample compositions*. *Food extrapolation rules* specify if insufficient data for a food can be suppleted with data from another food. From these basic data the list of *modelled-foods* is derived.

Active substance concentrations in modelled-foods are modelled in concentration models, optionally allowing for occurrence pattern models. In addition, processing factors and unit variability factors can be provided for further use in dietary exposure assessment.

3.3.1 Concentration distributions

Concentration distributions describe substance concentrations on foods in the form of summary statistics.

This module has as primary entities: Foods Substances

Output of this module is used by: Concentration models Dietary exposures

Concentration distributions from data

Concentration distributions data formats

Concentration distributions describe substance concentrations on foods in the form of summary statistics. These distributions can be characterised by a mean and a dispersion factor, the standard deviation or, preferably, a percentile point e.g. p95.

Download empty dataset template: Zipped CSV Excel

Concentration distributions

Substance concentrations on foods specified in the form of summary statistics.

Name	Туре	Description	Aliases	Required
idFood	AlphaNumeric (50)	Food code, the raw agricultural commodity.	idFood	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
Mean	Numeric	The mean of (monitoring) samples, on the original scale (in mg/kg).	Mean	Yes
CV	Numeric	Coefficient of variation, for samples of the size of the TDS pooled amount.	CV	No
Percentile	Numeric	The percentile at the point specified by the percentage.	Percentile	No
Percentage	Numeric	The percentage that belongs to the given the percentile, e.g., 95 (in mg/kg).	Percentage	No
Limit	Numeric	The specified norm value or limit value (in mg/kg).	Limit	No
Concentration- Unit	ConcentrationUnit	The unit of the limit value (default mg/kg).	Concentration- Unit, Unit	No

Table 5.00. Table definition for Concentration distributions.	Table 1	3.68:	Table	definition	for	Concentration	distributions.
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Accepted table names: ConcentrationDistributions, ConcentrationDistribution.

Concentration distributions as data

Concentration distributions describe substance concentrations on foods in the form of summary statistics.

- Concentration distributions data formats
- Concentration distributions from data

3.3.2 Concentration limits

Concentration limits specify (legal) limit values for substance concentrations on foods and are sometimes used as conservative values for concentration data. In the framework of pesticides the legal Maximum Residue Limit (MRL) is the best known example.

This module has as primary entities: Foods Substances

Output of this module is used by: Concentrations Single value concentrations Concentration models Modelled foods

Concentration limits from data

Concentration limits data formats

The concentration limits table describes limit values (e.g., MRLs) for specific food/substance combinations. This data may be used, for instance, for the food/substance combinations for which no concentration data is available. The food codes (idFood) and substance codes (idSubstance) should match the codes of the foods and substances table respectively.

Concentration limits are concentration limit values for specific food and substance combinations originating from regulations (e.g., MRLs). This data may be used, for instance, for the food/substance combinations for which no concentration data is available.

Download empty dataset template: Zipped CSV Excel

Concentration limits

The food codes (idFood) and substance codes (idSubstance) should match the codes of the foods and substances table respectively.

Name	Туре	Description	Aliases	Required
idFood	AlphaNumeric (50)	Code of the food of this residue limit definition.	idFood, FoodId, Food	Yes
idSubstance	AlphaNumeric (50)	Code of the substance of this residue limit definition.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
Value	Numeric	Residue limit value.	Value, Limit, Maximum- ResidueLimit, Maximum- ResidueLimits, MRL	Yes
StartDate	DateTime	Start date of the period during which the limit applies.	StartDate	No
EndDate	DateTime	End date of the period during which the limit applies.	EndDate	No
Concentration- Unit	ConcentrationUnit	<i>tionUnit</i> The unit of the limit value (default mg/kg).		No
ValueType	ConcentrationLimit- ValueType	Value type of the concentration value.	ValueType, Concentration- LimitValue- Type, Concentration- SingleValue- Type	No
Reference	AlphaNumeric (200)	Reference to the source from which this concentration single value is obtained.	Reference, References, Source, Sources	No

Table 3.69:	Table definition	for Concentration lin	nits.
1 uoie 5.07.	ruble definition	for concentration in	mus.

Accepted table names: ResidueLimits, ResidueLimit, MaximumResidueLimits, MaximumResidueLimit, MRLs, MRL.

Concentration limits as data

Maximum Residue Limits (MRL) are provided as data.

- Concentration limits data formats
- Concentration limits from data

3.3.3 Concentration models

Concentration models are distributional models of substance concentrations on foods. They describe both the substance presence (yes/no, with no representing an absolute zero concentration) and the substance concentrations. Concentration models are specified per food/substance combination. This module has as primary entities: Foods Substances Effects

Output of this module is used by: High exposure food-substance combinations Dietary exposures

Concentration models calculation

There are a number of *concentration model types* are available. A basic distinction is between using the empirical concentration data (empirical model), fitting a statistical model to the concentration data (parametric model), or to construct a model from (conservative) limit values. Settings relevant for some of these model types as well as other settings are described under *concentration model settings*.

Concentration data from *Total Diet Studies* does not include variability information. In order to include variability in the concentration models created for TDS samples, the variance of the TDS concentrations may be *estimated using variances of the foods making up the composite TDS food*.

Concentration model types

Empirical model

Data points are sampled at random from the available set. Censored values (non-detect, i.c. < LOD, non-quantifications, i.c. < LOQ) are handled by imputation. If *occurrence patterns* are used, a proportion p_0/p_{ND} of censored values is set as 0. See also *concentration models*.



Figure 3.16: Empirical distribution

Censored spike lognormal model

A binomial model is used to estimate the proportion p of positive values (detects). This is just the proportion observed in the data (unless *agricultural use* data have been used to set a proportion of true zeroes). A lognormal model is fitted to the positive data. This provides estimates of μ and σ , which are the mean and standard deviation of the natural logarithm of the concentration. Simulated concentrations are a censored value with probability $p_{ND} = 1 - p$ or a value sampled from the fitted lognormal distribution with probability p. Censored values (non-detects or nonquantifications) are handled by imputation. If occurrence patterns are used, a proportion p_0/p_{ND} of censored values is set as 0. Minimum requirements: at least two positive concentration values. See also *concentration models*.



NDSpike-LogN

Figure 3.17: Censored Spike Lognormal distribution

Censored-Spike Truncated lognormal model

A binomial model is used to estimate the proportion p of positive values (detects). This is just the proportion observed in the data (unless agricultural use data have been used to set a proportion of true zeroes in which case p is calculated on the remaining proportion). A truncated lognormal model, with LOR as the truncation limit, is fitted to the positive data, leading to estimates of μ and σ , which are the mean and standard deviation of the natural logarithm of the concentration. Simulated concentrations are a censored with probability $p_{ND} = 1 - p$ or a value sampled from the fitted lognormal distribution with probability p. Censored values (non-detects or non-quantifications) are handled by imputation. If occurrence patterns are used, a proportion p_0/p_{ND} of censored values is set as 0. Minimum requirements: at least two positive concentration values, all censored values must have one LOR value. See also *concentration models*.

Censored Lognormal model

A censored lognormal model, with LOR as the censoring limit, is fitted to the data, both positives and censored values. This provides estimates of μ and σ , which are the mean and standard deviation of the natural logarithm of the concentration. If agricultural use data are being used, then a proportion p_0/p_{ND} of censored values will be excluded, where p_0 will be lowered to p_{ND} if it would be higher. Simulated concentrations are sampled from the fitted lognormal distribution. If agricultural use data have been used, simulated concentrations are 0 with probability p_0 or are sampled from the fitted lognormal distribution with probability $1 - p_0$. Minimum requirements: at least one positive concentration value. See also *concentration models*.



NDSpike-TruncLogN

Figure 3.18: Censored Spike Truncated Lognormal distribution



Figure 3.19: Censored Lognormal distribution

Zero-spike censored lognormal model

A mixture distribution of a spike of true zeroes and a censored lognormal model, with LOR as the censoring limit, is fitted to the data (censored values and positives). This provides estimates of p_0 , which is the proportion of true zeroes, and of μ and σ , which are the mean and standard deviation of the natural logarithm of the concentration. Simulated concentrations are 0 with probability p_0 and are sampled from the fitted lognormal distribution with probability $1-p_0$. Minimum requirements: at least one positive concentration value, no agricultural use data for the food-substance combination (which directly specify p_0 , therefore it should not be estimated from the data). See also *concentration models*.



Figure 3.20: Zero Spike Censored Lognormal distribution

Censored spike MRL model

This model simply takes values specified in an input table as *Maximum Residue Levels* (MRL) to be used for the proportion of positive values in the concentration dataset, and can be used to force the use of a pessimistic value.

Summary statistics model

For this model, no individual measurements on raw agricultural commodities are needed. The final estimates of μ and σ are simply provided or pooled or estimated using e.g. a coefficient of variation. Specific use of this model is found in *Total Diet Study* assessments. In general, each TDS food sample is prepared only once, yielding one measurement for a TDS food sample. The variability of the underlying distribution is unknown. However, a rough guess can be made using the e.g. coefficient of variation of the subsamples (in general raw agricultural commodities) that compose the TDS food sample. The estimated standard deviation is *calculated as a pooled estimate using the coefficient of variation of the TDS food*.

Concentration models

Let x denote a random variable from a lognormal distribution. Then, the log transformed variable y = ln(x) is normally distributed with μ and variance σ . The probability density function (p.d.f.) of y may be expressed as:

$$f_y(y, p_0, \mu_y, \sigma_y^2) = p_0 I(y; 0) + (1 - p_0)(1 - I(y; 0)) \cdot \frac{1}{\sqrt{2\pi\sigma_y}} \exp \frac{(y - \mu_y)^2}{2\sigma_y^2}$$

where $p_0 = Pr(y < log(X_{lor})), x_{lor}$ is the limit of reporting and I(y; 0) is an indicator function for $y < log(X_{lor})$. For $p_0 = 0$ the p.d.f. of y reduces to the usual lognormal density. The left truncated density for $y \ge log(X_{lor})$ may be expressed as:

$$f_y(y;\mu_y,\sigma_y^2) = \frac{1}{\sqrt{2\pi\sigma_y}} \exp{\frac{(y-\mu_y)^2}{2\sigma_y^2}}/(1-\Phi(z))$$

with $\Phi(\cdot)$ the standard normal c.d.f. and $z = (\log(x_{lor}) - \mu_y)/\sigma_z$. Model parameters are estimated using maximum likelihood estimation based on the loglikelihood functions specified below. The loglikelihood functions are evaluated in R, using the **optim** algorithm to find estimates for μ_y, σ_y^2 and p_0 .

Mixture zero spike and censored lognormal

The loglikelihood may be expressed as:

$$\log L(p_0, \mu_y, \sigma_y^2) = \sum_{i=1}^{n_0} \log(p_0 + (1 - p_0) \Phi(z_i)) + n_1 \log(\frac{1 - p_0}{\sqrt{2\pi\sigma_y}}) - \sum_{i=n_0+1}^n \frac{(y_i - \mu_y)^2}{2\sigma_y^2} + \frac{1}{2\sigma_y^2} + \frac{1$$

where $y_i = \log(x_i)$, $\Phi(\cdot)$ is the standard normal c.d.f., $z = (\log(x_{i,lor}) - \mu_y)/\sigma_y$, $z_{lor} = (\log(lor) - \mu_y)/\sigma_y$ with n_0 number of censored values $(x_i < x_{i,lor})$, n_1 number of uncensored values $(x_i \ge x_{i,lor})$ and x_i , $i = 1 \cdots n$. Multiple values for LOR are allowed.

Censored lognormal

When $p_0 = 0$ the loglikelihood reduces to:

$$\log L(\mu_y, \sigma_y^2) = \sum_{i=1}^{n_0} \log(\Phi(z)) + n_1 \log(\frac{1}{\sqrt{2\pi\sigma_y}}) - \sum_{i=n_0+1}^n \frac{(y_i - \mu_y)^2}{2\sigma_y^2}$$

Multiple values for LOR are allowed.

Mixture censored spike and truncated lognormal

Ignoring the n_0 values below x_{lor} , the loglikelihood may be expressed as:

$$\log L(\mu_y, \sigma_y^2) = -n_1 \log(1 - \Phi(z)) + n_1 \log(\frac{1}{\sqrt{2\pi\sigma_y}}) - \sum_{i=n_0+1}^n \frac{(y_i - \mu_y)^2}{2\sigma_y^2}$$

Only one value for LOR is allowed.

Mixture censored spike and lognormal

Ignoring the n_0 values below x_{lor} , the loglikelihood may be expressed as:

$$\log L(\mu_y, \sigma_y^2) = n_1 \log(\frac{1}{\sqrt{2\pi\sigma_y}}) - \sum_{i=n_0+1}^n \frac{(y_i - \mu_y)^2}{2\sigma_y^2}$$

Only one value for LOR is allowed.

Imputation of non-detect measurements

A complication in concentration modelling occurs when concentration measurements are reported as below a certain limit. Different names may be used for such a limit, e.g. **limit of detection** (*LOD*), **limit of quantification** (*LOQ*) or the more general term **limit of reporting** (*LOR*). Results that are only reported to be below the LOD (non-detects) or below the LOQ (non-quantifications) are generally referred to as censored values or non-reports. Censored values are a very common phenomenon for some classes of substances like pesticides. When modelling the substance concentrations, they can be handled by incorporating them in a parametric model, or by replacing them with a given value (**imputation**). For imputation, different models are available:

- **Replace by zero**: All left censored measurement values are replaced by zero. This is an optimistic modelling choice.
- **Replace by factor x LOR**: Measurements reported below LOD or LOQ are replaced by a factor times LOQ. When the LOQ is unspecified, then the LOD is used. When the factor is equal to one, this method can be assumed to be a conservative approach.
- **Replace LOD by factor x LOD and LOQ by factor x (LOQ LOD)**: Measurements reported below LOD are replaced by a factor times LOD and measurements reported below LOQ are replaced by a factor times LOQ-LOD (i.e., a value between LOD and LOQ). the factor is set between zero and one.
- Replace non-detects by zero and non-quantifications by factor * LOQ: the factor is set between zero and one.

See also non-detects handling method.

Note, when LOD is not available then it is assumed to be 0. When LOQ is not available then it is assumed to be LOD (or zero if LOD is also not available).

An additional option of the imputation methods that replace the left censored values with some positive value is to *use occurrence frequencies for imputation*. When this option is used, a part of the left-censored values are replaced by a positive value and another part is replaced by zero, based on the *occurrence frequencies* of the modelled foods and substances. Another option is to restrict imputation with positive values to only the authorised substances.

In Figure 3.21 to Figure 3.24, the various scenarios are displayed. Two substances, Fenamidine and Hexythiazox are indicated with a brown box, these substances are authorised.

2,4-DB oleEo Amitrole Bixafen Buprofezin Carbofurar No positive No positive No positive Non Nor 100% 100% 100% 100% 100% Sninaches Zero Zero ero Zero Cens Cens ens Cens e ce ens Hexythiazox enamidone Folpet nazol Empirical No positives No positives Empirical Empirica Empirica 9.9 100% 100% Spinaches ont ent Zero Cens 2eD E C Zero ens loxyni Oryzalir Profenofos Ziran 100% 100% 100% 100% Spinaches Zero Cens Zero Cens Zero Cens Zero Cens Zerc 5

No imputation

Figure 3.21: Tier 1: Censored values are replaced by zero. For Fenamidine and Hexythiazox (brown boxes) authorized use is assumed.



Impute all censored values by factor times LOQ

Figure 3.22: All censored values are replaced by a **constant** x LOR. For Fenamidine and Hexythiazox (brown boxes) authorized use is assumed.



Impute censored values based on authorized uses

Figure 3.23: Censored values are replaced by a **constant** x LOR for authorized uses. For Fenamidine and Hexythiazox (brown boxes) authorized use is assumed.

No imputation except for authorized uses

Deriving the variance of TDS samples from monitoring

Variability of TDS food sample concentrations can be derived using *concentration distributions* for the sub-foods of the TDS food samples (defined by the *TDS compositions*). For each sub-food, e.g. *apple* (sub-food of TDS food *FruitMix*), a coefficient of variation (CV) is specified that is derived using the available monitoring samples. Note that monitoring samples may be composite samples. For *apple*, composite food samples are measured and each sample



Figure 3.24: Tier 1: Censored values are not replaced except for authorized uses (replaced by a **constant** x LOR). For Fenamidine and Hexythiazox (brown boxes) authorized use is assumed.

contains, for instance, 12 apples with unit weight 200 g. So monitoring concentrations, c_{mi} , are based on composite samples with a total weight w_{mi} = 2400 g each.

A TDS food sample is composed of w_i g of food i with i = 1...k, w_i represents the *PooledAmount* in *TDS food* sample compositions table. Then, the concentration of a TDS food sample may be represented as:

$$c_{TDS} = \sum_{i=1}^{k} (w_i \cdot c_i) / \sum_{i=1}^{k} w_i$$

with variance:

$$var(c_{\textit{TDS}}) = \sum_{i=1}^{k} (w_i \cdot var(c_i)) / \sum_{i=1}^{k} w_i$$

and $var(c_i)$ is the variance of concentrations c_i of food *i* with portion sample size w_i .

It is expected that increasing the number of units in a composite sample will have a reverse effect on the variation between concentrations. Suppose TDS food *FruitMix* is composed of $2 \times 200 = 400$ g *apple*. The expected variation between portion sizes of 400 g will be larger than between portion sizes of 2400 g:

$$var(c_i) = var(c_{mi}) \cdot w_{mi}/w_i$$

The variance of the monitoring samples are corrected as follows, calculate:

1. $var(c_{mi}) = \log(CV_{mi}^2 + 1)$

2.
$$var(c_i) = var(c_{mi}) \cdot w_{mi}/w_i$$

3.
$$CV_i = \sqrt{\exp(var(c+i)-1)}$$

Concentration models settings

Calculation settings

Name	Туре	Description
Seed for pseudo-random number generator	Numeric	A value of 0 will use a pseudo-random seed in each run, a value 0 will provide the same results in a repeated run.
Exposure type	ExposureType	The type of exposure considered in the assessment; acute (shor term) or chronic (long-term).
Selected tier	SettingsTemplateType	Specifies all module settings should be set according to a pre-defined tier or using custom settings.
Concentration model types per food-substance combination	ConcentrationModelType- FoodSubstance	The concentration model types used for food/substance combinations.
Default concentration model	<i>ConcentrationModelType</i>	The concentration model type that will be used as default for al food/substance combinations. If this model type cannot be fitte e.g., due to a lack of data, a simpler model will be chosen automatically as a fall-back.
Include MRL fallback model	Boolean	Use the MRL as fallback model in case the occurrence data is insufficient for other concentration modelling options.
Restrict LOR imputation to authorised uses	Boolean	Specifies whether imputation of factor x LOR should be limited authorised uses only.
Censored values replacement	NonDetectsHandlingMethod	How to replace censored values (when not co-modelled, as in censored models).
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	Numeric	Replace censored values by Limit of reporting (LOR), Non-detects (LOD) or Non-quantifications (LOQ) times this factor. Constant (f), e.g. 0.5.
MRL Factor (f x MRL)	Numeric	Use f x MRL as concentration estimate of the MRL models.
Sample based	Boolean	Include co-occurrence of substances in samples in simulations. checked, substance residue concentrations are sampled using th correlations between values on the same sample. If unchecked, any correlation between substances is ignored, substance residu concentrations are sampled ignoring the correlations between values on the same sample.
Impute missing values from available values (if unchecked, missing values are imputed with 0)	Boolean	If checked, in procedure of EFSA Guidance 2012, Appendix 1 impute missing values using substance based concentration models. If unchecked, missing values are imputed by 0.
Correlate imputed values with sample potency	Boolean	If checked, in procedure of EFSA Guidance 2012, Appendix 1 correlate high imputed values with high cumulative potency samples. If unchecked, random imputation.
Use occurrence frequencies for imputation	Boolean	Use of occurrence frequencies (e.g., agricultural use frequencie is relevant for imputation of censored values in the concentration data. Part of the observed censored values and missing values in be imputed with zero when the occurrence frequency is smaller than 100%. If checked, occurrence frequencies are expected as input of this action, otherwise 100% potential presence is assumed for all substances on all foods.
Total diet study concentration data	Boolean	Specifies whether exposure is based on sampling data from tota diet studies.
Multiple substances analysis	Boolean	Specifies whether the assessment involves multiple substances.
Compute cumulative exposures	Boolean	Specifies whether the assessment involves multiple substances a results should be cumulated over all substances.

Table 3.70: Calculation settings for module Concentration models.

Uncertainty settings

	, e	
Name	Туре	Description
Parametric uncertainty	Boolean	For resample concentrations: specifies whether the uncertainty assessment is based on a parametric approach.
Resample concentrations	Boolean	Specifies whether concentrations are resampled by empirical bootstrap or using a parametric uncertainty model.

Table 3.71: Uncertainty settings for module Concentration models.

Concentration models tiers

In addition to the possibility for users to work with their own choices for all settings, MCRA implements four tiers from two documents:

- The optimistic and pessimistic basic assessments from the EFSA 2012 Guidance on the Use of Probabilistic Methodology for Modelling Dietary Exposure to Pesticide Residues (EFSA (2012)).
- Tier 1 and 2 from the *European Commission working document SANTE-2015-10216 rev.* 7 (2018) on risk management aspects related to the assessment of cumulative exposure (EC (2018)).

Overview

Table 3.72: Tier overview for module Concentration mode

N EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
D Empirical fa co co tr ti n	Empirical	Empirical	Empirical	Empirical	NonDe- tect- SpikeLog- Normal	NonDe- tect- SpikeLog- Normal	Empirical	Empirical	Emp
Ir false cl M fa ba m	false	false	false	false	true	true	false	false	false
R false st L ir p ta ti ti tc au th ri u	false	false	false		false	false	false	false	false

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
C Se V U r C P n	Replace- ByLOR	Replace- ByLOR	Replace- ByLOR	Replace- ByLOR	Replace- ByZero	Replace- ByLOR	Replace- ByLOR	Replace- ByLOR	Replace- ByLOR	Rep ByL
F to f (1) x L o f x L o f x L to f x L to f L L	0.5	0.5	0.5	0.5		1	1	0.5	0.5	0.5
S p b	true	true	true	true	true	true	true	true	true	true

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
In p m ir v u f f a a a if a a if i a a if i a a a if i v v v v v v v v v v v v v	true	true	true	true	false	true	true	true	true	true
C rec la ir p va u w sa p p te	true	true	false	false	false	true	true	true	true	false
U o c r c f f f c i i r f c i i r f f f f c i r t c	true	true	true	true	false			true	true	true

Table 3.72 - continued from previous page
N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
P m ri un ce ta	false	false	false	false	false	true	false	false	false	false
F. tee saap ph ez cce inn th cce tr ti- ti- ti- ti- ti- ti-	false	false	false	false		false	false	false	false	false
U su st co vo si ru	true	true	true	true				true	true	true
Si st co vo si m	UseMost- Toxic	UseMost- Toxic	DrawRan- dom	DrawRan- dom				UseMost- Toxic	UseMost- Toxic	Drav dom

Table 3.72 – continued from previous page

 Table 3.72 – continued from previous page									ge
EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
<pre> true true true true true true true true</pre>	true	true	true				true	true	true
A false	false	true	true				false	false	true
F false	false	false	false				false	false	false

										-
N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
U ez tr o- la ti	true	true	true	true				false	false	false
T ol fc ez tr o- la ti	10	10	10	10						
R st ez tr o- la ti tc eq N	true	true	true	true						
R st ez tr o- la ti tc au tr tc au tr	true	true	true	true						
Ir pr w te co co tr ti	true	true	true	true				true	true	true

Table 3.72 - continued from previous page

						Tab	le 3.72 – cor	ntinued from	previous pag	le
N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202: Acu Tier
W te ce tr ti va (l	0.1	0.1	0.05	0.05				0.1	0.1	0.05
R st w te ir p ta ti tc tf fr m tc st	true	true	true	true				true	true	true
R st w te ir p ta ti tc au th ri u	false	false	false	false				false	false	false
R st w te ir p ta ti tc aj p s u st	false	false	false	false				true	true	true

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
A pl oc re pa te po ce ag	false	false	true	true				false	false	true
T ge le	External	External	External	External	External	External	External	External	External	Exte
Se uj us fr qu tc			true	true						true
R st us p c c e a g uj s c t c t c t c t c t t t t t t us			true	true						true
M F to (f x N						1	1			
C ccc tr ti li: fil te ex ccc fa										

N EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
Ir cl fc ca ca m it ca tr tr ti									
F ca co m it su su st co co co co co co co co co co co co co									
A ju m fa tc fc tt fc ca fc ca fc tt tt									

Table 3.72 - continued from previous page

N EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
U du te m is ti su st cu st cu si fc fc ca cu m it									
Ir cl pi cc dd ti of th fc cc fc in m su m rc pj m									

Table 3.72 - continued from previous page

The sections below describe the settings specified by each tier in detail.

Retrospective dietary CRA (EC 2018) - Acute / Tier I

Name	Setting	From input tier	ln module	
Default concentration model	Empirical			
Restrict LOR imputation to	false			
authorised uses				
Censored values replacement	Replace- ByLOR			
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5			
Sample based	true			
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true			
Correlate imputed values with sample potency	true			
Use occurrence frequencies for imputation	true			
Parametric uncertainty	false			
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations	
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations	
Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations	
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations	
Account for substance authorisations in substance conversions	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations	
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) -	Concen- Chapte trations	r 3. Modules

Table 3.73: Tier definition for Retrospective dietary CRA (EC 2018) - Acute / Tier I.

Retrospective dietary CRA (EC 2018) - Chronic / Tier I

Name	Setting	From input tier	ln module	
Default concentration model	Empirical			
Restrict LOR imputation to	false			
authorised uses	laise			
Censored values replacement	Replace- ByLOR			
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5			
Sample based	true			
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true			
Correlate imputed values with sample potency	true			
Use occurrence frequencies for imputation	true			
Parametric uncertainty	false			
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations	
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations	
Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations	
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations	
Account for substance authorisations in substance conversions	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations	
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) -	Concen- Chapte trations	r 3. Modules

Table 3.74: Tier definition for Retrospective dietary CRA (EC 2018) - Chronic / Tier I.

Retrospective dietary CRA (EC 2018) - Acute / Tier II

Name	Setting	From input tier	ln module	
Default concentration model	Empirical	[
Include MRL fallback model	false			
Restrict LOR imputation to	false			
authorised uses				
Censored values replacement	Replace- ByLOR			
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5			
Sample based	true			
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true			
Correlate imputed values with sample potency	false			
Use occurrence frequencies for imputation	true			
Parametric uncertainty	false			
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations	
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations	
Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations	
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations	
Account for substance authorisations in substance conversions	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations	
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) -	Concen- Chapte trations	r 3. Modules

Table 3.75: Tier definition for Retrospective dietary CRA (EC 2018) - Acute / Tier II.

Retrospective dietary CRA (EC 2018) - Chronic / Tier II

Name	Setting	From input tier	ln module	
Default concentration model	Empirical			
Include MRL fallback model	false			
Restrict LOR imputation to	false			
authorised uses				
Censored values replacement	Replace- ByLOR			
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5			
Sample based	true			
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true			
Correlate imputed values with sample potency	false			
Use occurrence frequencies for imputation	true			
Parametric uncertainty	false			
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations	
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations	
Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations	
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations	
Account for substance authorisations in substance conversions	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations	
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) -	Concen- Chapte trations	r 3. Modules

Table 3.76: Tier definition for Retrospective dietary CRA (EC 2018) - Chronic / Tier II.

Retrospective dietary CRA (EFSA 2012) - Optimistic

Use the optimistic model settings according to the EFSA Guidance 2012. Concentration values are sampled using a sample-based empirical distribution. Available processing factors are applied. No unit variability model should be applied.

Name	Setting	From input tier	In module
Default concentration model	Empirical		
Include MRL fallback model	false		
Censored values replacement	Replace- ByZero		
Sample based	true		
Impute missing values from available values (if unchecked, missing values are imputed with 0)	false		
Correlate imputed values with sample potency	false		
Use occurrence frequencies for imputation	false		
Parametric uncertainty	false		
Target level	External	Retrospec- tive dietary CRA (EFSA 2012) - Optimistic	Hazard character- isations

Table 3.77: Tier definition for Retrospective dietary CRA (EFSA 2012) - Optimistic.

Retrospective dietary CRA (EFSA 2012) - Acute / Pessimistic

Acute probabilistic exposure assessment using the pessimistic model settings according to the EFSA Guidance 2012. Only processing factors > 1 are applied. For unit variability, the Beta distribution is applied.

Name	Setting	From input tier	In module
Default concentration model	NonDe- tect- SpikeLog- Normal		
Include MRL fallback model	true		
Restrict LOR imputation to authorised uses	false		
Censored values replacement	Replace- ByLOR		
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	1		
MRL Factor (f x MRL)	1		
Sample based	true		
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true		
Correlate imputed values with sample potency	true		
Parametric uncertainty	true		
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Concen- trations
Target level	External	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Hazard character- isations

Table 3.78: Tier definition for Retrospective dietary CRA (EFSA 2012) - Acute / Pessimistic.

Retrospective dietary CRA (EFSA 2012) - Chronic / Pessimistic

Chronic probabilistic exposure assessment using the pessimistic model settings according to the EFSA Guidance 2012. Only processing factors > 1 are applied.

Name	Setting	From input tier	ln module
Default concentration model	NonDe- tect- SpikeLog- Normal		
Include MRL fallback model	true		
Restrict LOR imputation to authorised uses	false		
Censored values replacement	Replace- ByLOR		
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	1		
MRL Factor (f x MRL)	1		
Sample based	true		
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true		
Correlate imputed values with sample potency	true		
Parametric uncertainty	false		
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2012) - Chronic / Pessimistic	Concen- trations
Target level	External	Retrospec- tive dietary CRA (EFSA 2012) - Chronic / Pessimistic	Hazard character- isations

Table 3.79: Tier definition for Retrospective dietary CRA (EFSA 2012) - Chronic / Pessimistic.

Retrospective dietary CRA (EFSA 2022) - Acute / Tier I

Name	Setting	From input tier	In module	
Default concentration model Include MRL fallback model Pagtriat LOP imputation to	Empirical false			
authorised uses	Taise			
Censored values replacement	Replace- ByLOR			
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5			
Sample based	true			
values (if unchecked, missing values are imputed with 0)	true			
Correlate imputed values with sample potency	true			
Use occurrence frequencies for imputation	true			
Parametric uncertainty	false			
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations	
Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations	
Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations	
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations	
Account for substance authorisations in substance conversions	false	Retrospec- tive dietary	Concen- trations	
		CRA (EFSA 2022) - Acute / Tier I	Chapte	r 3. Module

Table 3.80: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier I.

Retrospective dietary CRA (EFSA 2022) - Chronic / Tier I

Name	Setting	From input tier	In module
Default concentration model Include MRL fallback model Restrict LOR imputation to authorised uses	Empirical false false		
Censored values replacement	Replace- ByLOR		
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5		
Sample based Impute missing values from available values (if unchecked, missing values	true true		
Correlate imputed values with sample potency	true		
Use occurrence frequencies for imputation	true		
Parametric uncertainty Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Account for substance authorisations in substance conversions	false	Retrospec- tive dietary	Concen- trations
		CRA (EFSA 2022) - Chronic / Tier I	Chapte

Table 3.81: Tier definition for Retrospective dietary CRA (EFSA 2022) - Chronic / Tier I.

Retrospective dietary CRA (EFSA 2022) - Acute / Tier II

Name	Setting	From input tier	In module	
Default concentration model Include MRL fallback model Restrict LOR imputation to authorized uses	Empirical false false			
Censored values replacement	Replace- ByLOR			
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5			
Sample based	true			
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true			
Correlate imputed values with sample potency	false			
Use occurrence frequencies for imputation	true			
Parametric uncertainty	false			
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations	
Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations	
Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations	
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations	
Account for substance authorisations in substance conversions	true	Retrospec- tive dietary	Concen- trations	
		CRA (EFSA 2022) - Acute / Tier II	Chapte	r 3. Mo

Table 3.82: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier II.

Retrospective dietary CRA (EFSA 2022) - Chronic / Tier II

Name	Setting	From input tier	In module	
Default concentration model	Empirical			
Include MRL fallback model	false			
Restrict LOR imputation to	false			
authorised uses	Deglass			
	ByLOR			
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5			
Sample based	true			
values (if unchecked, missing values are imputed with 0)	true			
Correlate imputed values with sample potency	false			
Use occurrence frequencies for imputation	true			
Parametric uncertainty	false			
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations	
Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations	
Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations	
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations	
Account for substance authorisations in substance conversions	true	Retrospec- tive dietary	Concen- trations	
		CRA (EFSA 2022) - Chronic / Tier II	Chapte	r 3. Modu

Table 3.83: Tier definition for Retrospective dietary CRA (EFSA 2022) - Chronic / Tier II.

Prospective dietary CRA (EFSA 2023) - Acute / Tier II

Table 3.84:	Tier	definition	for	Prospective	dietary	CRA	(EFSA	2023) -	-
Acute / Tier	·II.								

Name	Setting	From input tier	In module
Default concentration model	Empirical		1
Include MRL fallback model	false		
Restrict LOR imputation to	false		
authorised uses			
Censored values replacement	Replace- ByLOR		
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5		
Sample based	true		
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true		
Correlate imputed values with sample potency	false		
Use occurrence frequencies for imputation	true		
Parametric uncertainty	false		
Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations

Name	Setting	From input tier	In module
Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Use extrapolation rules	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Impute water concentrations	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations

Table 3.84 - continued from previous page

Name	Setting	From input tier	In module
Restrict water imputation to the five most toxic substances	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Restrict water imputation to approved substances	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Include focal commodity concentrations	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Focal commodity substance occurrence percentage	20	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Adjustment factor for the focal food/substance concentration	1	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations

Table 3.84 – continued from previous pac	vious page	nued from	3.84 - continu	Table
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Name	Setting	From input tier	In module
Use deterministic substance conversions for focal commodity	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Apply occurrence pattern percentages	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Occur- rence patterns
Target level	External	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Hazard character- isations

Table 3.84 - continued from previous page

Prospective dietary CRA (EFSA 2023) - Chronic / Tier II

Table 3.85: Tier definition for Prospective dietary CRA (EFSA 2023) - Chronic / Tier II.

Name	Setting	From input tier	In module
Default concentration model	Empirical		
Destaint LOD international	faise		
Restrict LOR imputation to	false		
authorised uses			

Name	Setting	From input tier	In module
Censored values replacement	Replace- ByLOR		
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5		
Sample based	true		
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true		
Correlate imputed values with sample potency	false		
Use occurrence frequencies for imputation	true		
Parametric uncertainty	false		
Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations

Name	Setting	From input tier	In module
Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Use extrapolation rules	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Impute water concentrations	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations

Table 3.85 - continued from previous page

Name	Setting	From input tier	ln module
Restrict water imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Restrict water imputation to approved substances	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Include focal commodity concentrations	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Focal commodity substance occurrence percentage	20	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Adjustment factor for the focal food/substance concentration	1	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Use deterministic substance conversions for focal commodity	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Apply occurrence pattern percentages	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Occur- rence patterns
Target level	External	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Hazard character- isations

Table 3.85 - continued from previous page

Prospective dietary CRA (EFSA 2024) - Acute / Tier II

Table 3.86: Tier definition for Prospective dietary CRA (EFSA 2024) -Acute / Tier II.

Name	Setting	From input tier	In module	
Default concentration model	Empirical			
Include MRL fallback model	false			
Restrict LOR imputation to authorised uses	false			
Censored values replacement	Replace- ByLOR			
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5			
Sample based	true			
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true			
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Name	Setting	From input tier	In module
Correlate imputed values with sample potency	false		
Use occurrence frequencies for imputation	true		
Parametric uncertainty	false		
Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Fix duplicate substance allocation inconsistencies	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Use extrapolation rules	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Impute water concentrations	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations

Table 3.86 - continued from previous page

Name	Setting	From input tier	ln module
Restrict water imputation to approved substances	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Include focal commodity concentrations	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Focal commodity substance occurrence percentage	20	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Adjustment factor for the focal food/substance concentration	1	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Use deterministic substance conversions for focal commodity	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Include processed derivatives of the focal food in measurement replacement	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations

Name	Setting	From input tier	In module
Apply occurrence pattern percentages	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Occur- rence patterns
Target level	External	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Hazard character- isations

Table 3.86 - continued from previous page

Prospective dietary CRA (EFSA 2024) - Chronic / Tier II

Table 3.87: Tier definition for Prospective dietary CRA (EFSA 2024) -Chronic / Tier II.

Name	Setting	From input tier	In module
Default concentration model	Empirical		
Include MRL fallback model	false		
Restrict LOR imputation to authorised uses	false		
Censored values replacement	Replace- ByLOR		
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5		
Sample based	true		
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true		
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Name	Setting	From input tier	In module
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Correlate imputed values with sample potency	false		
Use occurrence frequencies for imputation	true		
Parametric uncertainty	false		
Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Fix duplicate substance allocation inconsistencies	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Use extrapolation rules	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Impute water concentrations	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations

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Name	Setting	From input tier	ln module
Restrict water imputation to approved substances	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Include focal commodity concentrations	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Focal commodity substance occurrence percentage	20	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Adjustment factor for the focal food/substance concentration	1	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Use deterministic substance conversions for focal commodity	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Include processed derivatives of the focal food in measurement replacement	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Apply occurrence pattern percentages	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Occur- rence patterns
Target level	External	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Hazard character- isations

Table 3.87 - continued from previous page

Concentration models uncertainty

When using empirical distributions, concentration model uncertainty is covered by the the inputs. I.e., concentration models can be recomputed from *resampled/bootstrapped* concentration data. This happens for both the univariate concentration models, being recomputed from the bootstrapped residue collections for each food and substance, and also for the samples of the sample-based approach that are re-generated from the bootstrapped samples (including the necessary steps of missing value imputation and imputation of censored values).

When parametric uncertainty is preferred over empirical bootstrapping, the parameters of the univariate concentration models fitted as a parametric distributions can be *resampled parametrically*.

Let x denote a random variable from the specified distribution. The log transformed variable y = ln(x) is normally distributed with mean μ_y and variance σ_y . The maximum likelihood estimates are $\hat{\mu}_y$ and $\hat{\sigma}_y$. In each bootstrap sample, values are drawn from a normal distribution where the maximum likelihood estimates are replaced by ($\hat{\mu}_y^*$, $\hat{\sigma}_y^*$).

Calculation of concentration models

Concentration models can be computed from concentration data.

• Concentration models calculation

Inputs used: Concentrations Concentration limits Active substances Modelled foods Substance authorisations Occurrence frequencies Relative potency factors Concentration distributions Total diet study sample compositions

Settings used

• Calculation Settings

3.3.4 Concentrations

Concentrations data are analytical measurements of chemical substances occurring in food samples. In their simplest form, concentration data can just be used as provided by datasets. Optionally, concentrations data can be manipulated for active substances, extrapolated to other foods, and/or default values can be added for water.

This module has as primary entities: Foods Substances Populations

Output of this module is used by: Single value concentrations Occurrence patterns Concentration models Modelled foods

Concentrations from data

Concentrations calculation

Occasionally, concentrations of substances measured in food samples are exceeding a specified concentration limit e.g. the *Maximum Residue Limits* (MRL). An MRL is the highest level of a substance that is legally tolerated in or on food or feed when substances are applied correctly. *Filter samples* exceeding the concentration limits filter out all samples where one of the substances measured is exceeding the *MRL*.

Substance conversions rules may be used to *convert* concentration data at the level of measured substances to concentration data at the level of potentially active substances. These rules (provided as data) may be applicable, for example, when a measured substance represents multiple substances and these measurements should be converted into measurement values for these substances. This conversion may depend on *substance authorisations* which provides information on the likelihood of certain translations to occur. *points of departure* or *relative potency factors* might be needed when the substance conversion should select the most toxic candidate in case a measured substance translates to multiple active substances.

If there are only a few measurements in the concentration data, then *extrapolation of concentration data* may be desired. In that case, *food extrapolation rules* may be provided to specify per food the alternative foods from which extrapolation is allowed. The extrapolation of concentrations will then be performed within this module and the results are included in the resulting active substance concentrations data. *Substance authorisations* and/or *concentration limits* may be used to further restrict the to-food/from-food combinations per substance for which extrapolation is possible.

Concentration data for water are often not available in the concentration data, but it may be desirable to include them in the assessments. For this, *imputation* of low-tier, deterministic estimates of water concentrations of the most toxic substances may be used to include (typically conservative) estimates in the calculations.

In some scenarios it may be desired to perform a prospective analysis in which anticipated (or foreground) *focal commodity concentration data* for a particular focal commodity food (and substance) is added to, or replaces part of the background concentration data that is used for the null-scenario. The concentrations module offers various options to perform such *focal commodity scenario analyses*.

It is also possible to *filter (or subset) samples by specific sample properties* (e.g., year, location). This can be done by checking the option to *filter samples by specific property values (subset selection)*.

Filter samples exceeding the concentration limits

If the option **Filter samples exceeding the concentration limits** is checked, all samples with one or more substance concentrations exceeding the *MRL* are filtered out. Then a **concentration limit filter exceedance factor** (*factor*) is specified, which filters out samples with at least one substance concentration higher than $factor \cdot MRL$.

If the option **Filter samples exceeding the concentration limits** remains unchecked all samples are retained in the analysis.

Substance conversion

When concentration data at the level of measured substances have to be converted into concentration data at the level of *active substances* (or perhaps also inactive substances), then *substance conversion rules* are specified to provide the rules, the so-called residue definitions. This section describes the basic substance conversion, and then the refinements using available *substance authorisations*.

For each measured substance in the concentration data, there may be zero or more conversion rules, each one linking to an active or inactive substance. Each rule is represented by a record in the substance conversion data source. Substance conversion rules may specify a link to an exclusive substance or not. For an exclusive conversion it is assumed that only one substance is present in the sample, therefore the measured substance is considered to be just one of the linked substances. Another possibility is that measured substances link to one or more exclusive substances, plus one (non-exclusive) substance that is considered a metabolite of the other exclusive substances. The metabolite can occur together with any of the exclusive substances. It is assumed that either all conversion rules linked to a measured substance are marked as exclusive (case 1), or exactly one rule is marked as exclusive and the others are marked as not exclusive (case 2). If this does not apply for any set of rules linked to a measured substance, data are regarded as erroneous.

Four methods for substance conversion are implemented:

1. Allocate most potent: for each measured substance, the linked substances are restricted to the active substances of interest. The concentration of the measured substance is assigned to the most potent active substance in this set. Potency is specified by the *relative potency factors*. All other candidate active substances are assigned a zero concentration. I.e., the measured substance concentration is allocated for 100% to the most potent substance specified by the conversion rules and for this allocation, the concentration or LOQ/LOD is multiplied by the molecular weight correction factor.

2. Random allocation: one of the conversion rules is drawn randomly (with equal probability), including the rules of both active and other substances. Then, the rule is used as follows to generate active substance concentrations:

- If the conversion rule is marked as exclusive, the concentration or LOQ/LOD is allocated to the linked substance.
- If the conversion rule is marked as not exclusive, a proportion *p*, as specified by the rule, of the concentration or LOQ/LOD is allocated to the linked substance. The remaining proportion (*1-p*) is allocated to the substance that is linked to the measured substance in a conversion rule marked as exclusive (in this case it is assumed that exactly one record per measured substance is marked as exclusive).

All assigned concentrations are multiplied by the molecular weight correction factor. All unselected candidate substances are assigned a zero concentration.

3. Nominal estimate: the substances specified through the conversion rules are allocated with a nominal value based on all possible conversion rules. This may be regarded as the nominal or average allocation value of the random sampling method.

- All conversion rules are marked as exclusive: the measured substance concentration is allocated over all *n* active substances specified with equal proportions *1/n*, accounting for the molecular weight correction factor for all substances.
- Precisely one conversion rule is marked exclusive and n conversion rules are marked as not exclusive: the measured substance concentration is allocated over all active substances specified, with a proportion 1/2 + 1/n for the substance belonging to the exclusive conversion rule, and equal proportions 1/n for the other substances, accounting for the molecular weight correction factor for all substances.

4. Allocate all: the concentration of a measured substance is allocated to each active substance associated with the measured substance as if it were the most potent substance. I.e., the same measured substance is allocated to all associated active substances simultaneously. This method is not sensible when using it in a cumulative assessment, but it is of use in substance screening assessments, where in a combined analysis of multiple substances all active substances are considered independently.

Use of substance authorisations in substance conversion

When *substance authorisations* are available, these can be used to exclude conversions of measured substances to unauthorised substances on a given food. The information is used as follows in the substance conversion procedures:

1. Allocate most potent: the set of candidate active substances from which the most potent active substance is to be drawn is reduced to only the substances with authorised uses. However, if none of the candidate active substances is authorised, then the most potent of the unauthorised substances is selected for active substance allocation.

Active stance	sub-	RPF	Authorised use	Active stance	sub-	RPF	Authorised use	Measured substance allo- cated to
AS1		1	true	AS2		2	true	AS1
AS1		1	false	AS2		2	true	AS2
AS1		1	true	AS2		2	false	AS1
AS1		1	false	AS2		2	false	AS1

Table 3.88: Most toxic allocation of a measured substance to active substances based on RPFs and authorised use

In Table 3.88, results of most toxic allocation.

2. Random allocation: the set of conversion rules from which to draw is reduced to the rules linking to authorised substances or the non-exclusive substance (thus allowing the selection of a possibly unauthorised metabolite of an authorised substance). If none of the conversion rules links to an authorised substance, then one rule is drawn from the full set of all (unauthorised) conversion rules.

Table 3.89: Random allocation of a measured substance to active substances based on authorised use

Active stance	sub-	RPF	Authorised use	Active stance	sub-	RPF	Authorised use	Measured substance allo- cated to
AS1		1	true	AS2		2	true	AS1 or AS2
AS1		1	false	AS2		2	true	AS2
AS1		1	true	AS2		2	false	AS1
AS1		1	false	AS2		2	false	AS1 or AS2

In Table 3.89, results of random allocation.

3. Nominal estimate: the set of conversion rules is reduced in the same way as for random active substance allocation. Nominal calculation is performed on the resulting set of conversion rules.

4. Allocate all: For this method, the same rules apply as for *allocate most potent*. The set of candidate active substances that are to be allocated is reduced to only the substances with authorised uses. Hence, a substance is not allocated when it is not authorised and there is at least one other candidate active substance that is authorised. However, if none of the candidate active substances is authorised, then the most potent of the unauthorised substances is selected for active substance allocation.

Multiple allocations of the same active substance in one sample

In some datasets substance conversion can cause the same active substance to be allocated multiple times in one sample. For example, when an active substance is measured directly, but also a measurement is recorded for a (measured) substance that converts to the active substance. By default, MCRA does not accept such cases, because often it is associated with errors in the data. Therefore, an error will be reported with a message

"Unexpected substance translation in sample xxx: substance X is translated from multiple measured substances."

However, if such cases are known to exist and accepted in the data, then a method is available to *fix duplicate substance allocation inconsistencies*. If active substance allocation leads to multiple allocated measurements then the following procedure is implemented for resolving these inconsistencies:

- If any measurement is positive or zero then: clone one of these records to create the "aggregate measurement record"; to make this selection deterministic, prefer records that have measured-substance equal to active-substance over other records, and take the record with the highest residue. Update the measured value of the clone with the mean of all positive/zero measurements.
- If all allocated active substance measurements are censored, then clone one of the censored values. Here also, records that have measured-substance equal to active-substance are preferred over other records, and then the measurement with the smallest LOQ/LOD is used.

Note that these rules are quite generic. They work quite well even when there are many measurements for the same active substance. In practice, only a few (two) are expected.

Food extrapolation

If the *food extrapolation* setting has been checked, extrapolation of concentrations is performed for all food/*active substance* combinations for which:

- 1. the number of measurements in the analytical scope is smaller than a given threshold for extrapolation (default 10), and
- 2. there is an *extrapolation rule* allowing extrapolation of concentrations from one or more other foods (the from-food(s)) to the given food (the to-food), and
- 3. (optional criterion:) the substance is associated with authorised use for both foods, and
- 4. (optional criterion:) *concentration limits (e.g. MRLs)* on the from-food and to-food exist and are equal. Note: if the **active substance** is not a **measured substance**, then the MRL check has to be made per measurement at the level of the measured substance which provided the concentrations assigned to the active substance.

Food extrapolation is performed by one of the following procedures: 1) Substance-specific imputation of missing values by extrapolated measurements, or 2) Extrapolation of complete samples for multiple substances.

1. Substance-specific imputation of missing values by extrapolated measurements

The missing values in the active substance concentrations of the to-food are imputed in a random order by active substance concentrations (positive, censored or zero) from a randomised list obtained from the fromfood(s). By matching the randomised lists, each from-food measurement is assigned at most once, so after extrapolation there may still be missing values left, or not all measurements of the from-food(s) may have been used for extrapolation.

Note: In this method, it is assumed that the to-food has a sufficient number of samples. No extrapolation is applied for foods with no samples at all, and data gaps will also remain for foods with fewer than n samples, because no new samples are added.

Note: the resulting *occurrence patterns* will be random with respect to the extrapolated substances, i.e., observed occurrence patterns for the from-food are not extrapolated to the to-food.

1. Extrapolation of complete samples for multiple substances

(not yet implemented)

All samples of the from-food(s), i.e., complete samples with data for all active substances, are copied as samples for the to-food and added to the existing to-food samples. For example, extrapolate all apple sample records to the available pear sample records. However, measurements for substances that do not fulfil the (optional) criteria 3 and 4 above are non-valid extrapolations and are replaced by missing values. The status of the extrapolated samples is stored to distinguish between extrapolated and non-extrapolated sample records. Note that this method maintains correlations in the occurrence patterns and postpones imputation of MVs until the concentration models step.

Imputation of water concentrations

If water has been selected as an additional source of exposure, but concentration data is missing, then, fixed concentration values can be assigned to water for the five most toxic *active substances*, with the toxicity ranking being based on the *relative potency factors*. For all other substances, zero concentrations are imputed. The default imputation

value is 0.05 μ g/L, but this value can be chosen as a setting. If specified, *substance authorisations* may be used to restrict to the set of active substances for which water concentrations are imputed to only those for which concentrations may be expected from *authorised use*.

Focal commodity scenario analysis

There are different *methods* for modifying the (background) concentration data for specific (prospective) focal commodity scenario analyses. In the frontend these methods are accessible through the option *include focal commodity concentrations* (see Figure 3.25). Checking this option will open the focal commodity scenario analysis form (see Figure 3.26) where the method and the focal food/substance can be selected, and associated other settings can be configured, see also *focal food concentrations*.

Concentration settings	Save changes
Selected tier Custom	- i
Filter samples exceeding the concentration limits	0
Use substance conversion rules	0
Use extrapolation rules	0
Impute water concentrations	0
Include focal commodity concentrations	0
Filter samples by specific property values (subset selection)	0

Figure 3.25: Concentration settings: include focal commodity concentrations.

The following focal commodity replacement methods are available:

- **Replace samples with focal commodity samples:** This method will replace all samples for the selected focal commodity food by samples from the *focal commodity concentration data*. Food sample replacement is substance independent, therefore all substance concentrations of the focal commodity food in the background concentration data are replaced.
- **Append focal commodity samples:** This method adds the *focal commodity samples* of the selected focal commodity food to the background concentration data. This method is independent of the substance and may be a useful approach when the substances measured in the field trial do not overlap with the substances of the (background) concentration data. In this case, the focal commodity substance concentrations will be missing for the background concentration data and (also the other way around) the substance concentrations of all other substances will be considered missing for the focal commodity samples. These missing values may be imputed at a later stage according to the standard procedures.
- **Replace by focal commodity samples:** The focal food/substance measurements are obtained from *focal commodity samples*. Here, substance measurements of the focal commodity food in the background concentration data are replaced by randomly assigned substance measurements of the focal commodity samples.
- Replace by concentration limits from data: The focal food/substance measurements are obtained from

Focal commodity concentration scenario settings	Save changes	anges	
Focal commodity concentrations replacement method * Replace measurements of focal food/substance combinations with proposed concentration limit value from data	- 0		
Include focal commodity concentrations * Oranges (P0110020A)	- 0		
Focal commodity substances * Abamectin (RF-00000011-VET)	- 0		
Focal commodity substance occurrence percentage * 20	0		
Adjustment factor for the focal food/substance concentration * 1	0		
✓ Use deterministic substance conversions for focal commodity	0		
Filter processed focal commodity samples	0		

Figure 3.26: Focal commodity concentration scenario settings form. This form is a sub-form of the concentrations module panel.

focal commodity samples. Here, substance measurements of the focal commodity food in the background concentration data are replaced by the concentration limit value (e.g., *MRL* s) obtained from the provided *concentration limits data*.

- **Replace by proposed concentration limits:** Here, substance measurements of the focal commodity food in the background concentration data set are replaced by the proposed concentration limit values (e.g., MRLs in mg/kg) specified in the interface.
- **Remove measurements of focal food/substance combinations:** Simply remove all background concentrations for the selected focal commodity food and substance combination, values are not replaced. This method is useful when a separate analysis is desired for the background and foreground concentrations.

Depending on the selected *replacement method*, different additional modelling options become available.

Use the *focal commodity substance occurrence percentage* to specify an occurrence percentage for the combination of focal food and substance. For percentages less than 100%, a part of concentrations of the focal commodity food and substance is replaced with the focal concentrations, the remaining part of the concentrations is replaced with zero concentrations. E.g., when background concentrations of the substance Abamectine on oranges are replaced with an MRL value, then a specified occurrence percentage of 40% will replace 40% of the measurements with the MRL, and 60% of the measurements with a zero concentration. Note that the replacement is random, i.e., each substance measurement has a probability of being assigned to a focal concentration (>0) or a zero. So, the actual replacement percentage may differ from the specified percentage. This option is, for example, used to simulate a percentage of agricultural use.

Use the *adjustment factor for the focal food/substance concentration* to adjust the (positive) concentrations of the focal food and substance measurements. This factor is used when the focal commodity concentrations (e.g., from field trials) are assumed to be higher than what is expected in practice. In this case, this factor could be set for instance, to the expected ratio of mean monitoring concentration and mean field trial concentration. Note that for replacement by focal commodity measurements, this factor will only adjust the positive concentrations and not the LORs.

By default, the focal commodity substance measurements are replaced before the optional step of *converting the concentrations from measured to active substance concentrations*. This means that the same substance conversion rules apply for the background and foreground concentrations when converting measured substance concentrations to active substance measurements. Alternatively, substance measurements can also be replaced after allocation of the active substance. This can be done using the option to *use deterministic substance conversion factors*. When this option is checked, *deterministic substance conversion factors* are used for the focal commodity food and substance to convert measurements to the level of *active substances* and replacement of background concentrations by foreground concentrations is done after active substance allocation. Figure Figure 3.27 illustrates how active substance allocation



is done for the foreground and background concentration data with and without this option.

Figure 3.27: Active substance allocation of the background and foreground concentration data with and without the use of deterministic substance conversion factors.

When the occurrence data includes samples for processed derivatives of the selected focal food(s) (e.g., orange juice samples for focal food oranges), the substance measurements of the focal substance for the samples of the processed food can also be replaced. Checking the option *include processed derivatives of the focal food* replaces focal substance measurements of the processed derivatives of the focal food replaces focal substance for the focal food, focal substance, and processing type of the derivative, then this processing factor is used to first correct the focal concentrations for processing. Alternatively, it is also possible to just exclude the samples of the focal commodity, which can be done using the option to *filter processed focal commodity samples*.

Note that when also using *substance authorisations*, the focal food and substance combination will be treated as authorised, even if there is no authorisation supplied for the combination. The approved authorisation status is considered to be part of this scenario analysis.

Filter samples by specific sample properties

When the option to *filter samples by specific property values (subset selection)* is checked in the *main panel of the concentrations module*, a new form will appear in which various sample property subsets can be specified (see Figure 3.28). The visibility/availability of the sample property filters depends on the availability of these properties in the data. The following filters are available:

- The options to *filter samples by year* and *filter samples by month* allow the user to filter the samples by sampling date. The additional option to *include samples with missing sampling date* determines whether samples for which the sampling date is unknown/missing should be included or not.
- The options to *filter samples by location* and *filter samples by region* allow for subsets by location (country) and/or region within a location. Also for these options there is the possibility to include/exclude samples for which the location/region is unknown/missing.
- The option to *filter samples by production method* allows for subset selection on production method of the sampled product (e.g., organic or conventional).

• The option to filter samples by additional sample properties available in the data (see *data format*).

Sample subset settings	Save Changes
✓ Filter samples by year	6
Sample years 2018	- 0
Filter samples by month	0
Include samples with unspecified sampling date	0
Filter samples by location	0
Filter samples by region	0
Filter samples by production method	0
✓ Filter samples by additional sample properties	0
Filter samples by importseason	
✓ Filter samples by season	
Selected values * Winter	•

Figure 3.28: Sample subset selection form of the front end. This form is a sub-form of the concentrations module panel.

Concentrations data formats

Concentrations data are analytical measurements of chemical substances occurring in food samples. The following data formats are supported/available for providing concentration data in MCRA:

- 1. **Relational concentration data format:** Concentration data is provided using a number of relational tables describing (*food*) samples (e.g., sampling date and location), the *analytical methods*, with their properties for substances (e.g., LOQ and LOD), sample analyses, describing the analyses performed on the samples (e.g., analysis date and the used analytical method), and *concentration measurements* (e.g., the measurement type and value). This relational table structure is also used internally by MCRA.
- 2. EFSA Standard Sample Description (SSD) format: Concentration data according to the EFSA Standard Sample Description (SSD) format. During upload, *SSD data* are converted automatically to the relational data format used internally by MCRA.
- 3. **Tabulated data scheme:** This is a simplified data format where samples and analytical methods are not explicitly specified. *Tabulated concentration data* are converted automatically to the MCRA relational data format.

Relational concentration data format

The relational data format is the data format that is used internally in MCRA.

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Analytical methods

The analytical methods used for analysing the samples are recorded in the analytical methods table. Each analytical method should have a unique identification code (idAnalyticalMethod). The description field may be used for a more detailed description of the analytical method. The records of this table should be linked to one or more analytical method substance properties table, which record the substances that are measured by this method (and their limits of reporting).

Name	Туре	Description	Aliases	Required
idAnalytical- Method	AlphaNumeric (50)	The code for the method of analysis.	idAnalytical- Method, Analytical- MethodId, Analytical- MethodName, Id	Yes
Name	AlphaNumeric (100)	Name of the analytical method.	Name	No
Description	AlphaNumeric (255)	Additional description of method of analysis.	Description	No

Table 3.90:	Table	definition	for	Analytical	methods.
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Accepted table names: AnalyticalMethod, AnalyticalMethods.

Analytical method properties for substances

This table describes the substances analysed by an analytical method. For each substance analysed by an analytical method a record should be included that describes the unit of measurement and the reporting limits (LOQ/LOD).

Name	Туре	Description	Aliases	Required
idAnalytical- Method	AlphaNumeric (50)	The code of method of analysis.	idAnalytical- Method, Analytical- MethodName, Analytical- MethodId	Yes
idSubstance	AlphaNumeric (50)	The substance code.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
LOD	Numeric	The limit of detection (LOD) is the lowest concentration of an substance in a sample that can be consistently detected.	LOD	No
LOQ	Numeric	The limit of quantification (LOQ) is the lowest concentration of a substance that can be quantified. The LOQ should be larger than the LOD.	LOQ, LOR	No
Concentration- Unit	<i>ConcentrationUnit</i>	The unit used for reporting the LOD, LOQ, and the substance concentrations. When not specified, then a default unit of mg/kg is assumed.	Concentration- Unit, Units, Unit	No

Table 3.91: Table definition for Analytical method properties for substances.

 $\label{eq:accepted} Accepted \ table \ names: \ Analytical Method Substances, \ Analytical Method Substance, \ Analytical Method Compounds, \ Analytical Method Compound.$

Food samples

Food sample for analysis of concentrations. May be characterised by location and/or date of sampling. A sample can be analysed multiple times, the results per analysis are stored as analysis samples.

Name	Туре	Description	Aliases	Required
idFoodSample	AlphaNumeric (50)	The identification number of the food sample.	idFoodSample, idSample, SampleId, Id	Yes
idFood	AlphaNumeric (50)	The food code.	idFood, FoodId, Food, FoodCode	Yes
Location	AlphaNumeric (50)	The location or country code, sampling location.	Location, Location- Sampling, Sampling- Location, Country	No
Region	AlphaNumeric (50)	The area or region within the sampling location.	Region, Area, Sampling- Region, SamplingArea	No
DateSampling	DateTime	The date of sampling.	DateSampling, SamplingDate	No
Production- Method	AlphaNumeric (50)	Additional information on the type/method of production of the sampled food.	Production- Method, ProductionType	No
Name	AlphaNumeric (100)	Name of the food sample.	Name	No
Description	AlphaNumeric (200)	Additional description of the food sample.	Description	No

Table 3.92: Table definition for Food samples.

Accepted table names: FoodSamples, FoodSample, Samples, Sample, PrimarySamples.

Sample properties

Food sample properties, additional columns that can also be specified as additional columns in the food samples table

Name	Туре	Description	Aliases	Required
Name	AlphaNumeric (50)	The name of the property.	Id	Yes
Description	AlphaNumeric (200)	Additional description of the sample property.	Description	No

Table 3.93: Table definition for Sample properties.

Accepted table names: SampleProperties, SampleProperty.

Sample property values

Food sample property values, additional columns that can also be specified as additional columns in the food samples table

Name	Туре	Description	Aliases	Required
idSample	AlphaNumeric (50)	The identification number of the food sample.	Id, IdFoodSample	Yes
PropertyName	AlphaNumeric (50)	The name of the property.	IdProperty, Name	Yes
TextValue	AlphaNumeric (50)	The value of the property as text value.		No
DoubleValue	Numeric	The value of the property as number.		No

Accepted table names: SamplePropertyValues, SamplePropertyValue.

Sample Analyses

An analysis sample specifies the analysis of a sample by an analytical method. A sample can be analysed multiple times, the results per analysis are stored as analysis samples.

Name	Туре	Description	Aliases	Required
idSample- Analysis	AlphaNumeric (50)	The identification number of the analysed sample.	id, idSample- Analysis, SampleAnalysis, idAnalysis- Sample, AnalysisSample- Id	Yes
idFoodSample	AlphaNumeric (50)	The identification number of the food sample.	idFoodSample, idSample, SampleId, Sample	Yes
idAnalytical- Method	AlphaNumeric (50)	The code of method of analysis.	idAnalytical- Method, Analytical- MethodId	Yes
DateAnalysis	DateTime	The date of the analysis.	DateAnalysis, AnalysisDate, Date	No
Name	AlphaNumeric (100)	Name of the analysis sample.	Name	No
Description	AlphaNumeric (200)	Additional description of the the analysis sample.	Description	No

Table 3.9)5∙ Table	definition	for 9	Sample	Analyses
Table 5.7	5. Table	ucinition	101	Sampie	Analyses.

Accepted table names: AnalysisSamples, AnalysisSample, SampleAnalysis, SampleAnalyses.

Sample concentrations

This table contains substance concentration values specified in the unit defined by the analytical method. The analytical method contains the list of all substances which have been analyzed in the analysis sample. This ConcentrationsPerSample table contains the analysis results where substances with positive concentrations are included. Censored values (i.e. results 'less than LOQ or LOD') are reported as follows: 1) Substances for which an LOD (Limit of detection) is reported are included with ResType 'LOD', without a concentration value. 2) Substances for which only an LOQ (Limit of quantification) is reported are EXCLUDED, because the LOQ value from the analytical method substances table (AnalyticalMethodCompounds) is used by default. Explicitly missing concentration values are specified with ResType 'MV' (obligatory).

Name	Туре	Description	Aliases	Required
idSample- Analysis	AlphaNumeric (50)	The identification number of the analysed sample.	idSample- Analysis, SampleAnalysis, idAnalysis- Sample, AnalysisSample- Id	Yes
idSubstance	AlphaNumeric (50)	The substance code.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
Concentration	Numeric	The measured concentration.	Concentration	No
ResType	ResType	The type of residue. Should be VAL (= default), LOQ, LOD or MV.	ResType	No

 Table 3.96: Table definition for Sample concentrations.

Accepted table names: SampleConcentrations, ConcentrationsPerSample, ConcentrationPerSample.

SSD concentration data format

The Standard Sample Description (SSD) concentration is the standard data format proposed by EFSA. Optionally, additional sample properties may be specified of fields that are not part of the SSD format (e.g., season of sampling). For this, the sample properties table and the sample property values table can be used.

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SSD concentrations

MCRA uses the concept of samples analysed by analytical methods, where the analytical method contains the substances analysed and the LOQs and LODs for these substances. However, the SSD data do not provide information on the analytical methods at this level of detail. Therefore, the provided SSD sample records are used to generate analytical methods which are linked to the samples. All SSD records with the same labSampCode and labSubSampCode compose one MCRA analysis sample. All SSD samples that contain the same substance, LOQ/LOD values and resUnit combinations are linked to the corresponding generated analytical method. If both LOQ and LOD are provided, the LOQ is used as the LOR of the generated analytical method. It is highly recommended to supply LOQ/LOD values, even for positive measurements, because this reduces the number of gererated analytical methods.

Name	Туре	Description	Aliases	Required
labSampCode	AlphaNumeric (30)	Code of the laboratory sample. MCRA will use the combination of labSampCode and labSubSampCode as unique code for a sample.	labSampCode	Yes
labSubSamp- Code	AlphaNumeric (4)	Code of the laboratory sub-sample. MCRA will use the combination of labSampCode and labSubSampCode as unique code for a sample.	labSubSamp- Code	No
sampCountry	AlphaNumeric (2)	Two-letter code to identify the country of sampling.	sampCountry	No
sampArea	AlphaNumeric (5)	Area where the sample was collected.	sampArea	No
prodCode	AlphaNumeric (50)	Code identifying the modelled food. Should be equal to a code idFood in the Foods table.	prodCode	Yes
prodProdMeth	AlphaNumeric (50)	Code providing additional information on the type of production for the food under analysis.	prodProdMeth	No
sampStrategy	AlphaNumeric (5)	Sampling strategy code.	sampStrategy, progSamp- Strategy, sampling- Strategy	No
fieldTrialType	AlphaNumeric (50)	Field trial type code.	fieldTrialType	No
sampY	Integer (4)	Year of sampling.	sampY	No
sampM	Integer (2)	Month of sampling.	sampM	No
sampD	Integer (2)	Day of sampling.	sampD	No No
analysisM	Integer (2)	Month of analysis	analysisM	No
analysisD	Integer (2)	Day of analysis.	analysisD	No
paramCode	AlphaNumeric (50)	Code identifying the substance.	paramCode	Yes
resUnit	ConcentrationUnit	Unit of residue measurement.	resUnit	Yes
resLOD	Numeric	Residue Limit Of Detection. Required if resType is LOD. MCRA will use resLOD as LOR if resLOQ is not provided.	resLOD	No
resLOQ	Numeric	Residue Limit Of Quantification. Required if resType is LOQ. MCRA will use resLOQ as LOR if provided.	resLOQ	No
resVal resType	Numeric ResType	Required if resType is VAL. Type of residue data. Should be VAL, LOQ, LOD or MV.	resVal resType	No Yes

Table 3.97:	Table definition	n for SSD c	oncentrations.
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Accepted table names: ConcentrationsSSD, SSDConcentrations.

Sample properties

Food sample properties, additional columns that can also be specified as additional columns in the food samples table

Name	Туре	Description	Aliases	Required
Name Description	AlphaNumeric (50) AlphaNumeric (200)	The name of the property. Additional description of the sample property.	Id Description	Yes No

Table 3.98: Table definition for Sample properties.

Accepted table names: SampleProperties, SampleProperty.

Sample property values

Food sample property values, additional columns that can also be specified as additional columns in the food samples table

Name	Туре	Description	Aliases	Required
idSample	AlphaNumeric (50)	The identification number of the food sample.	Id, IdFoodSample	Yes
PropertyName	AlphaNumeric (50)	The name of the property.	IdProperty, Name	Yes
TextValue	AlphaNumeric (50)	The value of the property as text value.		No
DoubleValue	Numeric	The value of the property as number.		No

Table 3.99: Table definition for Sample property values.

Accepted table names: SamplePropertyValues, SamplePropertyValue.

Tabulated concentration data format

The tabulated concentration data format is an old data format for entering concentration data.

Download empty dataset template: Zipped CSV Excel

Tabulated concentrations

In the tabulated concentrations data table, each record represents one or multiple samples, and each sample contains a concentration value for a food/substance combination. Censored values (i.e. concentrations less than LOR) are specified as negative values, i.e. 'less than LOR' should be specified as minus the LOR value. MCRA uses the concept of samples analysed by analytical methods, where the analytical method is characterised by the substances analysed and the LORs for these substances. However, the tabulated data do not provide this information explicitly. Samples are reconstructed from the tabulated records using the NumberOfSamples field to create that number of single substance samples. Analytical methods are reconstructed from the data, with each analytical method having only one analysed substance with a LOR and concentration unit. When a negative concentration value is given (i.e., it is a censored measurement), this value is recorded as the LOR (negated). All censored measurements of the same substance with the same LOR and concentration unit are linked to the same analytical method. When a positive concentration value is given, this value is recorded as the measured concentration of the sample. All positive measurements of the same substance are linked to the same analytical method. When a positive than the lowest positive concentration. When a concentration of 0 (zero) is given, the measurement is considered to be a censored measurement and the LOR is set to the default value 1E-08.

Name	Туре	Description	Aliases	Required
GUID	AlphaNumeric (50)	Unique identifier of the analysis sample of this tabulated concentration record.	idAnalysis- Sample, SampleId, SampleCode, Code, Id	No
idSubstance	AlphaNumeric (50)	The code of the substance of this concentration value.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
idFood	AlphaNumeric (50)	The food code.	idFood, FoodId, FoodMeasured, Food	Yes
DateSampling	AlphaNumeric (10)	The date of sampling.	DateSampling	No
SamplingType	AlphaNumeric (50)	The type of sampling (monitoring).	SamplingType	No
Location	AlphaNumeric (50)	The location or country of sampling.	Location, Country	No
NumberOf- Samples	Integer	The count of the number of times the specified concentration or limit of reporting (LOR) occurs.	NumberOf- Samples	Yes
Concentration	Numeric	The concentration or LOR. LORs are specified using a minus (-) sign.	Concentration, Value	Yes
Concentration- Unit	ConcentrationUnit	The unit of the specified concentrations/LORs (default mg/kg).	Concentration- Unit, Unit	No

Table 3.100: T	Table definition	for Tabulated	concentrations.
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Accepted table names: ConcentrationTabulated, ConcentrationValues, TabulatedConcentrations, TabulatedConcentration.

Concentrations settings

Selection settings

Table 3.101: Selection settings for module Concentrations.

Name	Туре	Description
Seed for pseudo-random number generator	Numeric	A value of 0 will use a pseudo-random seed in each run, a valu 0 will provide the same results in a repeated run.
Selected tier	SettingsTemplateType	Specifies all module settings should be set according to a pre-defined tier or using custom settings.
Filter samples exceeding the concentration limits	Boolean	If checked, samples with at least one substance concentration higher than some factor (concentration limit filter exceedance factor) times the MRL are filtered out.
Concentration limit filter exceedance factor	Numeric	The multiplication factor for the concentration limit exceedance filter.
Use substance conversion rules	Boolean	If checked, concentrations are modelled in terms of active substances (using substance conversion).
Substance conversion method	SubstanceTranslationAlloca- tionMethod	Allocation method for assigning active substance concentration from measured substance concentrations based on substance translations.

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Name	Туре	Description
Retain all allocated substances after active substance allocation	Boolean	If checked, all allocated substances kept after substance conversion. Otherwise, the concentration data is restricted to the active substances of the assessment group
Account for substance authorisations in substance conversions	Boolean	Account for substance authorisations when allocating measured substances to active substance using substance conversions.
Fix duplicate substance allocation inconsistencies	Boolean	Resolve inconsistencies when active substance allocation leads to multiple concentration value estimates for the same active substance. This method uses the mean of the positives or zero concentrations when available, or else the lowest of the censore values.
Use extrapolation rules	Boolean	Use extrapolation rules to extrapolate food samples for foods w a limited amount of samples (data poor foods) from other food (data rich foods).
Threshold for extrapolation	Numeric	Threshold for extrapolation.
Restrict extrapolations to equal MRLs	Boolean	Restrict extrapolations to equal MRLs.
Restrict extrapolations to authorised uses	Boolean	Only extrapolate if substance use is authorised.
Impute water concentrations	Boolean	Impute constant concentration values on the selected (water) commodity.
Water commodity	AlphaNumeric	The commodity for which constant concentration values should added.
Water concentration value $(\mu g/kg)$	Numeric	Constant concentration value that should be used for water (in $\mu g/kg$).
Restrict water imputation to the five most toxic substances	Boolean	Restrict water imputation to the five most toxic substances.
Restrict water imputation to authorised uses	Boolean	Restrict water imputation to authorised uses.
Restrict water imputation to approved substances	Boolean	Specifies whether imputation of water should be limited to approved substances only.
Include focal commodity concentrations	Boolean	Specifies whether there is monitoring data that should replace p of the consumption data for the specified focal commodities.
substance combinations	rocali oou	are to be replaced by focal commodity concentrations
Focal commodity substances	AlphaNumeric	The substances for which background concentration data are to replaced by focal commodity concentrations.
Focal commodity concentrations replacement method	FocalCommodityReplacement- Method	Replacement method to be used for replacing base concentration data with concentration data of the focal commodity/commodition concentrations.
Focal commodity substance occurrence percentage	Numeric	Anticipated occurrence percentage / agricultural use percentage of the focal commodity.
Adjustment factor for the focal food/substance concentration	Numeric	Optional adjustment factor for the focal food/substance concentration. E.g., the expected ratio of mean monitoring concentration and mean field trial concentration.
Use deterministic substance conversions for focal commodity	Boolean	Convert measured substance concentrations of focal commodity to active substance concentrations using deterministic substance conversion factors.
Filter samples by specific property values (subset selection)	Boolean	Specifies whether a subset selection on specific sample properti should be made (e.g., by country or by year).
Restrict to specific modelled foods (modelled foods subset)	Boolean	If checked, then the assessment is restricted to the specified modelled foods.
Align sampling location subset with population	Boolean	If checked, the samples are filtered based on the location of the selected population.

Table 3.101 - continued from previous page

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Name	Туре	Description
Include samples with unspecified location	Boolean	If checked, then samples for which the sample location is not specified are also included by the sample location filter.
Align sample date subset with population	Boolean	If checked, the samples are filtered based on the period of the selected population.
Align sampling month subset with population	Boolean	If checked, the samples are filtered based on the month/period the selected population.
Include samples with unspecified sampling date	Boolean	If checked, then samples for which the sample date is not specified are also included by the sample date filter.
Filter samples by location	Boolean	If checked, samples are filtered based on the selected locations.
Filter samples by year	Boolean	If checked, samples are filtered based on the selected years.
Filter samples by month	Boolean	If checked, samples are filtered based on the specified sampling month.
Samples subset definitions	SamplesSubsetDefinition	Samples subset definitions filter the concentration samples on custom properties using a set of keywords.
Samples by location subset definition	LocationSubsetDefinition	The subset definition to filter samples by location.
Samples by time period subset definition	PeriodSubsetDefinition	The subset definition to filter samples by time period.
Filter processed focal commodity samples	Boolean	Filter (exclude) concentration data at the level of raw primary commodity (RPC) derivatives (processed RPC) for the focal for
Focal commodity proposed concentration limit (mg/kg)	Numeric	Proposed concentration value that should be used for the focal commodity substance combination (in mg/kg).
Include processed derivatives of the focal food in measurement replacement	Boolean	If checked, measurement replacement is also performed for processed derivatives of the focal food (e.g. for juiced oranges)

Table 3.101 - continued from previous page

Output settings

Name	Туре	Description
Lower percentage for variability (%)	Numeric	The default value of 25% may be overruled.
Upper percentage for variability (%)	Numeric	The default value of 75% may be overruled.

Uncertainty settings

	······	
Name	Туре	Description
Resample concentrations	Boolean	Specifies whether concentrations are resampled by empirical bootstrap or using a parametric uncertainty model.

Table 3.103: Uncertainty settings for module Concentrations.

Concentrations tiers

In addition to the possibility for users to work with their own choices for all settings, MCRA implements Tier 1 and 2 from the European Commission working document SANTE-2015-10216 rev. 7 (2018) on risk management aspects related to the assessment of cumulative exposure.

Overview

	N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFSA 2022 Acute Tier II	EFS 202 Chro Tier
	F te sa pl ez ce ir th co tr ti lit	false	false	false	false	false	false	false	false	false	false
	U st co vo si rt	true	true	true	true			true	true	true	true
	S st co vo si m	UseMost- Toxic	UseMost- Toxic	DrawRan- dom	DrawRan- dom			UseMost- Toxic	UseMost- Toxic	DrawRan- dom	Drav dom
	R ta al al lc ca st af te ad ti st al lc ca ti ti	true	true	true	true			true	true	true	true
	A co fc su	false	false	true	true			false	false	true	true
•	188 al th ri	;						Chapte	r 3. Module	S	

Table 3.104: Tier overview for module Concentrations.

Retrospective dietary CRA (EC 2018) - Acute / Tier I

Table 3.105: Tier definition for Retrospective dietary CRA (EC 2018) - Acute / Tier I.

Name	Setting
Filter samples exceeding the concentration limits	false
Use substance conversion rules	true
Substance conversion method	UseMostToxic
Retain all allocated substances after active substance allocation	true
Account for substance authorisations in substance conversions	false
Fix duplicate substance allocation inconsistencies	false
Use extrapolation rules	true
Threshold for extrapolation	10
Restrict extrapolations to equal MRLs	true
Restrict extrapolations to authorised uses	true
Impute water concentrations	true
Water concentration value (µg/kg)	0.1
Restrict water imputation to the five most toxic substances	true
Restrict water imputation to authorised uses	false
Restrict water imputation to approved substances	false

Retrospective dietary CRA (EC 2018) - Chronic / Tier I

Table 3.106: Tier definition for Retrospective dietary CRA (EC 2018) - Chronic / Tier I.

Name	Setting
Filter samples exceeding the concentration limits	false
Use substance conversion rules	true
Substance conversion method	UseMostToxic
Retain all allocated substances after active substance allocation	true
Account for substance authorisations in substance conversions	false
Fix duplicate substance allocation inconsistencies	false
Use extrapolation rules	true
Threshold for extrapolation	10
Restrict extrapolations to equal MRLs	true
Restrict extrapolations to authorised uses	true
Impute water concentrations	true
Water concentration value (µg/kg)	0.1
Restrict water imputation to the five most toxic substances	true
Restrict water imputation to authorised uses	false
Restrict water imputation to approved substances	false

Retrospective dietary CRA (EC 2018) - Acute / Tier II

Table	3.107:	Tier	definition	for	Retrospective	dietary	CRA	(EC	2018)	-
Acute	e / Tier I	I.								

Name	Setting		
Filter samples exceeding the concentration limits	false		
Use substance conversion rules	true		
Substance conversion method	DrawRandom		
Retain all allocated substances after active substance allocation	true		
Account for substance authorisations in substance conversions	true		
Fix duplicate substance allocation inconsistencies	false		
Use extrapolation rules	true		
Threshold for extrapolation	10		
Restrict extrapolations to equal MRLs	true		
Restrict extrapolations to authorised uses	true		
Impute water concentrations	true		
Water concentration value (µg/kg)	0.05		
Restrict water imputation to the five most toxic substances	true		
Restrict water imputation to authorised uses	false		
Restrict water imputation to approved substances	false		

Retrospective dietary CRA (EC 2018) - Chronic / Tier II

Table 3.108: Tier definition for Retrospective dietary CRA (EC 2018) - Chronic / Tier II.

Name	Setting
Filter samples exceeding the concentration limits	false
Use substance conversion rules	true
Substance conversion method	DrawRandom
Retain all allocated substances after active substance allocation	true
Account for substance authorisations in substance conversions	true
Fix duplicate substance allocation inconsistencies	false
Use extrapolation rules	true
Threshold for extrapolation	10
Restrict extrapolations to equal MRLs	true
Restrict extrapolations to authorised uses	true
Impute water concentrations	true
Water concentration value (µg/kg)	0.05
Restrict water imputation to the five most toxic substances	true
Restrict water imputation to authorised uses	false
Restrict water imputation to approved substances	false

Retrospective dietary CRA (EFSA 2012) - Acute / Pessimistic

Acute probabilistic exposure assessment using the pessimistic model settings according to the EFSA Guidance 2012. Only processing factors > 1 are applied. For unit variability, the Beta distribution is applied.

Table 3.109: Tier definition for Retrospective dietary CRA (EFSA 2012) - Acute / Pessimistic.

Name	Setting
Filter samples exceeding the concentration limits	false

Retrospective dietary CRA (EFSA 2012) - Chronic / Pessimistic

Chronic probabilistic exposure assessment using the pessimistic model settings according to the EFSA Guidance 2012. Only processing factors > 1 are applied.

Table 3.110: Tier definition for Retrospective dietary CRA (EFSA 2012) - Chronic / Pessimistic.

Name	Setting
Filter samples exceeding the concentration limits	false

Retrospective dietary CRA (EFSA 2022) - Acute / Tier I

Table 3.111: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier I.

Name	Setting
Filter samples exceeding the concentration limits	false
Use substance conversion rules	true
Substance conversion method	UseMostToxic
Retain all allocated substances after active substance allocation	true
Account for substance authorisations in substance conversions	false
Fix duplicate substance allocation inconsistencies	false
Use extrapolation rules	false
Impute water concentrations	true
Water concentration value (µg/kg)	0.1
Restrict water imputation to the five most toxic substances	true
Restrict water imputation to authorised uses	false
Restrict water imputation to approved substances	true

Retrospective dietary CRA (EFSA 2022) - Chronic / Tier I

Table 3.112: Tier definition for Retrospective dietary CRA (EFSA 2022)

⁻ Chronic / Tier I.

Name	Setting		
Filter samples exceeding the concentration limits	false		
Use substance conversion rules	true		
Substance conversion method	UseMostToxic		
Retain all allocated substances after active substance allocation	true		
Account for substance authorisations in substance conversions	false		
Fix duplicate substance allocation inconsistencies	false		
Use extrapolation rules	false		
Impute water concentrations	true		
Water concentration value (µg/kg)	0.1		
Restrict water imputation to the five most toxic substances	true		
Restrict water imputation to authorised uses	false		
Restrict water imputation to approved substances	true		

Retrospective dietary CRA (EFSA 2022) - Acute / Tier II

Table 3.113: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier II.

Name	Setting
Filter samples exceeding the concentration limits	false
Use substance conversion rules	true
Substance conversion method	DrawRandom
Retain all allocated substances after active substance allocation	true
Account for substance authorisations in substance conversions	true
Fix duplicate substance allocation inconsistencies	false
Use extrapolation rules	false
Impute water concentrations	true
Water concentration value (µg/kg)	0.05
Restrict water imputation to the five most toxic substances	true
Restrict water imputation to authorised uses	false
Restrict water imputation to approved substances	true

Retrospective dietary CRA (EFSA 2022) - Chronic / Tier II

Table 3.114: Tier definition for Retrospective dietary CRA (EFSA 2022)

⁻ Chronic / Tier II.

Name	Setting
Filter samples exceeding the concentration limits	false
Use substance conversion rules	true
Substance conversion method	DrawRandom
Retain all allocated substances after active substance allocation	true
Account for substance authorisations in substance conversions	true
Fix duplicate substance allocation inconsistencies	false
Use extrapolation rules	false
Impute water concentrations	true
Water concentration value (µg/kg)	0.05
Restrict water imputation to the five most toxic substances	true
Restrict water imputation to authorised uses	false
Restrict water imputation to approved substances	true

Prospective dietary CRA (EFSA 2023) - Acute / Tier II

Table 3.115: Tier definition for Prospective	dietary	CRA	(EFSA	2023)	-
Acute / Tier II.					

Name	Setting
Filter samples exceeding the concentration limits	true
Concentration limit filter exceedance factor	2
Use substance conversion rules	true
Substance conversion method	DrawRandom
Retain all allocated substances after active substance allocation	true
Account for substance authorisations in substance conversions	true
Fix duplicate substance allocation inconsistencies	false
Use extrapolation rules	false
Impute water concentrations	true
Water concentration value (µg/kg)	0.05
Restrict water imputation to the five most toxic substances	true
Restrict water imputation to authorised uses	false
Restrict water imputation to approved substances	true
Include focal commodity concentrations	true
Focal commodity substance occurrence percentage	20
Adjustment factor for the focal food/substance concentration	1
Use deterministic substance conversions for focal commodity	true

Prospective dietary CRA (EFSA 2023) - Chronic / Tier II

Table 3.116: Tier definition for Prospective dietary CRA (EFSA 2023) - Chronic / Tier II.

Name	Setting
Filter samples exceeding the concentration limits	true
Concentration limit filter exceedance factor	2
Use substance conversion rules	true
Substance conversion method	DrawRandom
Retain all allocated substances after active substance allocation	true
Account for substance authorisations in substance conversions	true
Fix duplicate substance allocation inconsistencies	false
Use extrapolation rules	false
Impute water concentrations	true
Water concentration value (µg/kg)	0.05
Restrict water imputation to the five most toxic substances	true
Restrict water imputation to authorised uses	false
Restrict water imputation to approved substances	true
Include focal commodity concentrations	true
Focal commodity substance occurrence percentage	20
Adjustment factor for the focal food/substance concentration	1
Use deterministic substance conversions for focal commodity	true

Prospective dietary CRA (EFSA 2024) - Acute / Tier II

Table 3.117: Tier definition for	Prospective	dietary	CRA	(EFSA	2024)	-
Acute / Tier II.						

Name	Setting
Filter samples exceeding the concentration limits	true
Concentration limit filter exceedance factor	2
Use substance conversion rules	true
Substance conversion method	DrawRandom
Retain all allocated substances after active substance allocation	true
Account for substance authorisations in substance conversions	true
Fix duplicate substance allocation inconsistencies	false
Use extrapolation rules	false
Impute water concentrations	true
Water concentration value (µg/kg)	0.05
Restrict water imputation to the five most toxic substances	true
Restrict water imputation to authorised uses	false
Restrict water imputation to approved substances	true
Include focal commodity concentrations	true
Focal commodity substance occurrence percentage	20
Adjustment factor for the focal food/substance concentration	1
Use deterministic substance conversions for focal commodity	true
Include processed derivatives of the focal food in measurement replacement	true

Prospective dietary CRA (EFSA 2024) - Chronic / Tier II

Name	Setting
Filter samples exceeding the concentration limits	true
Concentration limit filter exceedance factor	2
Use substance conversion rules	true
Substance conversion method	DrawRandom
Retain all allocated substances after active substance allocation	true
Account for substance authorisations in substance conversions	true
Fix duplicate substance allocation inconsistencies	false
Use extrapolation rules	false
Impute water concentrations	true
Water concentration value (µg/kg)	0.05
Restrict water imputation to the five most toxic substances	true
Restrict water imputation to authorised uses	false
Restrict water imputation to approved substances	true
Include focal commodity concentrations	true
Focal commodity substance occurrence percentage	20
Adjustment factor for the focal food/substance concentration	1
Use deterministic substance conversions for focal commodity	true
Include processed derivatives of the focal food in measurement replacement	true

Table 3.118: Tier definition for Prospective dietary CRA (EFSA 2024) - Chronic / Tier II.

Concentrations uncertainty

Uncertainty due to a limited number of samples can be accounted for by resampling/bootstrapping. Resampling is done on a sample-based basis preserving co-occurrence of substance residue values on the same sample for multiple-substance analyses.

Concentrations as data

Concentration data can be entered using the internal, relational data format or using the EFSA SSD format. Depending on the settings, the entered concentration data can be pre-processed for conversion to active substances, extrapolation to other foods, and/or default values can be added for water.

- Concentrations data formats
- Concentrations from data
- Concentrations calculation

Inputs used: Focal food concentrations Food extrapolations Substance conversions Deterministic substance conversion factors Relative potency factors Substance authorisations Active substances Concentration limits Substance approvals Processing factors

3.3.5 Deterministic substance conversion factors

Deterministic substance conversion factors are (possibly food specific) conversion factors for converting measured substance concentrations to equivalent active substance concentrations. They are used in the *concentrations module* for translation of foreground/focal commodity substance concentrations to active substance concentrations (see option *use deterministic substance conversions for focal commodity* and the section on *focal commodity scenario analyses*).

Data module for deterministic substance conversion factors. These are (possibly food specific) conversion factors for expressing measured substance concentrations to equivalent active substance concentrations.

This module has as primary entities: Substances Foods

Output of this module is used by: Concentrations Single value concentrations

Deterministic substance conversion factors from data

Deterministic substance conversion factors data formats

Deterministic substance conversion factors are (possibly food specific) conversion factors for expressing measured substance concentrations to equivalent active substance concentrations.

Download empty dataset template: Zipped CSV Excel

Deterministic substance conversion factors

Deterministic substance conversion factors for translating measured substance concentrations to active substance concentrations.

Name	Туре	Description	Aliases	Required
idMeasured- Substance	AlphaNumeric (50)	Substance code of the measured substance.	idMeasured- Substance, idResidue- Definition, Residue- Definition, Measured- Substance, ParamCode- Measured- Substance	Yes
idActive- Substance	AlphaNumeric (50)	Substance code of the active substance.	idActive- Substance, idSubstance, Active- Substance, Substance, ParamCode- ActiveSubstance	Yes
idFood	AlphaNumeric (50)	The unique identification code of the food.	idFood, Code, FoodId, FoodCode, Food	No
Conversion- Factor	Numeric	Specifies the conversion factor to translate concentrations of the measured substance to (equivalent) concentrations of the active substance according to e.g. the system used in PRIMo.	Factor, Conversion- Factor	Yes
Reference	AlphaNumeric (200)	Reference to the source from which this value is obtained.	Reference, References, Source, Sources	No

Table 3.119: Table definition for Deterministic substance conversion factors.

Accepted table names: SingleValueSubstanceConversionFactors, SingleValueConversionFactors, SingleValueConversions, SubstanceConversionsFixed, DeterministicSubstanceConversionFactors, DeterministicConversionFactors.

Deterministic substance conversion factors as data

Deterministic substance conversion factors are (possibly food specific) conversion factors for expressing measured substance concentrations to equivalent active substance concentrations.

- Deterministic substance conversion factors data formats
- Deterministic substance conversion factors from data

3.3.6 Dust concentration distributions

Dust concentration distributions data are analytical measurements of chemical substances occurring in dust samples. In their simplest form, concentration data can just be used as provided by datasets.

This module has as primary entities: Substances

Output of this module is used by: Dust exposures

Dust concentration distributions from data Dust concentration distributions data formats

Substance concentration distributions in dust. Download empty dataset template: Zipped CSV Excel

Dust concentration distributions

Substance concentrations in dust.

Name	Туре	Description	Aliases	Required
idSample	AlphaNumeric (50)	The identification number of the dust sample.	idSample, SampleId, idDustSample	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
Concentration	Numeric	The substance concentration value of the dust sample.	Concentration, Value	Yes
Concentration- Unit	ConcentrationUnit	The unit for the substance concentrations. When not specified, then a default unit of ug/g is assumed.	Unit, Concentration- Unit	No

Table 3.120:	Table	definition	for Dust	concentration	distributions.
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Accepted table names: DustConcentrationDistributions, ConcentrationDistributionsDust.

Dust concentration distributions as data

Dust concentration distributions data can be entered using the internal, relational data format.

- Dust concentration distributions data formats
- Dust concentration distributions from data

3.3.7 Soil concentration distributions

Soil concentration distributions data are analytical measurements of chemical substances occurring in soil samples. In their simplest form, concentration data can just be used as provided by datasets.

This module has as primary entities: *Substances*

Output of this module is used by: Soil exposures

Soil concentration distributions from data

Soil concentration distributions data formats

Substance concentration distributions in soil.

Download empty dataset template: Zipped CSV Excel

Soil concentration distributions

Substance concentrations in soil.

Name	Туре	Description	Aliases	Required
idSample	AlphaNumeric (50)	The identification number of the soil sample.	idSample, SampleId, idSoilSample	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
Concentration	Numeric	The substance concentration value of the soil sample.	Concentration, Value	Yes
Concentration- Unit	ConcentrationUnit	The unit for the substance concentrations. When not specified, then a default unit of ug/g is assumed.	Unit, Concentration- Unit	No

Table 3.121: Table definition for Soil	l concentration distributions.
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Accepted table names: SoilConcentrationDistributions, ConcentrationDistributionsSoil.

Soil concentration distributions as data

Soil concentration distributions data can be entered using the internal, relational data format.

- Soil concentration distributions data formats
- Soil concentration distributions from data

3.3.8 Consumer product concentrations

Consumer product concentration (distributions) data are analytical measurements of chemical substances occurring in PCPs. In their simplest form, concentration data can just be used as provided by datasets.

This module has as primary entities: Substances Consumer products

Output of this module is used by: *Consumer product exposures*

Consumer product concentrations from data

Consumer product concentrations data formats

Consumer products concentrations. Download empty dataset template: Zipped CSV Excel

Consumer product concentrations

Consumer product concentrations.

Name	Туре	Description	Aliases	Required
idProduct	AlphaNumeric (50)	The consumer product.	idProduct, ProductId	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
Concentration	Numeric	The substance concentration value of the consumer product.	Concentration, Value	Yes
Unit	<i>ConcentrationUnit</i>	The unit for the substance concentrations. When not specified, then a default unit of ug/kg is assumed.	Unit, Concentration- Unit	No
Distribution- Type	ConsumerProduct- Concentration- DistributionType	Distribution type (Uniform, or LogNormal). If not specified, the concentration is assumed to be a constant factor.	Distribution- Type, Distribution	No
CvVariability	Numeric	Coefficient of variation. CvVariability CV		No
Occurrence- Percentage	Numeric	Percentage (%) of consumer products with substance (default 100%).	Occurrence- Percentage, Percentage	No

Table 3.122: Table definition for Consumer product concentrations.

Accepted table names: ConsumerProductConcentrations, CPConcentrations.

Consumer product concentrations as data

Consumer product concentration (distributions) data can be entered using the internal, relational data format.

- Consumer product concentrations data formats
- Consumer product concentrations from data

3.3.9 Indoor air concentrations

Indoor air concentrations data are analytical measurements of chemical substances occurring in indoor air samples. In their simplest form, concentration data can just be used as provided by datasets.

This module has as primary entities: Substances

Output of this module is used by: Air exposures

Indoor air concentrations from data

Indoor air concentrations data formats

Substance concentration in indoor air.

Download empty dataset template: Zipped CSV Excel

Indoor air concentrations

Substance concentrations in indoor air.

Name	Туре	Description	Aliases	Required
idSample	AlphaNumeric (50)	The identification number of the indoor air sample.	idSample, SampleId, idAirSample	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
Location	AlphaNumeric (50)	The location or country code.	Country, Location	No
Concentration	Numeric	The substance concentration value of the indoor air sample.	Concentration, Value	Yes
Unit	AirConcentration- Unit	The unit for the substance concentrations. When not specified, then a default unit of ug/m3 is assumed.	Unit, Concentration- Unit, Air- Concentration- Unit	No

Table 3.123:	Table	definition	for	Indoor	air	concentrations.
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Accepted table names: IndoorAirConcentrations, IndoorConcentrationsAir.

Indoor air concentrations as data

Indoor air concentrations data can be entered using the internal, relational data format.

- Indoor air concentrations data formats
- Indoor air concentrations from data

3.3.10 Outdoor air concentrations

Outdoor air concentrations data are analytical measurements of chemical substances occurring in outdoor air samples. In their simplest form, concentration data can just be used as provided by datasets.

This module has as primary entities: Substances

Output of this module is used by: Air exposures

Outdoor air concentrations from data

Outdoor air concentrations data formats

Substance concentration in outdoor air.

Download empty dataset template: Zipped CSV Excel
Outdoor air concentrations

Substance concentrations in outdoor air.

Name	Туре	Description	Aliases	Required
idSample	AlphaNumeric (50)	The identification number of the outdoor air sample.	idSample, SampleId, idAirSample	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
Location	AlphaNumeric (50)	The location or country code.	Country, Location	No
Concentration	Numeric	The substance concentration value of the outdoor air sample.	Concentration, Value	Yes
Unit	AirConcentration- Unit	The unit for the substance concentrations. When not specified, then a default unit of ug/m3 is assumed.	Unit, Concentration- Unit, Air- Concentration- Unit	No

Accepted table names: OutdoorAirConcentrations, OutdoorConcentrationsAir.

Outdoor air concentrations as data

Outdoor air concentrations data can be entered using the internal, relational data format.

- Outdoor air concentrations data formats
- Outdoor air concentrations from data

3.3.11 Focal food concentrations

In some cases the attention in an assessment is to evaluate concentrations (e.g., from specific field trials) for a specific food (and substance), in combination with a background of concentration data for other foods. Focal food concentrations can be included to provide these separate (foreground) concentration data for one or more focal food commodities that should replace measurements in the (background) *concentration data* in *focal commodity scenario analyses*.

This module has as primary entities: Foods Substances

Output of this module is used by: Concentrations

Focal food concentrations from data

Focal food concentrations data formats

Focal food concentrations settings

Selection settings

Table 3.12	25: Selection	settings for	module Focal	food	concentrations.

Name	Туре	Description
Focal commodity food / substance combinations	FocalFood	The foods / substances for which background concentration dat are to be replaced by focal commodity concentrations.

Calculation settings

Name	Туре	Description
Focal commodity concentrations replacement method	FocalCommodityReplacement- Method	Replacement method to be used for replacing base concentration data with concentration data of the focal commodity/commodit concentrations.
Focal commodity substances	AlphaNumeric	The substances for which background concentration data are to replaced by focal commodity concentrations.
Filter samples by specific property values (subset selection)	Boolean	Specifies whether a subset selection on specific sample properti should be made (e.g., by country or by year).
Samples subset definitions	SamplesSubsetDefinition	Samples subset definitions filter the concentration samples on custom properties using a set of keywords.

Table 3.126: Calculation settings for module Focal food concentrations.

Output settings

Name	Туре	Description
Lower percentage for variability (%)	Numeric	The default value of 25% may be overruled.
Upper percentage for variability (%)	Numeric	The default value of 75% may be overruled.

Focal food concentrations as data

Focal food concentrations are concentration data and specified in the exact same manner. The difference is that this data will be used to replace part of the concentration data in order to combine specific concentration data with a background of ordinary concentration data.

- Focal food concentrations data formats
- Focal food concentrations from data

Settings used

• Calculation Settings

3.3.12 Food extrapolations

Food extrapolations data specify which foods (data rich foods) can be used to impute concentration data for other foods with insufficient data (data poor foods).

This module has as primary entities: Foods

Output of this module is used by: Concentrations Food conversions

Food extrapolations from data

Food extrapolations data formats

Food extrapolations (or read-across food translations) can be used to specify whether data (e.g, occurrence data) on a food for which this is missing (a data poor food) may be extrapolated from another food for which data is available (read-across food).

Download empty dataset template: Zipped CSV Excel

Food extrapolations

Food extrapolations are simply specified as combinations of two food codes. One code for the food for the data poor food, and one for the data rich food (or read-across food).

Name	Туре	Description	Aliases	Required
DataPoorFood	AlphaNumeric (50)	The code of the data poor food. I.e., the food for which missing data is allowed to be extrapolated.	IdFoodData- Poor, FoodDataPoor, idFromFood, FromFoodId, FromFood, FoodFrom, Food, IdFood	Yes
CodeDataRich- Food	AlphaNumeric (50)	The code of the read-across food (or data rich food). I.e., the food from which data is used for extrapolation.	IdFoodData- Rich, FoodDataRich, IdFoodRead- Across, FoodRead- Across, IdReadAcross- Food, ReadAcross- Food, idToFood, ToFoodId, ToFoodIo,	Yes

Table 3.128: Table definition for Food extrapolation
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Accepted table names: ReadAcrossFoodTranslations, ReadAcrossFoodTranslation, ReadAcrossTranslations, ReadAcrossTranslation, FoodExtrapolations, FoodExtrapolation.

Food extrapolations as data

Food extrapolations are specified as data in the form of simple tuples of data rich food and data poor food for which extrapolation is allowed/reasonable.

- Food extrapolations data formats
- Food extrapolations from data

3.3.13 Modelled foods

Modelled foods are foods within the foods scope for which concentration data or MRLs of substances are available (or expected).

This module has as primary entities: Foods Substances

Output of this module is used by: Concentration models Food conversions

Modelled foods calculation

Modelled foods are the foods within the foods scope for which concentration data or MRLs of substances are available (or expected). Modelled foods are derived primarily from *concentration data*. That is, all foods for which food samples are available in the concentration data or MRL data are considered to be modelled foods. In addition, this set may be extended when *concentration limits* such as MRLs are available (see *calculation settings*) and/or when *food extrapolation rules* are used. Foods for which such data is available are considered to be modelled foods. The set of foods can also be restricted by omitting foods with only censored measurements (see *calculation settings*).

Modelled foods settings

Calculation settings

Name	Туре	Description
Multiple substances analysis	Boolean	Specifies whether the assessment involves multiple substances.
Restrict to specific modelled foods (modelled foods subset)	Boolean	If checked, then the assessment is restricted to the specified modelled foods.
Selected modelled foods	AlphaNumeric	Set of modelled foods that are of particular interest.
Derive modelled foods from concentrations	Boolean	Derive modelled foods from sample based concentration data.
Derive modelled foods from single value concentrations	Boolean	Derive modelled foods from single value concentrations.
Derive modelled foods from concentration limits	Boolean	Derive modelled foods from concentration limits.
Include foods with only censored value measurements	Boolean	Specifies whether foods with only censored value measurement are part of the exposure assessment (default yes).
Include substances with only censored value measurements	Boolean	Specifies whether substances with only censored value measurements are part of the exposure assessment (default yes)
Include substances without measurements	Boolean	Specifies whether substances without any measurements should included.

Table 3.129: Calculation settings for module Modelled foods.

Calculation of modelled foods

Modelled foods are computed from concentration data (which may also be in the form of single-value concentrations) and/or derived from available maximum residue limits.

• Modelled foods calculation

Inputs used: Concentrations Single value concentrations Concentration limits

Settings used

• Calculation Settings

3.3.14 Occurrence frequencies

Occurrence frequencies specify how often substances occur on foods. Frequencies are expressed as percentages.

This module has as primary entities: Foods Substances

Output of this module is used by: Concentration models Single value dietary exposures

Occurrence frequencies from data

Occurrence frequencies calculation

Occurrence frequencies can be provided as data or computed from *occurrence patterns*. For a food and substance, they are computed by collecting all occurrence patterns of this food and summing up the frequencies of the occurrence patterns containing the substance. In the unlikely case that the total frequency of the occurrence patterns of a food exceeds 100%, then a rescaling is applied first. If the sum of the frequencies does not sum up to 100%, the interpretation of the remaining unspecified percentage can be designated as either "no use" or "all use". In the the first case it is assumed that none of the substances occur on this remaining percentage. In the latter it is assumed that all of the substances occur on this remaining percentage. This choice is available as the setting *associate the unspecified percentage with no-occurrence for foods with at least one specified occurrence pattern*.

Depending on the setting *apply occurrence pattern percentages*, occurrence frequencies can be computed in a crisp form in which the occurrence frequency is either 0% or 100% or as percentages ranging from 0% to 100%.

Occurrence frequencies data formats

Occurrence frequencies are described by one simple table, specifying for pairs of food and substance, the associated occurrence frequencies as percentages.

Download empty dataset template: Zipped CSV Excel

Occurrence frequencies

Occurrence frequencies are specified as percentages for pairs of food and substance. Optionally, a reference can be included in each record to specify the source (e.g., from literature) from which the percentage was obtained.

Name	Туре	Description	Aliases	Required
idFood	AlphaNumeric (50)	The food code.	idFood, CodeFood, FoodId, FoodCode, Food	Yes
idSubstance	AlphaNumeric (50)	Code of the substance.	idSubstance, CodeSubstance, SubstanceId, SubstanceCode, Substance	Yes
Percentage	Numeric	The occurrence frequency percentage.	Percentage, Frequency- Percentage	Yes
Reference	AlphaNumeric (200)	Reference to the source from which this use frequency value is obtained.	Reference, References, Source, Sources	No

Table 3.130: Table definition for Occurrence frequencies.

Accepted table names: OccurrenceFrequencies.

Occurrence frequencies Settings

Selection settings

Name	Туре	Description
Associate the unspecified percentage with no-occurrence for foods with at least one specified occurrence pattern	Boolean	If checked, for foods with at least one specified occurrence pattern, unspecified occurrence patterns for the same food are assumed to be associated with no use. If unchecked, all substances are considered to be authorised (potentially present samples). Note that this setting cannot be used for foods that h no specified AUs. These foods have 100% potential presence of all substances. To declare all AUs on such a food un-authorised include an empty AU with percentage 100% in the AU data tal (i.e., use an AU for this food, without specifying substances in AU Substances table)
Apply occurrence pattern percentages	Boolean	If checked, use the percentages of potential presence as specifi by the occurrence patterns. If unchecked, 100% potential presence in samples is assumed for all substances identified by occurrence patterns.
Selected tier	SettingsTemplateType	Specifies all module settings should be set according to a pre-defined tier or using custom settings.

Table 3.131:	Selection	settings for	module	Occurrence	frequencies.
10010 5.151.	Selection	Settings for	module	Occurrence	inequencies.

Calculation settings

Fable 3.132:	Calculation	settings f	for module	Occurrence	frequencies.
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Name	Туре	Description
Multiple substances analysis	Boolean	Specifies whether the assessment involves multiple substances.

Uncertainty settings

Name	Туре	Description
Recompute occurrence patterns	Boolean	Specifies whether occurrence patterns should be recomputed in the uncertainty runs.

Occurrence frequencies as data

Occurrence frequencies are described by one table, specifying for a food and substance the associated occurrence frequency as percentage.

- Occurrence frequencies data formats
- Occurrence frequencies from data

Inputs used: Active substances

Settings used

• Calculation Settings

Calculation of occurrence frequencies

Occurrence frequencies for a food and substance are computed according to the model that is part of the EC 2018 Tier II definition (see van Klaveren et al. 2019)

• Occurrence frequencies calculation

Inputs used: Occurrence patterns

Settings used

• Calculation Settings

3.3.15 Occurrence patterns

Occurrence patterns (OPs) are the combinations (or mixtures) of substances that occur together on foods and the frequencies of these mixtures occurring per food, expressed in percentages. In the context of pesticides, occurrence patterns are associated with agricultural use percentages. Occurrence patterns are relevant to account for co-occurrence of active substances in exposed individuals. Occurrence patterns may be specified as data or modelled based on observed patterns of positive concentrations.

This module has as primary entities: Foods Substances

Output of this module is used by: Occurrence frequencies Dietary exposures

Occurrence patterns from data

Occurrence patterns calculation

Assumptions can be made for each food on the basis of findings in concentration data.

Tier 1: 0% occurrence is assumed for all substances with no positive concentrations at all; 100% occurrence is assumed for all substances with at least one positive concentration;

Tier 2: 0% occurrence is assumed for all substances with no positive concentrations at all; for substancefood combinations with at least one positive (finding), use findings patterns to implement a specific interpretation of Option 5 in the SANTE document, as described below.

Therefore in both tiers, substance-food combinations without any positive finding are handled in the optimistic way by assuming absolute zeroes for any censored observation.

If Tier 2 is selected, then for each of the modelled foods a tabulation is made of the observed frequencies of positives for all substance combinations (including the empty set), based on the *active substance concentrations*. For an OP consisting of just one substance, the basic frequency is the number of samples with a positive concentration divided by the number of samples where the substance has been measured (i.e., is not a MV). For an OP consisting of multiple substances, the basic frequency is the number of samples with all concentrations positive for the members divided by the number of samples where all members of the set have been measured.

After calculation of the basic frequencies for all occurrence patterns, these frequencies are rescaled such that the overall sum of frequencies is 100%. When *substance authorisations* are available, then patterns involving unauthorised substances are not rescaled and only those patterns for which all substances are authorised are rescaled such that the sum of all frequencies is 100%.

Note: the Tier 2 procedure is not what is literally written in the SANTE document, but is an interpretation agreed upon by EFSA and RIVM. An alternative model, not yet implemented, but perhaps more in line with the text of the SANTE document, would be to double the basic frequencies to modelled occurrence pattern frequencies. Only if the sum of all frequencies becomes larger than 100%, the set of frequencies would be normalised to 100% sum.

Occurrence patterns data formats

Agricultural use percentages for plant protection products (PPPs) may be of use for concentration modelling, as they provide information about what substance mixtures are expected to be present simultaneously on food samples. Especially for censored concentration measurements, this information may aid to determine whether the censored measurement originated from a true zero or may be a positive measurement below the limit of detection. Agricultural use percentages are specified using the agricultural uses and agricultural use substances table. This data format expects agricultural use percentages to be specified for mixtures of substances. Each mixture has an id (idAgriculturalUse) and a list of substances that are part of this mixture (agricultural use substances). These agricultural uses are assumed to be exclusive (i.e., only one mixture or PPP is used per sample). Hence, the sum of the agricultural uses for one food should not exceed 100%.

Download empty dataset template: Zipped CSV Excel

Agricultural uses

The AgriculturalUses contains the definitions of the agricultural use mixtures, or PPPs and the specification of the percentage of the products treated with this mixture. Optionally also the time period of the use percentage may be specified.

Name	Туре	Description	Aliases	Required
idAgricultural- Use	AlphaNumeric (50)	The unique identification code of the agricultural use group / plant protection product (PPP).	idAgricultural- Use, AgriculturalUse- Id, Id	Yes
idFood	AlphaNumeric (50)	The food code.	idFood, FoodId, Food	Yes
Location	AlphaNumeric (50)	The location or country code, agricultural use location.	Country, Location	No
StartDate	DateTime		StartDate	No
EndDate	DateTime		EndDate	No
Percentage- CropTreated	Numeric	The percentage agricultural use (%).	PercentageCrop- Treated, Percentage, PercCrop- Treated, PercentageUse	Yes
Name	AlphaNumeric (100)	Name of the agricultural use.	Name	No
Description	AlphaNumeric (200)	Additional description of the agricultural use.	Description	No

Table 3.134: Table definition for Agricultural uses.

Accepted table names: AgriculturalUses, AgriculturalUse.

Agricultural use substances

The agricultural use substances table records the substances that are part of the agricultural use mixtures (PPPs).

Name	Туре	Description	Aliases	Required
idAgricultural- Use	AlphaNumeric (50)	The agricultural use code, normally a code for a combination of authorised substances.	idAgricultural- Use, AgriculturalUse- Id	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes

Table 3.135: Table definition for	r Agricultural use substances.
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Accepted table names: AgriculturalUseHasSubstances, AgriculturalUseSubstances, AgriculturalUseGroups, AgriculturalUseGroup.

Occurrence patterns settings

Selection settings

Name	Туре	Description
Selected tier	SettingsTemplateType	Specifies all module settings should be set according to a pre-defined tier or using custom settings.
Associate the unspecified percentage with no-occurrence for foods with at least one specified occurrence pattern	Boolean	If checked, for foods with at least one specified occurrence pattern, unspecified occurrence patterns for the same food are assumed to be associated with no use. If unchecked, all substances are considered to be authorised (potentially present samples). Note that this setting cannot be used for foods that he no specified AUs. These foods have 100% potential presence of all substances. To declare all AUs on such a food un-authorised include an empty AU with percentage 100% in the AU data tak (i.e., use an AU for this food, without specifying substances in AU Substances table)
Apply occurrence pattern percentages	Boolean	If checked, use the percentages of potential presence as specific by the occurrence patterns. If unchecked, 100% potential presence in samples is assumed for all substances identified by occurrence patterns.
Scale up use frequency to 100%	Boolean	Scale up use frequency to 100%.
Restrict use percentage up-scaling to authorised uses	Boolean	Restrict use percentage up-scaling to authorised uses.

Table 3.136: Selection settings for module Occurrence patterns.

Calculation settings

Name	Туре	Description
Compute cumulative exposures	Boolean	Specifies whether the assessment involves multiple substances a results should be cumulated over all substances.
Multiple substances analysis	Boolean	Specifies whether the assessment involves multiple substances.

Uncertainty settings

Table 3.138:	Uncertainty	settings for	module	Occurrence	patterns.
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Name	Туре	Description
Recompute occurrence patterns	Boolean	Specifies whether occurrence patterns should be recomputed in the uncertainty runs.

Occurrence patterns tiers

Overview

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFSA 2022 Acute Tier II	EFSA 2022 Chronic Tier II	EFSA 2023 Acute Prospec- tive Tier II	EFS 2023 Chros Pros tive II
A pl ci re pi te pi ce	false	false	true	true	false	false	true	true	true	true
F te sa pl ez ca in th ca tr ti lii it	false	false	false	false	false	false	false	false	true	true
U st co vo si	true	true	true	true	true	true	true	true	true	true
Si st co ve si m	UseMost- Toxic	UseMost- Toxic	DrawRan- dom	DrawRan- dom	UseMost- Toxic	UseMost- Toxic	DrawRan- dom	DrawRan- dom	DrawRan- dom	Drav dom
R ta al lc ca st af te ac ti	true	true	true	true	true	true	true	true	true	true
3.3 . al lc ca	Occurren	ce modules	5					21	1	

Table 3.139: Tier overview for module Occurrence patterns.

Retrospective dietary CRA (EC 2018) - Acute / Tier I

	Name	Setting	From input tier	In module
	Apply occurrence pattern percentages Filter samples exceeding the	false false	Retrospec-	Concen-
	concentration limits		tive dietary CRA (EC 2018) - Acute / Tier I	tranons
	Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
	Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
	Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
	Account for substance authorisations in substance conversions	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
	Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
	Use extrapolation rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
	Threshold for extrapolation	10	Retrospec- tive	Concen- trations
3.3. Occurre	nce modules		CRA (EC 2018) - Acute /	

Tier I

Table 3.140: Tier definition for Retrospective dietary CRA (EC 2018) - Acute / Tier I.

Retrospective dietary CRA (EC 2018) - Chronic / Tier I

	Name	Setting	From	ln module
	Apply occurrence pattern paraenteese	false		modulo
	Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
	Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
	Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
	Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
	Account for substance authorisations in substance conversions	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
	Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
	Use extrapolation rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
	Threshold for extrapolation	10	Retrospec- tive	Concen- trations
3.3. Occurre	nce modules		CRA (EC 2018) - Chronic /	

Tier I

Table 3.141: Tier definition for Retrospective dietary CRA (EC 2018) - Chronic / Tier I.

Retrospective dietary CRA (EC 2018) - Acute / Tier II

	Name	Setting	From input tier	In module
	Apply occurrence pattern percentages Scale up use frequency to 100% Restrict use percentage up-scaling to authorised uses	true true true		
	Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
	Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
	Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
	Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
	Account for substance authorisations in substance conversions	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
	Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
	Use extrapolation rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
3.3. Occurre	nce modules Threshold for extrapolation	10	Retrospec- tive dietary	Concen- trations

CRA (EC

Table 3.142: Tier definition for Retrospective dietary CRA (EC 2018) - Acute / Tier II.

Retrospective dietary CRA (EC 2018) - Chronic / Tier II

	Name	Setting	From input tier	In module
	Apply occurrence pattern percentages Scale up use frequency to 100%	true		
	Restrict use percentage up-scaling to authorised uses	true		
	Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
	Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
	Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
	Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
	Account for substance authorisations in substance conversions	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
	Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
	Use extrapolation rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic /	Concen- trations
3.3. Occurre	nce modules Threshold for extrapolation	10	Retrospec- tive dietary	Concen- trations

CRA (EC

Table 3.143: Tier definition for Retrospective dietary CRA (EC 2018) - Chronic / Tier II.

Retrospective dietary CRA (EFSA 2022) - Acute / Tier I

	Name	Setting	From input tier	In module
	Apply occurrence pattern percentages Filter samples exceeding the concentration limits	false false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
	Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
	Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
	Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
	Account for substance authorisations in substance conversions	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
	Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
	Use extrapolation rules	false	Retrospec- tive dietary	Concen- trations
3.3. Occurre	nce modules		CRA (EFSA 2022) - Acute / Tier I	

Table 3.144: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier I.

Retrospective dietary CRA (EFSA 2022) - Chronic / Tier I

	Name	Setting	From input tier	In module
	Apply occurrence pattern percentages Filter samples exceeding the concentration limits	false false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
	Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
	Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
	Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
	Account for substance authorisations in substance conversions	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
	Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
	Use extrapolation rules	false	Retrospec- tive dietary	Concen- trations
3.3. Occurre	nce modules		CRA (EFSA 2022) - Chronic / Tier I	

Table 3.145: Tier definition for Retrospective dietary CRA (EFSA 2022) - Chronic / Tier I.

Retrospective dietary CRA (EFSA 2022) - Acute / Tier II

	Name	Setting	From input tier	In module
	Apply occurrence pattern percentages Scale up use frequency to 100% Restrict use percentage up-scaling to authorised uses	true true true		
	Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
	Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
	Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
	Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
	Account for substance authorisations in substance conversions	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
	Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
3.3. Occurr	Use extrapolation rules ence modules	false	Retrospec- tive dietary CRA	Concen- trations

(EFSA

Table 3.146: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier II.

Retrospective dietary CRA (EFSA 2022) - Chronic / Tier II

	Name	Setting	From input tier	In module
	Apply occurrence pattern percentages Scale up use frequency to 100% Restrict use percentage up-scaling to authorised uses	true true true		
	Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
	Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
	Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
	Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
	Account for substance authorisations in substance conversions	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
	Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
3.3. Occurre	Use extrapolation rules Use modules	false	Retrospec- tive dietary CRA	Concen- trations

(EF

Table 3.147: Tier definition for Retrospective dietary CRA (EFSA 2022) - Chronic / Tier II.

Prospective dietary CRA (EFSA 2023) - Acute / Tier II

	Name	Setting	From input tier	In module
	Apply occurrence pattern percentages Scale up use frequency to 100% Restrict use percentage up-scaling to authorised uses	true true true		
	Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
	Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
	Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
	Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
	Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
	Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
3.3. Occurre	Fix duplicate substance allocation nce modules inconsistencies	false	Prospec- tive dietary CRA	Concen- trations

(EFSA

Table 3.148: Tier definition for Prospective dietary CRA (EFSA 2023) - Acute / Tier II.

Prospective dietary CRA (EFSA 2023) - Chronic / Tier II

	Name	Setting	From input tier	In module
	Apply occurrence pattern percentages Scale up use frequency to 100% Restrict use percentage up-scaling to authorised uses	true true true		
	Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
	Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
	Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
	Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
	Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
	Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
3.3. Occurre	Fix duplicate substance allocation nce modules inconsistencies	false	Prospec- tive dietary CRA	Concen- trations

(EFS

Table 3.149: Tier definition for Prospective dietary CRA (EFSA 2023) - Chronic / Tier II.

233

Prospective dietary CRA (EFSA 2024) - Acute / Tier II

	Name	Setting	From input tier	In module
	Apply occurrence pattern percentages Scale up use frequency to 100% Restrict use percentage up-scaling to authorised uses	true true true		
	Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
	Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
	Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
	Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
	Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
	Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
3.3. Occurre	Fix duplicate substance allocation nce modules inconsistencies	false	Prospec- tive dietary CRA	Concen- trations

(EFSA

Table 3.150: Tier definition for Prospective dietary CRA (EFSA 2024) - Acute / Tier II.

Prospective dietary CRA (EFSA 2024) - Chronic / Tier II

	Name	Setting	From input tier	In module
	Apply occurrence pattern percentages Scale up use frequency to 100%	true true		
	Restrict use percentage up-scaling to authorised uses	true		
	Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
	Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
	Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
	Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
	Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
	Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
3.3. Occurre	Fix duplicate substance allocation nce modules inconsistencies	false	Prospec- tive dietary CRA	Concen- trations

(EFS

Table 3.151: Tier definition for Prospective dietary CRA (EFSA 2024) - Chronic / Tier II.

Occurrence patterns as data

Occurrence patterns are provided as data by specification of the occurrence mixtures and their associated occurrence/agricultural use percentages.

- Occurrence patterns data formats
- Occurrence patterns from data

Inputs used: Substance authorisations Active substances

Settings used

• Calculation Settings

Calculation of occurrence patterns

Occurrence patterns are computed from the observed patterns of positive concentrations in the concentration data.

Occurrence patterns calculation

Inputs used: Concentrations

Settings used

• Calculation Settings

3.3.16 Processing factors

Processing factors are multiplication factors to derive the concentration in a processed food from the concentration in an unprocessed food and can be specified for identified processing types (e.g., cooking, washing, drying). Processing factors are primarily used in dietary exposure assessments to correct for the effect of processing on substance concentrations in dietary exposure calculations.

This module has as primary entities: Foods Substances

Output of this module is used by: Concentrations Food conversions Dietary exposures Single value dietary exposures

Processing factors from data

Processing factors calculation

Processing factors fixed or distribution based

Processing factors can be specified as fixed factors (nominal) or as statistical distributions for the variability across samples.

- The distribution is either *the logistic-normal distribution* for processing types with factors restricted between 0 and 1 (e.g. washing),
- or the lognormal distribution *the lognormal distribution* for processing types with non-negative factors (e.g. drying).

Variability distribution parameters are specified indirectly via the 50th and 95th percentile. Uncertainty for processing factors can be specified using uncertainty distributions of the same form as for variability. Uncertainty distribution parameters are specified indirectly via the 95th uncertainty percentiles on the 50th and 95th variability distribution percentiles.

For distribution based processing factors specify $f_{k,nominal}$ and $f_{k,upper}$ (Nominal and Upper in table Processing factors). Two situations are distinguished depending on the type of transformation.
Nonnegative processing factors

Equate the logarithms of $f_{k,nominal}$ and $f_{k,upper}$ to the mean and the 95% one-sided upper confidence limit of a normal distribution. This normal distribution is specified by a mean

$$ln(f_{k,nominal})$$

and a standard deviation

$$ln(f_{k,upper}) – ln(f_{k,nominal}) / 1.645$$

Processing factors between 0 and 1

Equate the logits of $f_{k,nominal}$ and $f_{k,upper}$ to the mean and the 95% one-sided upper confidence limit of a normal distribution. This normal distribution is specified by a mean

$$logit(f_{k,nominal})$$

and a standard deviation

$$logit(f_{k,upper}) – logit(f_{k,nominal}) / 1.645.$$

See also processing correction

Processing factors data formats

Processing factors connect to a food id and a processing type or (food) facet (FoodEx 2). The specification of a substance (id) is optional. Specify the unprocessed food code and the processing type or facet. The combination of idFood-idProcessingType represents a processed food.

Processing factors are defined for triplets of processing type, food, and substance. The processing types are defined in the processing types table and the processing factors are defined in the processing factors table.

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Processing factors

Processing factor records should be linked to processing types (or facets) using the processing type (or facet) code (idProcessingType) and for the foods and substances. The codes of the processing factor records should match the codes of the foods, substances, and processing type (of facets) definitions.

Name	Туре	Description	Aliases	Required
idProcessing- Type	AlphaNumeric (50)	The code of the processing type.	idProcessing- Type, ProcessingType- Id, ProcessingType, ProcType, facet, idFacet, codeFacet	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	No
idFood- Unprocessed	AlphaNumeric (50)	The code of the unprocessed food.	idFood- Unprocessed, Food- UnprocessedId, idFood, FoodId, Food- Unprocessed	Yes
Nominal	Numeric	The nominal value (best estimate of 50th percentile) of processing factor (defines median processing factor).	Nominal, ProcNom	Yes
Upper	Numeric	The upper value (estimate of 95th percentile or "worst case" estimate) of processing factor due to variability.	Upper, ProcUpp	No
Nominal- Uncertainty- Upper	Numeric	The upper 95th percentile of nominal value (Nominal) due to uncertainty. A standard deviation for uncertainty of the nominal value (Nominal) is derived using the nominal value (Nominal) and upper 95th percentile (NominalUncertaintyUpper).	Nominal- Uncertainty- Upper, ProcNomUnc- Upp	No
Upper- Uncertainty- Upper	Numeric	The upper 95th percentile of upper value (Upper) due to uncertainty. From the nominal value (Nominal), upper value (Upper) and the specified uncertainties of these values (NominalUncertaintyUpper and UpperUncertaintyUpper, respectively) the degrees of freedom of a chi-square distribution describing the uncertainty of the standard deviation for variability is derived.	Upper- Uncertainty- Upper, ProcUppUnc- Upp	No

Table 3.152: Table definition for Processing factors.

Accepted table names: ProcessingFactors, ProcessingFactor, Processing.

Processing factors settings

Calculation settings

Name	Туре	Description
Selected tier	SettingsTemplateType	Specifies all module settings should be set according to a pre-defined tier or using custom settings.
Apply processing factors	Boolean	Specified in table ProcessingFactor. If checked, processing fact are applied. Concentrations in the consumed food may be different from concentrations in the modelled food in monitorin programs (typically raw food) due to processing, such as peelin washing, cooking etc. If unchecked, no processing information used. This is in most (though not all) cases a worst-case assumption
Use distribution	Boolean	Probabilistic specifications of processing factors will be used
Ignore processing factors less than 1	Boolean	This setting will suppress the use of processing factors lower th 1 (it is used in the EFSA 2012 Pessimistic tier).
Use default processing factor for missing processing types	Boolean	Use default processing factor for missing processing types
Default processing factor for missing processing types (except foods with processing type equal to 'Unspecified' (default = 1))	Numeric	Specify a processing factor for missing processing types. When the processing type is 'Unspecified', the default processing factor is equal to 1.

Table 3.153: Calculation settings for module Processing factors.

Uncertainty settings

Table 3.154:	Uncertainty	settings for	module Proc	cessing factors.
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Name	Туре	Description
Resample processing factors	Boolean	Specifies whether processing factors are resampled from a
		parametric uncertainty distribution.

Processing factors tiers

Overview

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
A pl pl ce in fa to	true	true	true	true	true	true	true	true	true	true
U di tr bi ti	false	false	false	false	false	false	false	false	false	false
Ig no pr co in fa to le th 1	false	false	false	false	false	true	true	false	false	false

Table 3.155: Tier overview for module Processing factors.

Retrospective dietary CRA (EC 2018) - Acute / Tier I

Table 3.156: Tier definition for Retrospective dietary CRA (EC 2018) -

Acute / Tier I.

Name	Setting
Apply processing factors	true
Use distribution	false
Ignore processing factors less than 1	false

Retrospective dietary CRA (EC 2018) - Chronic / Tier I

Table 3.157: Tier definition for Retrospective dietary CRA (EC 2018) - Chronic / Tier I.

Name	Setting
Apply processing factors	true
Use distribution	false
Ignore processing factors less than 1	false

Retrospective dietary CRA (EC 2018) - Acute / Tier II

Table 3.158: Tier definition for Retrospective dietary CRA (EC 2018) - Acute / Tier II.

Name	Setting
Apply processing factors	true
Use distribution	false
Ignore processing factors less than 1	false

Retrospective dietary CRA (EC 2018) - Chronic / Tier II

Table 3.159: Tier definition for Retrospective dietary CRA (EC 2018) - Chronic / Tier II.

Name	Setting
Apply processing factors	true
Use distribution	false
Ignore processing factors less than 1	false

Retrospective dietary CRA (EFSA 2012) - Optimistic

Use the optimistic model settings according to the EFSA Guidance 2012. Concentration values are sampled using a sample-based empirical distribution. Available processing factors are applied. No unit variability model should be applied.

Table 3.160: Tier definition for Retrospective dietary CRA (EFSA 2012) - Optimistic.

Name	Setting
Apply processing factors	true
Use distribution	false
Ignore processing factors less than 1	false

Retrospective dietary CRA (EFSA 2012) - Acute / Pessimistic

Acute probabilistic exposure assessment using the pessimistic model settings according to the EFSA Guidance 2012. Only processing factors > 1 are applied. For unit variability, the Beta distribution is applied.

Table 3.161: Tier definition for Retrospective dietary CRA (EFSA 2012) - Acute / Pessimistic.

Name	Setting
Apply processing factors	true
Use distribution	false
Ignore processing factors less than 1	true

Retrospective dietary CRA (EFSA 2012) - Chronic / Pessimistic

Chronic probabilistic exposure assessment using the pessimistic model settings according to the EFSA Guidance 2012. Only processing factors > 1 are applied.

Table 3.162: Tier definition for Retrospective dietary CRA (EFSA 2012)

- Chronic / Pessimistic.

Name	Setting
Apply processing factors	true
Use distribution	false
Ignore processing factors less than 1	true

Retrospective dietary CRA (EFSA 2022) - Acute / Tier I

Table 3.163: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier I.

Name	Setting
Apply processing factors	true
Use distribution	false
Ignore processing factors less than 1	false

Retrospective dietary CRA (EFSA 2022) - Chronic / Tier I

Table 3.164: Tier definition for Retrospective dietary CRA (EFSA 2022) - Chronic / Tier I.

Name	Setting
Apply processing factors	true
Use distribution	false
Ignore processing factors less than 1	false

Retrospective dietary CRA (EFSA 2022) - Acute / Tier II

Table 3.165: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier II.

Name	Setting
Apply processing factors	true
Use distribution	false
Ignore processing factors less than 1	false

Retrospective dietary CRA (EFSA 2022) - Chronic / Tier II

Table 3.166: Tier definition for Retrospective dietary CRA (EFSA 2022) - Chronic / Tier II.

Name	Setting
Apply processing factors	true
Use distribution	false
Ignore processing factors less than 1	false

Prospective dietary CRA (EFSA 2023) - Acute / Tier II

Table 3.167: Tier definition for Prospective dietary CRA (EFSA 2023) - Acute / Tier II.

Name	Setting
Apply processing factors	true
Use distribution	false
Ignore processing factors less than 1	false

Prospective dietary CRA (EFSA 2023) - Chronic / Tier II

Table 3.168: Tier definition for Prospective dietary CRA (EFSA 2023) - Chronic / Tier II.

Name	Setting
Apply processing factors	true
Use distribution	false
Ignore processing factors less than 1	false

Prospective dietary CRA (EFSA 2024) - Acute / Tier II

Table 3.169: Tier definition for Prospective dietary CRA (EFSA 2024) - Acute / Tier II.

Name	Setting
Apply processing factors	true
Use distribution	false
Ignore processing factors less than 1	false

Prospective dietary CRA (EFSA 2024) - Chronic / Tier II

Table 3.170: Tier definition for Prospective dietary CRA (EFSA 2024) - Chronic / Tier II.

Name	Setting
Apply processing factors	true
Use distribution	false
Ignore processing factors less than 1	false

Processing factors uncertainty

Processing effects are modelled either by a fixed processing factor, or by a lognormal or logistic-normal distribution (depending on the distribution type of the *processing type*). In case of a fixed factor, the uncertainty distribution is lognormal or logistic-normal with the same mean μ as the fixed value, and with a standard deviation σ_{unc} which is calculated from the specified central value μ (or nominal) and an estimate of the p95 of the *uncertainty distribution* (set *NominalUncertaintyUpper* in the *table for Processing factors*).

The calculation is:

$$\sigma_{unc} = \frac{f(\textit{NominalUncertaintyUpper}) - f(\mu)}{1.645}$$

with f() = logit for the logistic-normal distribution (distribution type 1) and f() = ln for the lognormal distribution (distribution type 2). Values lower than 0.01 or higher than 0.99 (distribution type 1 only) are replaced by default

values (0.01 and 0.99); this is useful computationally to avoid problems. In each iteration of the uncertainty analysis a new value is drawn from this distribution to be used as a fixed factor in the Monte Carlo calculation. In case of distribution based processing factors (describing the variability of processing factors) two uncertainties can be specified. For σ_{unc} , specification and calculation is as before (set *NominalUncertaintyUpper* in the *table for Processing factors*).

The uncertainty about the variability standard deviation

$$\sigma_{var} = \frac{f(Upper) - f(\mu)}{1.645}$$

can be specified by the UpperUncertaintyUpper value. This value is specified as the p95 upper limit on Upper. The specified value is used to derive in a iterative search the number of degrees of freedom df (van der Voet et al. (2009)). In the uncertainty analysis, a modified chi-square distribution with df degrees of freedom is used to generate new values of σ_{var} . A very high value of df means little uncertainty and σ_{var} will be almost equal in all iterations of the uncertainty analysis. A df close to 0 means a large uncertainty and very different values of σ_{var} will be obtained in the iterations of the uncertainty analysis. The p95 upper limit on Upper is set through parameter UpperUncertaintyUpper.

Processing factors as data

Specify for a combination of processing type, food and substance the processing factor (nominal, upper).

- Processing factors data formats
- Processing factors from data
- Processing factors calculation

Settings used

• Calculation Settings

3.3.17 Single value concentrations

Single value concentrations data are the single value estimates (High Residue, Maximum Residue Limit, Supervised Trials Median Residue) of residue concentrations on modelled foods.

This module has as primary entities: Foods Substances

Output of this module is used by: Modelled foods Single value dietary exposures

Single value concentrations from data

Single value concentrations calculation

Single value concentrations as data are supplied as mean concentrations, median concentrations, highest residues, percentiles, LOQs or maximum residue limits. Specify the 'Use data' option in the interface. In a retrospective context, the single values are computed based on the concentration distributions available for the modelled food as supplied in the *Concentrations module*. Specify option 'Compute' in the Single value concentrations action.

Single value concentrations data formats

Single value concentrations data provides a single value concentration for a substance.

Download empty dataset template: Zipped CSV Excel

Concentration single values

The food codes (idFood) and substance codes (idSubstance) should match the codes of the foods and substances table respectively.

Name	Туре	Description	Aliases	Required
idFood	AlphaNumeric (50)	Code of the food of this concentration single value.	idFood, FoodId, Food	Yes
idSubstance	AlphaNumeric (50)	Code of the substance of this concentration single value.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
Value	Numeric	Concentration single value.	Value, Concentration, Concentration- Value	Yes
ValueType	ConcentrationLimit- ValueType	Value type of the concentration value.	Concentration- SingleValue- Type, Concentration- ValueType, SingleValue- Type, Concentration- Type, ValueType, Type	Yes
Percentile	Numeric	Percentile.	Percentile	No
Concentration- Unit	ConcentrationUnit	The unit of the concentration single value (default mg/kg).	Concentration- Unit, Unit	No
Reference	AlphaNumeric (200)	Reference to the source from which this concentration single value is obtained.	Reference, References, Source, Sources	No

Table 3.171:	Table definition	for Concentration	single values.
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 $\label{eq:accepted} Accepted \ table \ names: \ ConcentrationSingleValues, \ SingleValueConcentrations.$

Single value concentrations settings

Selection settings

Table 3.172:	Selection	settings for	module Single	value concentrations.
		0	U	

Name	Туре	Description
Use substance conversion factors	Boolean	Specifies whether to use substance conversion factors to conver measured substance concentrations to active substance concentrations.

Single value concentrations as data

Single value concentrations data are the single value concentrations of residues on modelled foods.

- Single value concentrations data formats
- Single value concentrations from data

Inputs used: Active substances

Calculation of single value concentrations

Single value concentrations are calculated as a percentile (p50, p97.5 or maximum residue limit) of the modelled food concentration distribution.

• Single value concentrations calculation

Inputs used: Concentrations Concentration limits Deterministic substance conversion factors

3.3.18 Substance authorisations

Substance authorisations specify which food/substance combinations are authorised for (agricultural) use. If substance authorisations are used, then only the food/substance combinations that are specified in the data are assumed to be authorised and all other combinations are assumed to be not authorised. This information may, for instance, be used to determine whether concentration measurements below the LOQ or LOD could be assumed true zeros. I.e., if a food/substance combinations is assumed to be unauthorised, then the LOQ, LOD may be assumed to be a zero.

This module has as primary entities: Foods Substances

Output of this module is used by: Concentrations Occurrence patterns Concentration models

Substance authorisations from data

Substance authorisations data formats

Authorised uses data provides information about whether substance use is allowed for specified foods. For cumulative exposure assessments, this information is used for imputation of censored values/missing values.

Download empty dataset template: Zipped CSV Excel

Authorised uses

The authorised uses table

Name	Туре	Description	Aliases	Required
idFood	AlphaNumeric (50)	The food code.	idFood, FoodId, Food	Yes
idSubstance	AlphaNumeric (50)	The substance code.	idSubstance, Substance, SubstanceId	Yes
Reference	AlphaNumeric (200)	External reference(s) to sources containing more information about the effect (key event) relationships.	Reference, References	No

Table 3.173: Table definition for Authorised uses	•
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Accepted table names: AuthorisedUses, AuthorisedUse.

Substance authorisations as data

Substance authorisations are specified as data in the form of a list of authorised food/substance combinations, with combinations not on the list associated with no authorised use.

• Substance authorisations data formats

• Substance authorisations from data

3.3.19 Substance approvals

Substance approvals specify which substances are approved within the definition under regulation (EC) No 1107/2009. This information may, for instance, be used to to restrict water imputation to approved substances only.

This module has as primary entities: *Substances*

Output of this module is used by: Concentrations

Substance approvals from data

Substance approvals data formats

Substance approvals specify which substances are approved within the definition under regulation (EC) No 1107/2009. This information may, for instance, be used to restrict water imputation to approved substances only.

Download empty dataset template: Zipped CSV Excel

Substance approvals

Substance approval records are tuples consisting of an identifier of the substance and a boolean value specifying the approval status of the substance. Substances that are not included in the table are assumed to be NOT approved.

Name	Туре	Description	Aliases	Required
idSubstance	AlphaNumeric (50)	The substance code.	idSubstance, Substance, SubstanceId	Yes
IsApproved	Boolean	Specifies whether the substance is approved or not. Substances not included in the table are assumed to be NOT approved.	IsApproved, Approved	Yes

Table 3.174: Table definition for Substance approvals.

Accepted table names: SubstanceApprovals, ApprovedSubstances.

Substance approvals as data

Substance approvals are specified as data in the form of a list of approved substances. Substances not on the list are assumed to be not approved.

- Substance approvals data formats
- Substance approvals from data

3.3.20 Substance conversions

Substance conversions specify how measured substances are converted into active substances, which are the substances assumed to cause health effects. In pesticide legislation such measured substances and the substance conversion rules are known as residue definitions.

This module has as primary entities: *Substances*

Output of this module is used by: Concentrations

Substance conversions from data

Substance conversions data formats

Two types of substance conversions are implemented, with two subtypes for the first type:

1a) The measured substance is one or more of a set of possible substances (e.g. isomers or metabolites), and the toxicity of all substances in this set is assumed to be the same and is expressed in one active substance. Example: The measured substance Parathion-methyl(RD) is either Parathion-methyl or paraoxon-methyl, but both are expressed as the active substance Parathion-methyl.

1b) The measured substance is one or more of a set of possible substances (e.g. isomers or metabolites), and the toxicity of all substances in this set is assumed to relate with equal probability to one of a subset of active substances. Example: The measured substance Dithiocarbamates includes the active substances maneb, mancozeb, metiram, propineb, thiram and ziram, one of which will be assumed to be the active substance present with equal probability.

2) If *n active substances* all metabolise to the same active substance (the metabolite), it is assumed that all n+1 substances have equal probability of being the source of the measured concentration. The measured substance then is either one active substance (the metabolite) or a mixture of two active substances, one being the metabolite and the other one of the possible parent substances. Example: the measured substance Carbofuran(RD) is either the active substance Carbufuran or a mixture of Carbofuran and one of the possible active parent substances Benfuracarb or Carbosulfan.

Note, it is not allowed to have conversion factors equal to 0.

Substance conversions are described by a single substance conversions table.

Download empty dataset template: Zipped CSV Excel

Substance conversion rules

The records of the substance translations definitions table specify which active substances (idActiveSubstance) link to a measured substance (idMeasuredSubstance). Each record contains a conversion factor that specifies how a concentration of the measured substance translates to a concentration of the active substance, a flag that states whether the residue definition should be assumed to translate exclusively to one of its active substances, and a proportion. The proportion specifies the proportion of the samples that should translate to this specific active substance in case the translation is exclusive, otherwise it specifies the proportion of the concentration that is assumed to be attributed to the active substance.

Name	Туре	Description	Aliases	Required
idMeasured- Substance	AlphaNumeric (50)	Substance code of the measured substance.	idResidue- Definition, Residue- Definition, Measured- Substance	Yes
idActive- Substance	AlphaNumeric (50)	Substance code of the active substance.	idActive- Substance, idSubstance, Active- Substance, Substance	Yes
Conversion- Factor	Numeric	Specifies the (molecular weight) conversion factor to translate the concentration of the residue definition to a concentration of the active substance	Conversion- Factor	Yes
IsExclusive	Boolean	Specifies whether a measurement of the residue substance should be translated exclusively to this active substance, or if the residue definition represents/breaks down to a mixture of active substances.	IsExclusive	Yes
Proportion	Numeric	Only applicable for non-exclusive conversions. The proportion of the concentration that is assumed to be attributed to the active substance.	Proportion	No

Гаble 3.175: Та	able definition	for Substance	conversion rules.
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Accepted table names: ResidueDefinitions, ResidueDefinition.

Substance conversions as data

Substance conversions are provided as data.

- Substance conversions data formats
- Substance conversions from data

Inputs used: Active substances

3.3.21 Total diet study sample compositions

Total diet study sample compositions specify the composition of mixed food samples, such as used in a total diet study (TDS), in terms of their constituting foods.

This module has as primary entities: Foods

Output of this module is used by: Concentration models Food conversions

Total diet study sample compositions from data

Total diet study sample compositions data formats

Total diet studies (TDS) complement traditional monitoring of substance concentrations on raw commodities by measuring substance occurrence in main foods prepared as consumed and pooled into representative food groups. To include occurrence data from TDS for exposure assessment, the composition of the TDS samples is needed in order to link the composite samples to the consumed foods (either directly or indirectly). TDS composition data describes the composition of TDS samples by specifying the foods (and the amounts) of TDS samples.

Download empty dataset template: Zipped CSV Excel

TDS food sample compositions

The TDS food sample compositions table contains the descriptions of the TDS samples and specifications of the foods (with amounts) included in the TDS samples.

Name	Туре	Description	Aliases	Required
idTDSFood	AlphaNumeric (50)	The code of the TDS food.	idTDSFood	Yes
idFood	AlphaNumeric (50)	Sub-food of the TDS food.	idFood	Yes
PooledAmount	Numeric	Total weight (in g) or volume (in ml) of the food.	PooledAmount, Weight	Yes
Description	AlphaNumeric (200)	Additional description of the TDS sample (e.g. number of subsamples).	Description	No
Regionality	AlphaNumeric (250)	Regionality information.	Regionality	No
Seasonality	AlphaNumeric (250)	Seasonality information.	Seasonality	No

Fable 3 176 .	Table	definition	for	TDS	food	samr	le	com	osition	2
1 auto 5.170.	rabic	ucilition	101	IDS	1000	Samp	лс	comp	Jositiona	5.

Accepted table names: TDSFoodSampleCompositions, TDSFoodSampleComposition, CompositionTDSFoodSamples, CompositionTDSFoodSample.

Total diet study sample compositions as data

Total diet study sample compositions are provided as data.

- Total diet study sample compositions data formats
- Total diet study sample compositions from data

3.3.22 Unit variability factors

Unit variability factors specify the variation in concentrations between single units of the same food, which have been put together in a mixture sample on which the concentration measurements have been made. Unit variability factors are used for *modelling unit variability* in acute *(individual) dietary exposures calculations* to account for the fact that concentration data often relate to composite samples, whereas an acute risk may result from consumption of single food units. For the same purpose, they are also used in the *IESTI model* for *single value dietary exposures calculations*.

This module has as primary entities: Foods Substances

Output of this module is used by: Dietary exposures Single value dietary exposures

Unit variability factors from data

Unit variability factors data formats

Unit variability factors specify the unit-to-unit variation of substance concentrations on foods. Unit variability factors are described by a single unit variability factors table.

Download empty dataset template: Zipped CSV Excel

Unit variability factors

Unit variability factors are defined for a food, and may possibly also be specified for a specific substance and/or processing type. The unit variability factors are linked to the foods by means of the food code (idFood). Unit variability factors can be specified as unit variability factors (P97.5/mean) or as coefficients of variation of a statistical distribution.

Name	Туре	Description	Aliases	Required
idFood	AlphaNumeric (50)	The food code.	idFood, FoodId, Food	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	No
idProcessing- Type	AlphaNumeric (50)	The processing type code.	idProcessing- Type, ProcessingType- Id, ProcessingType, ProcType	No
Factor	Numeric	The variability factor.	Factor, VarFac, VariabilityFactor	No
UnitsIn- Composite- Sample	Numeric	The number of units in the composite sample.	UnitsIn- Composite- Sample, NoUnitComp	Yes
Coefficient	Numeric	The coefficient of variation.	Coefficient, Variability- Coefficient, CoefVar, VarCoef	No

Table 3.177: Table definition for Unit variability factors.

Accepted table names: UnitVariabilityFactors, UnitVariabilityFactor, VariabilityFactor, VariabilityFactors, VariabilityProcCompProd, UnitVariability.

IESTI special cases

IESTI special cases for specified combinations of food, substance. The application type (post-harvest or pre-harvest) determines whether Case 1 or Case 3 should be used.

Name	Туре	Description	Aliases	Required
idFood	AlphaNumeric (50)	The unique identification code of the food.	idFood, Code, FoodId, FoodCode, Food, Id	Yes
idSubstance	AlphaNumeric (50)	The unique identification code of the substance. This code may be from an existing coding system, such as CAS-codes or Param codes of EFSA, or it may be a used-defined code.	idSubstance, SubstanceId, Substance, Code, Id	Yes
Application- Type	HarvestApplication- Type	Harvest application type (pre-harvest or post-harvest).	Application- Type, Harvest- ApplicationType	Yes
Reference	AlphaNumeric (200)	External reference(s) to pre-harvest use.	Reference	No

Table 3.178: Table definition for IESTI special	cases.
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Accepted table names: IestiSpecialCases.

Unit variability factors

Calculation settings

Name	Туре	Description
Selected tier	SettingsTemplateType	Specifies all module settings should be set according to a pre-defined tier or using custom settings.
Unit variability model	UnitVariabilityModelType	Describes variation between single units when concentration da are from composite samples.
Estimates nature	EstimatesNature	Simulated unit concentrations can be higher or lower than composite value (realistic) or only equal or higher (conservative
Unit variability parameter	UnitVariabilityType	Use Coefficient of variation or Variability factor, specified in VariabilityFactor table.
Mean of LogNormal simulated values (biasing)	MeanValueCorrectionType	Unbiased: correct unit simulations for difference between medi and mean.
Default variability factor for unit weight <= 25g	Numeric	Default variability factor 1 (unit weight <= 25 g, small crops). S requires specification of unit weight (FoodProperties table) and case of beta model, also the Number of units in a composite sample (UnitVariability table).
Correlation between substances on the same units	UnitVariabilityCorrelationType	Specifies the type of correlation between substances on the samunits; no correlation or full correlation.
Default variability factor for unit weight > 25g	Numeric	Default variability factor 5 (unit weight > 25 g, medium/large crops). Still requires specification of unit weight (FoodPropertitable) and, in case of beta model, also the Number of units in a composite sample (UnitVariability table).

Table 3.179: Calculation settings for module Unit variability factor	ors.
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Unit variability factors tiers

Overview

N	EC 2018 Acute Tier I	EC 2018 Acute Tier II	EFSA 2012 Acute Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Acute Tier II	EFSA 2023 Acute Prospec- tive Tier II	EFSA 2024 Acute Prospec- tive Tier II
U va al it m	BetaDis- tribution	BetaDis- tribution	BetaDis- tribution	BetaDis- tribution	BetaDis- tribution	BetaDis- tribution	BetaDis- tribution
E ti m n: tu	Realistic	Realistic	Realistic	Realistic	Realistic	Realistic	Realistic
U va al it pa ra e- te	Variabili- tyFactor	Variabili- tyFactor	Variabili- tyFactor	Variabili- tyFactor	Variabili- tyFactor	Variabili- tyFactor	Variabili- tyFactor

Table 3.180: Tier overview for module Unit variability factors.

Retrospective dietary CRA (EC 2018) - Acute / Tier I

Table 3.181: Tier definition for Retrospective dietary CRA (EC 2018) - Acute / Tier I.

Name	Setting
Unit variability model	BetaDistribution
Estimates nature	Realistic
Unit variability parameter	VariabilityFactor

Retrospective dietary CRA (EC 2018) - Acute / Tier II

Table 3.182: Tier definition for Retrospective dietary CRA (EC 2018) - Acute / Tier II.

Name	Setting
Unit variability model	BetaDistribution
Estimates nature	Realistic
Unit variability parameter	VariabilityFactor

Retrospective dietary CRA (EFSA 2012) - Acute / Pessimistic

Acute probabilistic exposure assessment using the pessimistic model settings according to the EFSA Guidance 2012. Only processing factors > 1 are applied. For unit variability, the Beta distribution is applied.

Name	Setting
Unit variability model	BetaDistribution
Estimates nature	Realistic
Unit variability parameter	VariabilityFactor

Table 3.183: Tier definition for Retrospective dietary CRA (EFSA 2012) - Acute / Pessimistic.

Retrospective dietary CRA (EFSA 2022) - Acute / Tier I

Table 3.184: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier I.

Name	Setting
Unit variability model	BetaDistribution
Estimates nature	Realistic
Unit variability parameter	VariabilityFactor

Retrospective dietary CRA (EFSA 2022) - Acute / Tier II

Table 3.185: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier II.

Name	Setting
Unit variability model	BetaDistribution
Estimates nature	Realistic
Unit variability parameter	VariabilityFactor

Prospective dietary CRA (EFSA 2023) - Acute / Tier II

Table 3.186: Tier definition for Prospective dietary CRA (EFSA 2023) - Acute / Tier II.

Name	Setting
Unit variability model	BetaDistribution
Estimates nature	Realistic
Unit variability parameter	VariabilityFactor

Prospective dietary CRA (EFSA 2024) - Acute / Tier II

Table 3.187: Tier definition for Prospective dietary CRA (EFSA 2024) - Acute / Tier II.

Name	Setting
Unit variability model	BetaDistribution
Estimates nature	Realistic
Unit variability parameter	VariabilityFactor

Unit variability factors as data

Unit variability factors are provided as data.

- Unit variability factors data formats
- Unit variability factors from data

Settings used

• Calculation Settings

3.4 Exposure modules

Exposures are, in the simplest applications, *dietary exposures*, which combine consumption and occurrence data, either for single or for multiple *substances* causing the same adverse *effect*. Links between the foods-as-eaten and the *modelled foods* are made using *food conversions*, and the consumptions are expressed as *consumptions per modelled food*. For large assessment groups, the use of *dietary exposures screening* may be used to reduce the complexity of the calculations and only focus calculations on the risk drivers.

In aggregate exposure assessments, *exposures* combine *dietary exposures* with *non-dietary exposures*, which have to be entered as pre-calculated data.

Human monitoring data can be compared to exposures using human monitoring analysis.

In cumulative assessments, important mixtures of substances can be identified using exposure mixtures.

3.4.1 Consumptions by modelled food

Consumptions by modelled food are consumptions of individuals expressed on the level of the foods for which concentration data are available (i.e., the modelled-foods). These are calculated from consumptions of foods-as-eaten and food conversions that link the foods-as-eaten amounts to modelled-foods amounts.

This module has as primary entities: Populations Foods Substances

Output of this module is used by: Single value consumptions High exposure food-substance combinations Dietary exposures

Consumptions by modelled food calculation

Consumptions by modelled food are calculated from *consumptions* of *modelled foods* and *food conversions* that link the foods-as-eaten amounts to modelled-foods amounts. Given that the food conversion is already available, the procedure for computing the consumptions by modelled-food is straightforward. For each consumption of each individual, a modelled-food consumption record is created for each modelled-food that is linked to the consumed foods through the food conversion, with the amount being the total consumption amount multiplied by the proportion indicated by the food conversion. Also, if in the *food conversion algorithm* one or more *processing types* are found, then these types are recorded in the consumption by modelled food record.

Consumptions by modelled food settings

Calculation settings

Table 3.188:	Calculation	settings for	or module	Consumptions	by modelled
food.					

Name	Туре	Description
Restrict population to consumers or consumer days only (food-as-measured)	Boolean	Specifies whether the population should be restricted to the individuals (chronic) or individual days (acute) with consumpti containing any of the modelled foods.
Exposure type	ExposureType	The type of exposure considered in the assessment; acute (shor term) or chronic (long-term).
Restrict population to consumers or consumer days with consumptions of specified modelled foods only	Boolean	Specifies whether the population should be restricted to the individuals (chronic) or individual days (acute) with consumpti containing any of the specified modelled food subset.
Selected modelled foods	AlphaNumeric	Set of consumed modelled foods that are of particular interest restricting the consumers / consumption days.
Apply processing factors	Boolean	Specified in table ProcessingFactor. If checked, processing factor are applied. Concentrations in the consumed food may be different from concentrations in the modelled food in monitorin programs (typically raw food) due to processing, such as peelir washing, cooking etc. If unchecked, no processing information used. This is in most (though not all) cases a worst-case assumption

Output settings

Name	Туре	Description
Lower percentage for variability (%)	Numeric	The default value of 25% may be overruled.
Upper percentage for variability (%)	Numeric	The default value of 75% may be overruled.

Calculation of consumptions by modelled food

Consumptions by modelled food are calculated from consumptions of foods-as-eaten and food conversions that link the foods-as-eaten amounts to modelled-foods amounts.

• Consumptions by modelled food calculation

Inputs used: Consumptions Food conversions

Settings used

• Calculation Settings

3.4.2 Dietary exposures

Dietary exposures are the amounts of substances, expressed per kg bodyweight or per individual, to which individuals in a population are exposed from their diet per day. Depending on the exposure type, dietary exposures can be short-term/acute exposures and then contain exposures for individual-days, or they can be long-term/chronic exposures, in which case they represent the average exposure per day over an unspecified longer time period.

This module has as primary entities: Populations Foods Substances Effects

Output of this module is used by: Internal exposures Exposure mixtures Dust exposures Soil exposures Air exposures Risks Environmental burden of disease

Dietary exposures calculation

In probabilistic exposure assessment we consider a population of individuals. Exposure assessment with MCRA can address *acute exposure* or *chronic exposure*. Acute exposure is relevant when the short-term effect on individuals is relevant, chronic exposure when the long-term effects on the individuals matter. In MCRA short-term is operationalised as one day, so effectively acute exposure assessment is concerned with a population of person-days, whereas chronic exposure assessment is concerned with a population of persons.

The basic operation in exposure assessment is integrating consumptions and concentrations per food. With multiple foods, consumptions are typically correlated, therefore MCRA works with the multivariate distribution of a consumption vector, as represented by the consumption data of individuals in a consumption survey. In contrast, the distributions of concentration for each food are typically considered to be independent between foods. E.g., eating an apple with an accidentally high residue concentration does not predict that another food eaten on the same day will also have a high residue concentration. As a consequence of this assumption, concentrations of substances are modelled for each food independently.

For large assessment groups, the use of *dietary exposures screening* may be used to reduce the complexity of dietary exposures calculations and only focus calculations on the risk drivers. In this case, only detailed information is recorded for the risk drivers. With or without screening MCRA produces the same estimated cumulative exposure distribution summarized by percentiles and exceedance percentages, the same contributions of all substances and all modelled foods. After screening, contributions related to food-as-eaten are available for the risk drivers only.

In cumulative exposure calculations two simple approaches are used to identify and select mixtures contributing to the exposure of a target population:

1. qualitative approach: counting of co-exposure. To which combinations of substances are individuals exposed?

Co-exposure of substances is a qualitative approach where the number of combinations of substances to which an individual is exposed is recorded. There is no cut-off level, the only criterion is the presence of a substance in the simulated daily diet or not. For an *acute* or short term exposure assessment, a simulated individual day is the smallest entity to determine co-exposure. For a *chronic* or long term exposure assessment, co-exposures are summarized at the individual level, e.g. co-exposure is determined combining all consumption days of an individual. For more information see *co-exposure of substances*.

2. quantitative approach: *maximum cumulative ratio (MCR)*. To what degree are mixtures more important than single substances?

For a quantitative approach, see also the *exposure mixtures module*.

Acute exposure assessment

In an acute exposure assessment, the short term exposure to a substance or group of substances is estimated. The interest is in the distribution of individual day exposures and derived statistics like the fraction of days that exceed an intake limit or point of departure (PoD). The PoD is calculated as the acute reference dose (ARfD) * safety factor (SF). The basic model for the exposure to a substance in an acute exposure assessment is:

$$y_{ij} = \frac{\sum_{k=1}^{p} x_{ijk} c_{ijk}}{b w_i}$$

where y_{ij} is the intake by individual *i* on day *j* (in microgram substance per kg body weight), x_{ijk} is the consumption by individual *i* on day *j* of food *k* (in g), c_{ijk} is the (*simulated*) concentration of that substance in food *k* eaten by individual *i* on day *j* (in mg/kg), and bw_i is the body weight of individual *i* (in kg). Finally, *p* is the number of foods accounted for in the model. Within parenthesis, the default unit definitions are assumed, but decimal multiples or submultiples of units are easily specified using the relevant tables.

In the exposure assessment, individual days enter the Monte Carlo sample using the inverse of the sampling weights w_i when the number of MC iterations is > 0 (see *table for Individuals*, field *SamplingWeight*).



Contribution to total exposure distribution for foods as measured

Figure 3.29: Example MCRA dietary exposure contributions modelled foods.

Contribution to total exposure distribution for foods as eaten



Figure 3.30: Example MCRA dietary exposure contributions foods as eaten



Contribution to total exposure distribution for substances

Figure 3.31: Example MCRA dietary exposure contributions substances

Contribution to total exposure distribution for foods as modelled x substances (MSCC)



Figure 3.32: Example MCRA dietary exposure contributions modelled foods x substances

Modelling unit-to-unit variation

The basic model for an acute exposure assessment assumes that the concentration of the substance displays the variation of residues between units in the marketplace. In general, both monitoring data and controlled field trial data are obtained using composite samples. As a result some of the unit-to-unit variation is averaged out. The model for unit variability aims to adjust the composite sample mean such that sampled concentrations represent the originally unit-to-unit variation of the units in the composite sample.

MCRA offers three distributions to sample from:

- 1. the beta distribution,
- 2. the lognormal distribution,
- 3. and the bernoulli distribution.

The beta distribution simulates values for a unit in the composite sample. It requires knowledge of the number of units in a composite sample and of the variability between units.

The lognormal distribution simulates values for a new unit in the batch. It requires only knowledge of the variability between units.

The bernoulli distribution is considered as a limiting case of the beta distribution when knowledge of the variability between units is lacking and only the number of units in the composite sample is known. For the beta and lognormal distribution, estimates of unit variability are either realistic (no censoring at the value of the monitoring residue) or conservative (unit values are left-censored at the value of the monitoring residue). For the lognormal distribution sampled concentrations have no upper limit. Whereas for the beta distribution, sampled concentration values for a unit are never higher than the monitoring residue times the number of units in the composite sample.

Variability between units is specified using a variability factor v (defined as 97.5th percentile divided by mean) or a coefficient of variation c_v (standard deviation divided by mean). Following FAO/WHO recommendations, the default variability factor v = 1 for small crops (unit weight < 25 g). For large crops (unit weight > 25 g) v = 5. For foods which are processed in large batches, e.g. *juicing, marmalade/jam, sauce/puree, bulking/blending* the variability factor v = 1 is proposed.

Estimation of intake values using the concept of unit variability

A composite sample for food k is composed of nu_k units with nominal (whole food/RAC) unit weight wu_k . The weight of a composite sample is $wm_k = nu_k \cdot wu_k$ with mean residue value cm_k .

- For each iteration i in the MC-simulation, obtain for each food k a simulated intake x_{ik} , and a simulated composite sample concentration cm_{ik} .
- Calculate the number of unit intakes nux_{ik} in x_{ik} (round upwards) and set weights w_{ikl} equal to unit weight wu_k , except for the last partial intake, which has weight $w_{ikl} = x_{ik} (nux_{ik} 1)wu_k$.
- For the beta or bernoulli distribution: draw nux_{ik} simulated values bc_{ikl} from a beta or bernoulli distribution. Calculate concentration values as $c_{ikl} = bc_{ikl} \cdot cm_{ik,max} = bc_{ikl} \cdot cm_{ik} \cdot nu_k = svf_{ikl} \cdot cm_{ik}$, where nu_k is the number of units in a composite sample of food k, and svf_{ikl} is the stochastic variability factor for this simulated unit, i.e. the ratio between simulated concentration c_{ikl} and the simulated composite sample concentration cm_{ik} . Sum to obtain the simulated concentration in the consumed portion:

$$c_{ik} = \sum_{l=1}^{nux_{ik}} w_{ikl} c_{ikl} / x_{ik}$$

• For the lognormal distribution: draw nux_{ik} simulated logconcentration values lc_{ikl} from a normal distribution with (optional) a biased mean $\mu = ln(cm_{ik})$ or (default) unbiased mean $\mu = ln(cm_{ik}) - 1/2\sigma^2$ and standard deviation σ . Calculate concentration values as

$$c_{ikl} = \exp(lc_{ikl}) = svf_{ikl} * cm_{ikl}$$

where svf_{ikl} is the stochastic variability factor for this simulated unit, i.e. the ratio between simulated concentration c_{ikl} and the simulated composite sample concentration cm_{ik} . Back transform and sum to obtain the simulated

concentration in the consumed portion:

$$c_{ik} = \sum_{l=1}^{nux_{ik}} w_{ikl} c_{ikl} / x_{ik}$$

For cumulative exposure assessments, a sensitivity analysis may be performed by specifying a full correlation between concentrations from different substances on the same unit. As a result, high (or low) concentrations from different substances occur together on the same unit. In MCRA, for each unit the random sequence is repeatedly used to generate concentration values for all substances.

Beta distribution

Under the beta model simulated unit values are drawn from a bounded distribution on the interval $(0, c_{max})$ with $c_{max} = nu_k \cdot cm_k$. The standard beta distribution is defined on the interval (0, 1) and is usually characterised by two parameters a and b, with a > 0, b > 0 (see e.g. Mood et al. (1974)). Alternatively, it can be parameterised by the mean

$$\mu = a/(a+b)$$

and the variance

$$\sigma^2 = ab/(a+b+1)^{-1}(a+b)^{-2}$$

or, as applied in MCRA, by the mean μ and the squared coefficient of variation

$$c_{\rm v}^2 = ba^{-1}(a+b+1)^{-1}$$

For the simulated unit values in *each iteration of the program* we require an expected value cm_k . This scales down to a mean value $\mu = cm_k/c_{max} = 1/nu_k$ in the (standard) beta distribution. From this value for μ and an externally specified value for cv_k the parameters a and b of the beta distribution are calculated as:

$$a = b(nu_k - 1)^{-1}$$

and

$$b = \frac{(nu_k - 1)(nu_k - 1 - cv_k^2)}{nu_k cv_k^2}$$

From the second formula it can be seen that cv_k should not be larger than $\sqrt{nu_k - 1}$ in order to avoid negative values for b. When the unit variability is specified by a variability factor

$$v_k = \frac{p97.5_k}{cm_k}$$

instead of a coefficient of variation cv_k then MCRA applies a bisection algorithm to find a such that the cumulative probability

$$P[Beta(a,b)] = 0.975$$

for $b = a(nu_k - 1)$.

Sampled values from the beta distribution are rescaled by multiplication with cm_{max} to unit concentrations c_{ijk} on the interval $(0, cm_{max})$.

Lognormal distribution

The lognormal distribution is characterised by μ and σ , which are the mean and standard deviation of the logtransformed concentrations. The unit log-concentrations are drawn from a normal distribution with mean $\mu = ln(cm_{ik}) - 1/2\sigma^2$. The coefficient of variation cv is turned into the standard deviation σ on the log-transformed scale with:

$$\sigma = \sqrt{\ln(cv^2 + 1)}$$

The variability factor is defined as the 97.5th percentile of the concentration in the individual measurements divided by the corresponding mean concentration seen in the composite sample. A variability factor v is converted into the standard deviation σ as follows:

$$v = \frac{p97.5}{mean} = \frac{e^{\mu + 1.96\sigma}}{e^{\mu + 1/2\sigma^2}} = e^{1.96\sigma - 1/2\sigma^2}$$

with μ and σ representing the mean and standard deviation of the log-transformed concentrations. So

$$ln(v) = 1.96\sigma - 1/2\sigma^2$$

Solving for σ gives:

$$\sigma^2 - 2 \cdot 1.96\sigma + 2\log(v) = 0$$

with roots for σ according to:

$$\sigma = 1.96 \pm \sqrt{(1.96^2 - 2log(v))}$$

The smallest positive root is taken as an estimate for σ .

Bernoulli distribution

The bernoulli model is a limiting case of the beta model, which can be used if no information on unit variability is available, but only the number of units in a composite sample is known (see van der Voet et al. 2001). As a worst case approach we may take the coefficient of variation cv as large as possible. When cv is equal to the maximum possible value $\sqrt{nu_k - 1}$, the (unstandardised) beta distribution simplifies to a bernoulli distribution with probability

 $(nu_k-1)/nu_k$

or

$$(v_k-1)/v_k$$

 $1/nu_k$

 $1/v_k$

for the value 0 and probability

or

for the value $c_{max} = nu_k \cdot cm_k$.

In MCRA values 0 are actually replaced by cm_k , to keep all values on the conservative side. For example, with $nu_k = 5$, there will be 80% probability at $c_{ijk} = cm_k$ and 20% probability at $c_{ijk} = c_{max}$. When the number of units nu_k in the composite sample is missing, the nominal unit weight wu_k is used to calculate the parameter for unit variability.

Chronic exposure assessment

In a chronic exposure assessment, usual exposure is defined as the long-run average of daily exposure to a substance or group of substances by an individual. The interest is in the distribution of individual exposures and derived statistics like the fraction of individuals that exceed an intake limit or point of departure *PoD*). The PoD is calculated as the average daily intake (ADI) * safety factor (SF). Usually, for an individual, dietary recall data are available on 2 (or more) consecutive days. We assume an equal number of days for each individual, unless specified differently in *table for Individuals*.

For a chronic exposure assessment the available data are used to calculate exposures per person-day (daily exposure):

$$y_{ij} = \frac{\sum_{k=1}^{p} x_{ijk} c_{ijk}}{b w_i}$$

where y_{ij} , x_{ijk} and bw_i are defined as before but now concentrations of the substance found in food k enter the model as the *estimated mean substance concentration value* c_k . Using the person-day exposures MCRA, provides a number of *exposure models* to calculate the distribution of usual exposure at the person level.

Chronic exposure models

Using the person-day exposures MCRA uses one of the following models to calculate the distribution of usual exposure at the person level:

- 1. The observed individual means observed individual means (OIM) model;
- 2. The *logisticnormal-normal (LNN) model*, in a full version that includes the estimation of correlation between exposure frequency and amount, and in a simpler version without this estimation;
- 3. The betabinomial-normal (BBN) model;
- 4. The *discrete/semi-parametric* model known as the Iowa State University Foods (ISUF) model. For this model, an equal number of days per individual is assumed.

In modelling usual exposure, two situations can be distinguished. Foods are consumed on a *daily basis* or foods are *episodically consumed*. For the logistic normal-normal model and the betabinomial-normal model, the latter requires fitting of a two-part model,

- 1. a model for the frequency of consumption, and
- 2. a model for the exposure amount on consumption days.

In the final step, both models are integrated in order to obtain the usual exposure distribution. For daily consumed foods, fitting of the frequency of consumption is skipped and modelling resorts to fitting the model to daily exposure amounts only. Note that the distinction between BNN and LNN disappears and modelling will give equivalent results.

Observed individual means (OIM)

The usual exposure distribution for a population is estimated with the empirical distribution of individual means. Each mean is the average of all single-day exposures for an individual. The mean value for an individual still contains a considerable amount of within-individual variation. As a consequence, the distribution of within-individual means has larger variance than the true usual exposure distribution and estimates using the OIM-method are biased, leading to a too high estimate of the fraction of the population with a usual exposure above some standard. Despite its known tendency to over-estimate high-tail exposures, the OIM method is the method to be used in EFSA (2012) basic assessments.

Model based and model assisted

Following Kipnis et al. (2009), some of the models available in MCRA are extended to predict individual usual exposures. This model assisted approach has been added to BBN and LNN when used without correlation) and may be a useful extension in evaluating the relationship between health outcomes and individual usual exposures of foods. In contrast, the estimation of the usual exposure distribution in the general population is called the model based approach. Summarizing, we get Table 3.190:

F	
Model based approach	Model assisted approach
	observed individual means (OIM)
betabinomial-normal (BBN)	betabinomial-normal (BBN)
logisticnormal-normal (LNN) without correlation	logisticnormal-normal (LNN) without correlation
logisticnormal-normal (LNN) with correlation	
Iowa State University Foods (ISUF)	

Table 3.190: Model based and assisted approach available for chronic exposure models

The model assisted approach builds on the proposal of Kipnis et al. (2009), but is modified to ensure that the population mean and variance are better represented. The method is based on shrinkage of the observed individual means (modified BLUP estimates) and shrinkage of the observed exposure frequencies. The model-assisted usual exposure distribution applies to the population for which the consumption data are representative, and automatically integrates over any covariates present in the model. Model-assisted exposures are not yet available for LNN, and when a covariable is modelled by a spline function of degree higher than 1. In case of a model with covariates the usual exposure is presented in graphs and tables as a *function of the covariates* (conditional usual exposure distributions).

Betabinomial-Normal model (BBN)

The *Betabinomial-Normal (BBN)* model for chronic risk assessment is described in de Boer et al. (2009), including its near-identity to the STEM-II model presented in Slob (2006). The BBN model combines a betabinomial model for the exposure frequencies with a normal model for transformed positive exposures.

Logisticnormal-Normal model (LNN with and without correlation)

In the logistic normal-normal (LNN) model, exposure frequencies are modelled by a logistic normal distribution. In notation, for probability p:

 $logit(p) = log(p/1 - p) = \mu - i + \underline{c}_i$

where μ_i represents the person specific fixed effect model and \underline{c}_i represent person specific random effects with estimated variance component $\sigma_{between}^2$. Similarly as in the BBN model, the positive exposure amounts are modelled, after transformation (logarithmic or *Box-Cox*), with a normal distribution. This model is referred to as the *LogisticNormal-Normal (LNN)* model. The full *LNN model* model includes the estimation of a correlation between exposure frequency and exposure amount. This is similar to the NCI model described in Tooze et al. (2006). A simple and computationally less demanding version of the LNN method does not estimate the correlation between frequency and amount. The models are fitted by maximum likelihood, employing *Gauss-Hermite integration*.

For chronic models amounts are usually transformed before the statistical model is fit. The power transformation, given by y^p , has been replaced by the equivalent Box-Cox transformation. The Box-Cox transformation is a linear function of the power transformation, given by $(y^p - 1)/p$, and has a better numerical stability. *Gauss-Hermite integration* is used for back-transformation (see also *Box Cox power transformation*).

Discrete/semi-parametric model (ISUF)

Nusser et al. (1996) described how to assess chronic risks for data sets with positive exposures (a small fraction of zero exposures was allowed, but then replaced by a small positive value). The modelling allowed for heterogeneity of variance, e.g. the concept that some people are more variable than others with respect to their consumption habits. However, a disadvantage of the method was the restricted use to contaminated foods which were consumed on an almost daily basis, e.g. dioxin in fish, meat or diary products. The estimation of usual exposure from data sets with a substantial amount of zero exposure became feasible by modelling separately zero exposure on part or all of the days via the estimation of exposure probabilities as detailed in Nusser et al. (1997) and Dodd (1996). In MCRA, a discrete/semi-parametric model is implemented allowing for zero exposure and heterogeneity of variance following the basic ideas of Nusser et al. (1996), Nusser et al. (1997) and Dodd (1996). This implementation of the ISUF model for chronic risk assessment is fully described in de Boer et al. (2009).

Model-Then-Add

The traditional approach can be termed the Add-Then-Model approach, because adding over foods precedes the statistical modelling of usual exposure. MCRA offers, as an advanced option, an alternative approach termed Model-Then-Add (van der Voet et al. (2014)). In this approach the statistical model is applied to subsets of the diet (single foods or food groups), and then the resulting usual exposure distributions are added to obtain an overall usual exposure distribution. The advantage of such an approach is that separate foods or food groups may show a better fit to the normal distribution model as assumed in all common models for usual exposure (including MCRA's *betabinomialnormal* (BBN) model and *logisticnormal-normal* model (LNN)). That this principle can work in practice was shown in previous work (de Boer et al. (2009), Slob et al. (2010), Goedhart et al. (2012)) and a simulation model was developed and implemented in MCRA 7.1 to show how multimodal distributions can arise from adding unimodal distributions of foods that are not always consumed (Slob et al. (2010), de Boer and van der Voet (2011)). For specific cases involving separate modelling of dietary supplements and the rest of the diet, proposals have been made (Verkaik-Kloosterman et al. (2011)). However, a practical approach to apply the Model-Then-Add approach to general cases of usual exposure estimation was still missing. Therefore a module in MCRA was developed to implement such an approach based on a visual inspection of a preliminary estimate of the usual exposure distribution using the *Observed Individual Means* (OIM) method.

The Model step

At this stage of development the division of foods into a number of food groups is performed in an interactive process, where the MCRA user is presented with a visual display (see example in Figure 3.33) which shows:

- 1. The OIM distribution represented as a histogram, where each bar shows the frequency of exposures (summed over foods) of individuals in a certain exposure interval; each bar is subdivided according to the contributions of the individual foods contributing to those exposures (left panel Figure 3.33).
- 2. The contributions graph, where each of the bars in the OIM histogram is expanded to 100%. This graph allows a better view of the lower bars in the OIM histogram.

The visual display identifies the nine foods that contribute most to the total exposure; the remaining foods are grouped in a rest category to avoid identification problems because of too many colours (right panel Figure 3.33).



Figure 3.33: Left panel: OIM usual exposure distribution to smoke flavours via the different foods (excluding the zero exposures) in young children; right panel: Contribution of foods to exposures within each bar of the OIM distribution histogram.

The user has now the possibility to select one or more foods and to split these from the main exposure histogram. A separate graph shows the OIM distribution for the split-off food or food group. The graphs for the main group (now called the rest group) are adapted to show the OIM distribution and the contributions for the remaining foods only (see Figure 3.34 upper two panels). This splitting-off can be repeated several times for other foods or food groups. In this way the user can try to obtain foods or food groups that show unimodal OIM distributions. If the result is not what is intended, a food or food group can be added again to the rest group. Per split-off food or food group the usual exposure can be modelled using either BBN or LNN, with a logarithmic or power transformation. The rest group will always be modelled as OIM. It is possible that the rest group is empty, when the total exposure via the different split-off foods and /or food groups is modelled with BBN or LNN.

After a split-off selection has been made, the OIM distribution is summarised in terms of the defined grouping (Figure 3.35), and the usual exposure distribution per split-off food or food group is fitted according to the chosen modelling settings.

The Add step

Consumptions of foods may be correlated. In the traditional Add-Then-Model approach the Add step automatically reflects any correlations that are apparent in the consumptions at the individual-day or individual level. In the Model-Then-Add approach the estimated usual exposure distributions for different foods or food groups have to be combined to assess the total usual exposure. Two approaches are available for this:

- 1. *Model-based approach*: adds independent samples from the usual exposure distribution per food or food group, ignoring any correlations in consumption;
- 2. *Model-assisted approach*: adds the model-assisted, person-specific usual exposure estimates per food or food group, taking correlations in consumptions into account.



Figure 3.34: Result of a selection into two split-off groups and a rest group. The graph bottom left represents the exposure via a food group containing 'Sausage, frankfurter' and 'Sausage, smoked cooked'. The graph bottom right represents the exposure via a food group containing 'Sausage, luncheon meat', Herbs, mixed, main brands, not prepared', 'Soup, pea', 'Ham', and 'Bacon'. The top graph represents the exposure via the rest group.



Usual exposures per model

Figure 3.35: OIM usual exposure distribution showing the contributions from the three food groups as constructed in Figure 3.34.

See also, episodically consumed foods, model-based, model-assisted.

Before the addition is made, in the model-based approach, model-based estimates of the usual exposure amounts distribution per food or food group are back-transformed values from the normal distribution assumed for transformed amounts per food or food group, and the *model-based frequency* distribution is sampled to decide if a simulated individual has exposure via the food or food group or not. Model-assisted estimates of the usual exposure distribution are back-transformed values from a shrunken version of the transformed OIM distribution, also done per food or food group, where the shrinkage factor is based on the variance components estimated using the linear mixed model for amounts at the transformed scale (van Klaveren et al. (2012)). For individuals with no observed exposure (OIM=0) no model-assisted estimate of usual exposure can be made and a model-based replacement is used.

The model-based approach was investigated in Slob et al. (2010) and performed surprisingly well, even if correlations in consumptions of foods were present. The model-assisted approach adds exposures at the individual level, and therefore retains effects of correlations between foods in the usual exposure distribution.

MCRA calculates both the model-based and model-assisted usual intake distributions.



Figure 3.36: Model-assisted estimated usual exposure distributions (excluding the zero exposures).

Chronic exposure as a function of covariates

The intake frequency and transformed intake amounts may be modelled as a function of covariates. MCRA allows one covariable and/or one cofactor.



Figure 3.37: Model-based estimated usual exposure distributions (excluding the zero exposures).

Table 3.191: Intake frequencies and amounts, modelled as a function of covariates.

	Frequencies	Amounts
cofactor	$\mathit{logit}(\pi) = \beta_{0l}$	$transf(y_{ij}) = \beta_{0l} + c_i + u_{ij}$
covariable	$\textit{logit}(\pi) = \beta_0 + \beta_1 f(x_1; df)$	$transf(y_{ij}) = \beta_0 + \beta_l f(x_1; df) + c_i + u_{ij}$
both	$\textit{logit}(\pi) = \beta_{0l} + \beta_1 f(x_1; df)$	$transf(y_{ij}) = \beta_{0l} + \beta_l f(x_1; df) + c_i + u_{ij}$
interaction	$\textit{logit}(\pi) = \beta_{0l} + \beta_{1l} f(x_1; df)$	$transf(y_{ij}) = \beta_{0l} + \beta_{1l}f(x_1; df) + c_i + u_{ij}$

Here $l = 1 \cdots L$ and L is the number of levels of the cofactor, y_{ij} , the intake amount, x_1 is the covariable, f is a polynomial function with the degrees of freedom df, c_i and u_{ij} are the individual effect and interaction effect, respectively. These effects are assumed to be normally distributed $N(0, \sigma_{between}^2)$ resp. $N(0, \sigma_{within}^2)$. The degree of the function is determined by backward or forward selection. In the output, the usual intake is displayed for a specified number of values of the covariable and/or the levels of the cofactor.

Total Diet Study

In Total Diet Studies (TDS), substance occurrence data is obtained from measuring food products as consumed. TDS offers a more direct measure of substance concentrations compared to traditional monitoring and surveillance programs that are concerned with contamination of raw agricultural commodities. In a TDS, food selection is based on national consumption data in such a way that 90 to 95% of the usual diet is represented by the samples. Selected foods are collected, prepared as consumed and related foods are pooled prior to analysis. The compositions these TDS food samples are described by the *TDS food sample compositions* data module.

In MCRA, TDS concentration data can also be used in *dietary exposure assessments*, using it as an alternative type of concentration data where the modelled foods are not the raw primary commodities (RACs), but these are TDS food compositions. To link the concentration data to the consumed foods, the *TDS food sample composition information* is used in the *food conversion algorithm* in a manner analogous to the use of *food recipes* describing the composition of a composite food. The main difference is that the translation proportion is always 100% (default). Take, as an example, a TDS food *FruitMix* that is composed of *apple, orange* and *pear*, then a consumed food (food-as-eaten) *apple-pie* is converted to *apple, wheat* and *butter* (in some specific proportions) and subsequently, *apple* is converted to modelled food *FruitMix* (100%). Not necessarily all foods as consumed are represented in a TDS food sample. In addition to the TDS food sample compositions, there may be additional foods that are not officially part of a TDS food, but which can be extrapolated to a TDS food sample. Through the use of *food extrapolations* (read across translations), these foods may be directly linked to a TDS food sample, e.g., by specifying that *pineapple* is translated to *FruitMix*, *pineapple* or foods containing *pineapple* will also be matched to a *FruitMix* concentration.

Because TDS samples only contain one single, average measurement, TDS occurrence data can currently only be used for only applicable for chronic exposures assessments. However, when variability information is available for

the raw primary foods in the TDS food samples (e.g., from monitoring), this information may be used *to approximate the variance of TDS samples*.

For more information about Total Diet Studies, visit the TDS-Exposure website http://www.tds-exposure.eu.

TDS reduction to limit scenario analysis

The outcome of a MCRA risk assessment may be that some foods dominate the right upper tail of the exposure distribution. A scenario analysis answers the question to what extent the risk of foods with a high exposure would have been diminished by an intervention or by taking any precautions. To be able to do so, some information is needed about the concentration distributions of the raw agricultural commodities (RACs) that make up the TDS food sample. The decision to intervene or not can be based on comparison between the p95 percentile point of the concentration distribution and a concentration limit value that associated with a high risk.

- For p95 ≤ limit, most concentration values are below the value that is considered as a potential risk, so there is no urgency to take any precautions.
- When the opposite is true, i.c. p95 > limit, there may be an argument to intervene for this specific food.

In MCRA, limits and p95's are supplied by the *concentration distributions module*. In the MCRA interface, a scenario analysis can be checked and the scroll down list allows to select the foods that should be included in the scenario analysis. For the selected foods, concentration reduction factors are computed based on the p95 percentile and the limit value:

$$f_{\rm reduction} = limit/p95$$

These reduction factors (computed for the RACs) are applied to the simulated concentrations on the level of the TDS composite foods of which the conversion paths contain the RACs included in the scenario analysis.

$$f_{\rm reduction} \cdot c_{TDS}$$

Here, c_{TDS} is the concentration value of the TDS food.

Substance concentrations generation

Both *chronic* and *acute* dietary exposure assessments rely on assigning substance concentrations to consumed modelled foods. For chronic exposure assessments, this concentration should be the mean concentration of the food x substance combination, as obtained from the concentration models. For acute, these concentrations are obtained through random sampling. For acute, two approaches are available: sample-based and substance-based.

Sample-based concentrations generation

This approach is based on the analytical samples behind the concentration data. For each modelled food, substance concentration values are generated by taking a random sample from the set of analytical samples. For each analytical sample, the corresponding substance concentration values are kept together, maintaining the correlations between the substance concentrations.

Occasionally, one or more substances have censored or missing concentration values. Then, apply imputation first.

For imputation of missing values there are two methods:

- 1. Imputation by zero: all missing values are assumed zero.
- 2. **Imputation using substance-based concentration models:** all missing values are imputed by drawing a concentration value from the substance-based concentration models.

For imputation of censored values, three methods exist:

- 1. Replace by zero: Censored values are imputated by a zero concentration value. This is an optimistic approach.
- 2. Replace by factor times LOR: Each censored value is replaced by a factor f (e.g., 1 or 1/2) times its LOR.
- 3. **Replace by factor times LOD LOQ system:** Non-detects are replaced by f * LOD; non-quantifications are replaced by LOD + f * (LOQ LOD) and factor f is e.g., 1 or 1/2. For f = 0, non-detects are replaced by zero, non-quantifications are replaced by LOD; for f = 1, non-detects are replaced by LOD, non-quantifications are replaced by LOQ.

Substance-based concentrations generation

In this approach, substance concentrations for a given food are drawn independently per substance from the food/substance concentration models.

Processing correction

Concentrations in the consumed food (food as eaten) may be different from concentrations in the modelled food in monitoring programs (typically raw food) due to processing, such as peeling, washing, cooking etc. Concentrations are therefore corrected according to

$$c'_{jhk} = pf_{jhk} \cdot c_{jhk} = \left(\frac{PF_k}{cf_k}\right) \cdot c_{jhk}$$

where c_{jhk} is the concentration of substance k in the food j with processing type h, and where $pf_{jhk} = \frac{PF_{jhk}}{cf_{jhk}}$ is a factor indicating the mass change for a specific combination k of modelled food and processing. The processing correction factor cf_{jhk} is used to correct for the fact that the processing factors PF_{jhk} as commonly available from the input data describe both the effects of chemical alteration and weight change. E.g. for a dried food with a consumption of 100 gram which is translated to 300 gram raw agricultural commodity, the correction factor is 3. Note that the weight change is already included when calculating the consumption amounts of the modelled foods.

Chronic exposure assessment, daily consumed foods

Model based usual intake

Foods are consumed on a daily basis.

For individual i on day j let Y_{ij} denote the 24 hour recall of a food $(i = 1...n; j = 1...n_i)$. In most cases withinindividual random variation is dependent on the individual mean and has a skewed distribution. It is therefore customary to define a one-way random effects model for Y_{ij} on some transformed scale

$$Y_{ij}^* = g(Y_{ij}) = \mu_i + b_i + w_{ij}$$

with $b_i \sim N(0, \sigma_b^2)$ and $w_{ij} \sim N(0, \sigma_w^2)$

Note that b_i represents variation between individuals and w_{ij} represents variation within individuals between days. The mean μ_i may depend on a set of covariate $Z_i = (Z_{i1}, ..., Z_{ip})$:

$$\mu_i = \beta_0 + \beta_1^t Z_i$$

where β_0 and β_1 are regression coefficients.

The usual intake T_i for an individual *i* is defined as the mean consumption over many many days. This assumes that the untransformed intakes Y_{ij} are unbiased for true usual intake rather than the transformed intakes Y_{ij}^* . In mathematical terms T_i is the expectation of the intake for this individual where the expectation is taken over the random day effect:

$$T_i = E_w[g^{-1}(\mu_i + b_i + w_{ij})|b_i] = F(b_i)$$

Model based usual intake on the transformed scale

For the model based usual intake first note that the conditional distribution

$$(\mu_i + b_i + w_{ij}|b_i) \sim N(\mu_i + b_i, \sigma_w^2)$$

It follows that the usual intake T_i is given by

$$T_i = E_w[g^{-1}(\mu_i + b_i + w_{ij}|b_i)] = \int_{-\infty}^{\infty} g^{-1}(\mu_i + b_i + w_{ij}) \frac{1}{\sqrt{2\pi\sigma_w^2}} \exp\left(-\frac{w^2}{2\sigma_w^2}\right) \mathrm{d}w$$

Model based using a logarithmic transformation

For the logarithmic transform the usual intake T_i can be written in closed form using the formula for the mean of the lognormal distribution:

$$T_i = \exp(\mu_i + b_i + \sigma_w^2/2)$$

In this case T_i follows a log-normal distribution with mean $\mu_i + \sigma_w^2/2$ and variance σ_b^2 . This fully specifies the usual intake distribution, e.g. the mean and variance of the usual intake are given by

$$\begin{split} \mu_{iT} &= E[T_i] = \exp(\mu_i + \sigma_w^2/2 + \sigma_b^2/2) \\ \sigma_{iT}^2 &= Var[T_i] = [\exp(\sigma_b^2) - 1] \exp(2\mu_i + \sigma_w^2 + \sigma_b^2) \end{split}$$

Model based using a power transformation

For the *power transformation* the integral can be approximated by means of N-point Gauss-Hermite integration. This results in the following usual intake

$$T_i \approx \frac{1}{\sqrt{\pi}} \sum_{j=1}^N w_j (\mu_i + b_i + \sqrt{2} \sigma_w x_j)^p$$

with p the inverse of the power transformation. A similar approximation can be used for the Box-Cox transformation. There can be a small problem with Gauss-Hermite integration. The summation term $(\mu_i + b_i + \sqrt{2}\sigma_w x_j)^p$ can not be calculated when the factor between round brackets is negative and the power p is not an integer. This can happen when $(\mu_i + b_i)$ is small relative to the between day standard error σ_w . In that case the corresponding term is set to zero. This is not a flaw in the numerical method but in the statistical model since the model allows negative intakes on the transformed scale which cannot be transformed back to the natural scale. The mean and variance of T_i can be approximated again by using Gauss-Hermite integration:

$$\begin{split} \mu_{iT} &= E[T_i] = \frac{1}{\sqrt{\pi}} \sum_{k=1}^N w_k \frac{1}{\sqrt{\pi}} \sum_{j=1}^N w_j (\mu_i + \sqrt{2}\sigma_w x_j + \sqrt{2}\sigma_b x_k) \\ \\ _{iT} &= Var[T_i] = \frac{1}{\sqrt{\pi}} \sum_{k=1}^N w_k \left[\frac{1}{\sqrt{\pi}} \sum_{j=1}^N w_j (\mu_i + \sqrt{2}\sigma_w x_j + \sqrt{2}\sigma_b x_k) \right]^2 - \mu_T^2 \end{split}$$

An alternative method for obtaining model based usual intakes for the power transformation employs a Taylor series expansion for the power, see e.g. Kipnis et al. (2009). This is however less accurate than Gauss-Hermite integration. For the power transformation simulation is required to derive the usual intake distribution: simulate a random effect b_i for many individuals and then approximate T_i for these individuals. The T_i values then form a sample form the usual intake distribution.

Model assisted usual intake on the transformed scale

The model assisted approach employs a prediction for the usual intakes of every individual in the study. This requires a prediction of the individual random effect b_i for every individual.

In the one-way random effects model the Best Linear Unbiased Prediction for $(\mu_i + b_i)$ is given by

$$\textit{BLUP}_i = \mu_i + (\bar{Y}^*_i - \mu_i) \left(\frac{\sigma_b^2}{\sigma_b^2 + \sigma_w^2/n_i} \right)$$

in which \bar{Y}_i^* is the mean of the transformed intakes for individual *i*. BLUPs have optimal properties for some purposes, but not for the purpose of representing the variation σ_b^2 between individuals. This can be seen by noting that

$$Var(\bar{Y}_i^*) = \sigma_b^2 + \sigma_w^2 / n_i$$

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and thus

$$\textit{Var}(\textit{BLUP}_i) = \left(\frac{\sigma_b^4}{\sigma_b^2 + \sigma_w^2/n_i}\right)$$

which is smaller than the between individual variance σ_b^2 . As an alternative a modified BLUP can be defined by means of

$$\textit{modifiedBLUP}_i = \mu_i + (\bar{Y}_i^* - \mu_i) \sqrt{\left(\frac{\sigma_b^2}{\sigma_b^2 + \sigma_w^2/n_i}\right)}$$

which has the correct variance σ_b^2 and also the correct mean μ_i . However these optimal properties disappear when modified BLUPs are directly backtransformed to the original scale.

Model assisted using a logarithmic transformation

For the logarithmic transformation the usual intake T_i follows a log-normal distribution with mean $\mu_i + \sigma_w^2/2$ and variance σ_b^2 . If we can construct a BLUP like stochastic variable with the same mean and variance, then this variable be an unbiased predictor with the correct variance. It is easy to see that the following variable has the same distribution as T_i

$$\textit{modelassistedBLUP}_{i} = \mu_{i} + \frac{\sigma_{w}^{2}}{2} + (\bar{Y}_{i}^{*} - \mu_{i}) \sqrt{\left(\frac{\sigma_{b}^{2}}{\sigma_{b}^{2} + \sigma_{w}^{2}/n_{i}}\right)}$$

So the model assisted individual intake $\exp(modelassistedBLUP_i)$ has the same distribution as the usual intake and is thus the best predictor for usual intake.

Kipnis et al. (2009) employs the conditional distribution of b_i given the observations Y_{i1}, \dots, Y_{in_i} to obtain a prediction. First note that

$$(b_i|Y_{i1},\cdots,Y_{in_i})=(b_i|Y_{i1}^*,\cdots,Y_{in_i}^*)=(b_i|\bar{Y}_i^*)$$

Since all distributions in the one-way random effects model are normal it follows that:

$$(b_i, \bar{Y}_i^*) \sim BivariateNormal(0, \mu_i, \sigma_h^2, \sigma_h^2 + \sigma_w^2/n_i, \sigma_h^2)$$

where the last parameter represents the covariance between b_i and \bar{Y}_i^* . It follows that the conditional distribution

$$(b_i | \bar{Y}_i^*) \sim N(\mu_c, \sigma_c^2)$$

with

$$\mu_c = \frac{\sigma_b^2}{\sigma_b^2 + \sigma_w^2/n_i} (\bar{Y}_i^* - \mu_i)$$

and

$$\sigma_c^2 = \frac{\sigma_b^2 \sigma_w^2/n_i}{\sigma_b^2 + \sigma_w^2/n_i}$$

A prediction for the usual intake $T_i = F(b_i)$ is then obtained by the expectation

$$E[F(b_i)|\bar{Y}_i^*] = \int F(b)\phi(b;\mu_c,\sigma_c^2)\mathrm{d}b$$

For the logarithmic transform $F(b_i) = \exp(\mu_i + b_i + \sigma_w^2/2)$ and the expectation reduces to

$$E[F(b_i)|\bar{Y}_i^*] = \exp(\mu_i + \mu_c + \sigma_c^2/2 + \sigma_w^2/2)$$

which is a function of \bar{Y}_i^* through μ_c . To obtain the mean and variance of the prediction note that

$$\mu_i + \mu_c + \sigma_c^2/2 + \sigma_w^2/2 \sim N\left(\mu_i + \frac{\sigma_b^2 \sigma_w^2/n_i}{2(\sigma_b^2 + \sigma_w^2/n_i)} + \frac{\sigma_w^2}{2}, \frac{\sigma_b^4}{\sigma_b^2 + \sigma_w^2/n_i}\right)$$
It follows that the expectation of the prediction equals

$$\begin{split} E[E[F(b_i)|\bar{Y}_i^*]] &= \exp\left(\mu_i + \frac{\sigma_b^2 \sigma_w^2/n_i}{2(\sigma_b^2 + \sigma_w^2/n_i)} + \frac{\sigma_w^2}{2} + \frac{\sigma_b^4}{2(\sigma_b^2 + \sigma_w^2/n_i)}\right) \\ &= \exp\left(\mu_i + \frac{\sigma_b^2}{2} + \frac{\sigma_w^2}{2}\right) \end{split}$$

which equals the mean of the usual intake. However the variance of the prediction equals

$$Var[E[F(b_i|\bar{Y}^*_i]] = \left[\exp\left(\frac{\sigma_b^4}{(\sigma_b^2 + \sigma_w^2/n_i)}\right) - 1\right]\exp(2\mu_i + \sigma_b^2 + \sigma_w^2)$$

Which is less than the variance of the usual intake. The approach of Kipnis et al. (2009) will therefor result in too much shrinkage of the model assisted usual intake.

Model assisted using a power transformation

For the *power transformation* a model assisted BLUP with optimal properties, as derived above, cannot be constructed. The approach of Kipnis et al. (2009) can however be used to obtain a prediction in the following way. First approximate $T_i = F(b_i)$ by *Gauss-Hermite integration*:

$$F(b_i) = T_i \approx \frac{1}{\sqrt{\pi}} \sum_{j=1}^N w_i (\mu_i + b_i + \sqrt{2} \sigma_w x_i)^p$$

Secondly again use Gauss-Hermite to approximate the expectation of the conditional distribution giving the prediction P_i .

$$P_i = E[F(b_i)|\bar{Y}_i^*] = \int F(b_i)\phi(b;\mu_c,\sigma_c^2) \mathrm{d}b \approx \frac{1}{\pi} \sum_{k=1}^N w_k \sum_{j=1}^N w_j (\mu_i + \mu_c + \sqrt{2}\sigma_w x_j + \sqrt{2}\sigma_c x_k)^p$$

which is a function of Y_i^* through μ_c . It is likely that the thus obtained predictions P_i have a variance that is too small. If we would know the mean μ_{iP} and variance σ_{iP}^2 of the predictions, the predictions could be linearly rescaled to have the correct mean μ_{iT} and variance $\frac{2}{iT}$. The mean and variance of the prediction can be calculated using *Gauss-Hermite integration*.

$$\mu_{iP} = \frac{1}{\sqrt{\pi}} \sum_{l=1}^{N} w_l \frac{1}{\pi} \sum_{k=1}^{N} w_k \sum_{j=1}^{N} w_j (\mu_i + \sqrt{2} \frac{\sigma_b^2}{\sigma_b^2 + \sigma_w^2/n_i} x_l + \sqrt{2} \sigma_w x_j + \sqrt{2} \sigma_c x_k)^p$$

$$\sum_{iP}^{2} = \frac{1}{\sqrt{\pi}} \sum_{l=1}^{N} w_l \left[\frac{1}{\pi} \sum_{k=1}^{N} w_k \sum_{j=1}^{N} w_j (\mu_i + \sqrt{2} \frac{\sigma_b^2}{\sigma_b^2 + \sigma_w^2/n_i} x_l + \sqrt{2} \sigma_w x_j + \sqrt{2} \sigma_c x_k)^p \right]^2 - \mu_{iP}^2$$

The proposed prediction then equals

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$$P_i^* = \mu_{iT} + \frac{\sigma_{iT}}{\sigma_{iP}}(P_i - \mu_{iP})$$

Chronic exposure assessment, episodically consumed foods

For episodically consumed foods we need to take the probability of consumption into account. Define p_i as the probability that individual *i* consumes the food on any given day. The usual intake for this individual is then given by the product of p_i and T_i which is now defined as the usual amount on consumption days. Since individuals will vary in their probability pi, besides modelling the amounts as for daily consumed foods, it is also necessary to model the frequency of consumption. A three stage analysis of 24-hour recall data is the necessary:

- 1. A model for the frequency of consumption
- 2. A model for the intakes on consumption days
- 3. Integration of both models in order to obtain a usual intake distribution.

Step 2 uses the analysis outlined in the previous section for the positive intakes only. For step 1 two popular models which describe between-individual variation for the probability of consumption are the beta-binomial model and the logistic-normal model.

Beta-Binomial model for frequencies (BBN)

Let n_i be the total number of recall days for individual i and X_i the number of days with a positive intake. The distribution of X_i , with p_i the probability of consumption for individual i, is given by

$$X_i = Binomial(n_i, p_i)$$

In this model the probability p_i varies among individuals according to the Beta distribution:

$$f(p)=B^{-1}(\alpha,\beta)p^{\alpha-1}(1-p)^{\beta-1}$$

with

$$B(\alpha, \beta) = \frac{\Gamma(\alpha)\Gamma(\beta)}{\Gamma(\alpha + \beta)}$$

Combining the binomial and the Beta distribution results in the betabinomial distribution:

$$P(X_i=x) = \left(\begin{array}{c} n_i \\ r \end{array} \right) \frac{B(\alpha+x,n_i+\beta-x)}{B(\alpha,\beta)}$$

The mean and variance of the betabinomial distribution are given by

$$E[X_i] = n_i \frac{\alpha}{\alpha + \beta}$$

and

$$Var[X_i] = n_i \frac{\alpha\beta(\alpha + \beta + n_i)}{(\alpha + \beta)^2(\alpha + \beta + 1)}$$

Using the reparameterization $\pi = \alpha/(\alpha + \beta)$ and $\phi = 1/(\alpha + \beta + 1)$, it follows that

$$E[X_i] = n_i \pi$$

and

$$Var[X_{i}] = n_{i}\pi(1-\pi)[1+(n_{i}-1)\phi]$$

This reparameterization enables to model the probability π_i of consumption for individual *i* directly as a logistic regression:

$$logit(\pi_i) = \gamma_0 + \gamma_1^t Z_i$$

Note that the dispersion parameter ϕ : is assumed to be equal for all individuals. The betabinomial logistic regression model can be fitted by means of maximum likelihood.

Model based frequencies for usual intake

For the model based usual intake distribution the estimated parameters π_i and ϕ are backtransformed using $\alpha_i = \pi_i \phi/(1-\phi)$ and $\beta_i = (1-\pi_i)\phi/(1-\phi)$. These can then be used to draw from the Beta distribution.

Model assisted frequencies for usual intake

For the model assisted usual intake distribution a prediction of the consumption probability is required for every individual. Simple predictions are

- 1. the observed frequencies for every individual or
- 2. the fitted probability for every individual. When there are no covariables the fitted probability is the same for every individual.
- 3. Alternatively one can use the approach outlined in Kipnis et al (2009) employing the conditional expectation of the probability given the observed frequency:

$$\begin{split} E(p_i|X_i = x) &= \int_p pf(p|X_i = x)\mathrm{d}p \\ &= \int_p p\frac{f(X_i = x|p)f(p)}{\int f(X_i = x|p)f(p)\mathrm{d}p}\mathrm{d}p \\ &= \frac{1}{P(x_i = x)}\int_p p\left(\begin{array}{c}n_i\\r\end{array}\right)p^x(1-p)^{n_i-x}B^{-1}(\alpha_i,\beta_i)p^{\alpha_i-1}(1-p)^{\beta_i-1}\mathrm{d}p \\ &= \frac{B^{-1}(\alpha_i,\beta_i)}{P(x_i = x)}\left(\begin{array}{c}n_i\\r\end{array}\right)\int_p p^{\alpha_i+x}(1-p)^{n_i+\beta_i-x-1}\mathrm{d}p \\ &= \frac{B(\alpha_i + x + 1, n_i + \beta_i - x)}{B(\alpha_i + x, n_i + \beta_i - x)} \\ &= \frac{\alpha_i + x}{\alpha_i + \beta_i - x} \end{split}$$

For individual with zero intakes on all recall days a prediction for the random individual amount effect b_i is not available. There seem to be two option for predicting the usual intake for such individuals:

- Set the individual intake to zero
- Simulate a model based prediction for the amount and combine this with the conditional expected probability given above to obtain an individual usual intake.

Logistic-Normal model for frequencies (LNN0)

In this model the distribution of X_i is again binomial:

$$X_i = Binomial(n_i, p_i)$$

The probability p_i is now given by a logistic regression with a random effect in the linear predictor which represents the between-individual variation in the probability p_i

 $\textit{logit}(p_i) = \lambda_i + v_i \text{ with } v_i \sim N(0, \sigma_v^2) \text{ and the regression equation } \lambda_i = \gamma_0 + \gamma_1^t Z_i$

The marginal probability π_i is obtained by integrating over the random effect v_i , i.e. using Gauss-Hermite integration

$$\pi_i = \int H(\lambda_i + v) f(v) dv \approx \frac{1}{\sqrt{\pi}} \sum_{j=1}^N w_j H(\lambda_i + \sqrt{2}\sigma_v x_j)$$

in which H() is the inverse of the logit transformation. Note that this is different from $logit^{-1}(\lambda_i)$ which is the median probability. The model can be fitted by maximum likelihood using Gauss-Hermite integration. An (approximate) maximum likelihood procedure is implemented in routine glmer of the lme4 package in R. For a new vector of covariates Z_i^* the linear predictor λ_i^* can be calculated along with its standard error $Se(\lambda_i^*)$. The marginal predicted probability π_i^* can be calculated by means of Gauss-Hermite integration and the standard error of the predicted probability can be calculated by means of the usual Taylor series expansion:

$$\begin{split} & Se(\pi_i^*) \approx \frac{Se(\lambda_i^*)}{\sqrt{\pi}} \sum_{j=1}^N w_j \frac{d}{d\lambda_i^*} H(\lambda_i^* + \sqrt{2}\sigma_v x_j) \\ &= \frac{Se(\lambda_i^*)}{\sqrt{\pi}} \sum_{j=1}^N w_j H(\lambda_i^* + \sqrt{2}\sigma_v x_j) [1 - H(\lambda_i^* + \sqrt{2}\sigma_v x_j)] \end{split}$$

Model based frequencies for usual intake

For the model based usual intake distribution the estimated parameters λ_i and σ_v^2 can be used to generate individual probabilities.

Model assisted frequencies for usual intake

For the model assisted usual intake distribution simple predictors are (a) the observed frequencies and (b) the marginal probability π_i . The conditional expectation (c) is given by

$$\begin{split} E(p_i|X_i = x) &= \int_v H(\lambda_i + v)f(v|X_i = x)\mathrm{d}v \\ &= \int_v H(\lambda_i + v)\frac{f(X_i = x_i|v)f(v)}{\int f(X_i = x_i|v)f(v)\mathrm{d}v}\mathrm{d}v \\ &= \frac{\int_v H(\lambda_i + v)[H(\lambda_i + v)]^{x_i}[1 - H(\lambda_i + v)]^{n_i - x_i}f(v)\mathrm{d}v}{\int_v [H(\lambda_i + v)]^{x_i}[1 - H(\lambda_i + v)]^{n_i - x_i}f(v)\mathrm{d}v} \end{split}$$

and both nominator and denominator can be approximated by means of the *Gauss-Hermite integration*. For individual with zero intakes on all recall days see above for the two options.

Logistic-Normal model for frequencies correlated with amounts (LNN)

This model is extends the LNN0 model with a correlation between the individual random effect b_i for amounts and the individual random effect v_i for frequencies. This model is also known as the NCI model and is introduced by Tooze et al. (2006) with further mathematical details in Kipnis et al. (2009). The model can be written as

$$\textit{logit}(P(Y_{ij} > 0)) = \lambda_i + v_i$$

$$g(Y_{ij}) = \mu_i + b_i + w_{ij}$$

and $(v_i, b_i) \sim \textit{BivariateNormal}(0, 0, \sigma_v^2, \sigma_b^2, \rho)$ and $w_{ij} \sim N(0, \sigma_w^2)$

The model can be fitted by maximum likelihood employing two-dimensional Gauss-Hermite integration.

Model based usual intake

Model based usual intake requires generation of the pair (v_i, b_i) for many hypothetical individual. The usual intake U_i for such a hypothetical individual is then given by

$$\begin{split} U_i &= H(\lambda_i + \nu_i)T_i \\ &= H(\lambda_i + \nu_i)E_w[g^{-1}(\mu_i + b_i + w_{ij})|b_i] \\ &= H(\lambda_i + \nu_i)F(b_i) \end{split}$$

The second term can be calculated using the method outlined for daily intakes.

Model assisted usual intake

This requires simultaneous prediction of the random effect for frequency and for amount as outlined in Kipnis et al. (2009). We have for individual *i* in the study $(U_i|Y_{i1}, \dots, Y_{in_i}) = (U_i|Y_{i1}^*, \dots, Y_{in_i}^*) = (U_i|x_i, \bar{Y}_i^*)$ where x_i is the number of positive intakes and \bar{Y}_i^* is the mean of the transformed **positive** intakes. It follows that the required conditional expectation P_i equals

$$\begin{split} P_i &= E[U_i|x_i,\bar{Y}_i^*] \\ &= E_{v_i,b_i}[H(\lambda_i+v_i)F(b_i)|x_i,\bar{Y}_i^*] \\ &= \frac{\int \int H(\lambda_i+v_i)F(b_i)f(x_i,\bar{Y}_i^*|v_i,b_i)\phi(v_i,b_i)dv_idb_i}{\int \int f(x_i,\bar{Y}_i^*|v_i,b_i)\phi(v_i,b_i)dv_idb_i} \end{split}$$

where

$$f(x_i, \bar{Y}_i^* | v_i, b_i) = [H(\lambda_i + v_i)]^{x_i} [1 - H(\lambda_i + v_i)]^{n_i - x_i} \phi(\bar{Y}_i^* - \mu_i - b_i; 0, \sigma_w^2 / x_i)$$

Both nominator and denominator can be approximated by a *two-dimensional Gauss-Hermite integration*. Note that for the log-transform $F(b_i) = T_i = \exp(\mu_i + b_i + \sigma_w^2)/2)$ can be calculated exactly; for the *power transformation* an approximation must be used. It can be expected that the predicted usual intake will not have the correct variance. This can possibly be remedied by equating the mean and variance of U_i and P_i . These are however rather involved to calculate.

For individual with zero intakes on all recall days the model assisted usual intake can be set to zero, or can be simulated as follows

- 1. Calculate the Model assisted frequency P_0 for usual intake (see LNN0)
- 2. Transform P_0 back to the logistic scale, i.e. $L_0 = logit(P_0)$. Get the conditional distribution of

$$(b|v=L_0-\lambda_i) \sim N\left(\frac{\sigma_b}{\sigma_v}\rho(L_0-\lambda_i),(1-\rho^2)\sigma_b^2\right)$$

3. Simulate a draw b_0 from this conditional distribution and obtain the usual intake as $P_0 \exp(\mu_i + b_0 + \sigma_w^2)$

Note that the backtransformation from P_0 to L_0 is according to the median of the distribution rather than the mean.

Dietary exposures settings

Selection settings

Name Type		Description			
Scenario analysis foods AlphaNumeric		The foods of interest for the scenario analysis.			

Calculation settings

Name	Туре	Description
Selected tier	SettingsTemplateType	Specifies all module settings should be set according to a pre-defined tier or using custom settings.
Exposure type	ExposureType	The type of exposure considered in the assessment; acute (shor term) or chronic (long-term).
Total diet study concentration data	Boolean	Specifies whether exposure is based on sampling data from tota diet studies.
Multiple substances analysis	Boolean	Specifies whether the assessment involves multiple substances.
Compute cumulative exposures	Boolean	Specifies whether the assessment involves multiple substances a results should be cumulated over all substances.
Sample based	Boolean	Include co-occurrence of substances in samples in simulations. checked, substance residue concentrations are sampled using th correlations between values on the same sample. If unchecked, any correlation between substances is ignored, substance residue concentrations are sampled ignoring the correlations between values on the same sample.
Consumptions on the same day come from the same sample	Boolean	If checked, in procedure of EFSA Guidance 2012, section 4.1. all consumptions of a raw commodity of an individual on the same day are assumed to come from the same sample. If unchecked, all consumptions of a raw commodity of an individ on the same day are assumed to come from different samples.

Name	Туре	Description
Maximise co-occurrence of high values in simulated samples	Boolean	Within each pattern of substance presence. If checked, substant residue concentrations are sorted within co-occurrence patterns substances on the same samples. After sorting, high residue val occur more frequently on the same sample. This choice is conservative. If unchecked, substance residue concentrations are sampled at random, ignoring any co-occurrence patterns of substances on the same samples. This choice is less conservative
Apply processing factors	Boolean	Specified in table ProcessingFactor. If checked, processing fact are applied. Concentrations in the consumed food may be different from concentrations in the modelled food in monitorin programs (typically raw food) due to processing, such as peelin washing, cooking etc. If unchecked, no processing information used. This is in most (though not all) cases a worst-case assumption
Use distribution	Boolean	Probabilistic specifications of processing factors will be used
Ignore processing factors less than 1	Boolean	This setting will suppress the use of processing factors lower th 1 (it is used in the EFSA 2012 Pessimistic tier).
Default processing factor for missing processing types (except foods with processing type equal to 'Unspecified' (default = 1))	Numeric	Specify a processing factor for missing processing types. When the processing type is 'Unspecified', the default processing factor is equal to 1.
Use default processing factor for missing processing types	Boolean	Use default processing factor for missing processing types
Perform MCR analysis	Boolean	Perform a Maximum Cumulative Ratio (MCR) analysis to determine co-exposure between substances.
Substance weighting in mixtures	ExposureApproachType	Risk based: exposures in equivalents of the reference substance standardised: standardised exposures per substance have varian 1; or unweighted exposures: RPFs are equal to 1.
Use unit variability	Boolean	Controls whether to use unit variability.
Unit variability model	UnitVariabilityModelType	Describes variation between single units when concentration da are from composite samples.
Estimates nature	EstimatesNature	Simulated unit concentrations can be higher or lower than composite value (realistic) or only equal or higher (conservative
Unit variability parameter	UnitVariabilityType	Use Coefficient of variation or Variability factor, specified in VariabilityFactor table.
Mean of LogNormal simulated values (biasing)	Mean ValueCorrection Type	Unbiased: correct unit simulations for difference between medi and mean.
Default variability factor for unit weight <= 25g	Numeric	Default variability factor 1 (unit weight <= 25 g, small crops). S requires specification of unit weight (FoodProperties table) and case of beta model, also the Number of units in a composite sample (UnitVariability table).
Correlation between substances on the same units	UnitVariabilityCorrelationType	Specifies the type of correlation between substances on the samunits; no correlation or full correlation.
Default variability factor for unit weight > 25g	Numeric	Default variability factor 5 (unit weight > 25 g, medium/large crops). Still requires specification of unit weight (FoodPropertitable) and, in case of beta model, also the Number of units in a composite sample (UnitVariability table).
Model type	IntakeModelType	The parametric model for between-and within-individual variation, and possibly covariates.
Model-then-add	Boolean	Specifies whether to create separate exposure models for specific groups of modelled foods (model-then-add).
Covariate modelling	Boolean	Specifies whether to model exposures as a function of covariate at individual level.

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Name	Туре	Description
Model-then-add sub-category models	IntakeModelPerCategory	Sub-model specifications for foods groups that should be model separately.
Grid precision frequency model	Numeric	The discrete frequency distribution (ISUF) is approximated via number of classes.
Number of iterations (x 1000)	Numeric	The number of iterations that is used to estimate the discrete frequency distribution for the ISUF model.
Spline-fit	Boolean	To achieve a better normality, a second transformation is performed: a spline function is fitted to the logarithmically or power transformed data as a function of the normal Blom score
Amount model covariate model	CovariateModelType	Specifies whether, and how to model exposures amounts as function of covariates.
Function	FunctionType	Functional relation between exposure and covariable.
Transformation	TransformType	The data transformation used to approximate normality for amounts.
Testing level	Numeric	Significance level for testing the degree of the function. e.g., 0.
Testing method	TestingMethodType	Starting from a full model (backward) or empty model (forward
Maximum degrees of freedom	Numeric	Order of function. Determines the maximum degree of complexity of the function.
Minimum degrees of freedom	Numeric	Order of function. Determines the minimum degree of complexity of the function.
Amounts model: variance ratio (between/within)	Numeric	Estimate of the ratio of the variance components of the amount model (only relevant for data with only 1 day per individual)
Frequency model covariates model	CovariateModelType	Specifies whether, and how to model exposure frequency as function of covariates.
Function	FunctionType	Functional relation between exposure and covariable.
Testing level	Numeric	Significance level for testing the degree of the function. e.g., 0.
Testing method	TestingMethodType	Starting from a full model (backward) or empty model (forward
Minimum degrees of freedom	Numeric	Order of function. Determines the minimum degree of complexity of the function.
Maximum degrees of freedom	Numeric	Order of function. Determines the maximum degree of complexity of the function.
Frequency model dispersion	Numeric	Frequency model dispersion estimate for (only relevant for data with only 1 day per individual).
Use occurrence patterns for generating simulated samples	Boolean	When selected, this simulated samples will be based on occurrence patterns.
Associate the unspecified percentage with no-occurrence for foods with at least one specified occurrence pattern	Boolean	If checked, for foods with at least one specified occurrence pattern, unspecified occurrence patterns for the same food are assumed to be associated with no use. If unchecked, all substances are considered to be authorised (potentially present samples). Note that this setting cannot be used for foods that ha no specified AUs. These foods have 100% potential presence o all substances. To declare all AUs on such a food un-authorised include an empty AU with percentage 100% in the AU data tak (i.e., use an AU for this food, without specifying substances in f AU Substances table)
Details level dietary exposures	DietaryExposuresDetailsLevel	Level of detail for summarizing dietary exposure/intakes.
Iterate survey	Boolean	Instead of (re-)sampling the individual days, loop over the entir survey (= 1 iteration). The number of iterations for a survey is calculated as round (number of Monte Carlo iterations /(number of individuals * surveys days)).
Monte Carlo iterations	Numeric	The number of iterations for Monte Carlo simulations, e.g. 100,000 (maximum is 100,000).
Impute exposure distributions	Boolean	Impute exposure distributions for substances with missing concentrations

Table 3.193 - continued from previous page

Name	Туре	Description
Include diagnostics analysis for variability	Boolean	For each percentile the variability (standard deviation) of the estimated percentiles versus sample size are plotted.
Cofactor name	AlphaNumeric	Specify the name of the cofactor.
Covariable name	AlphaNumeric	Specify the name of the covariable.
Allow conversion using food extrapolations	Boolean	Try to find read across codes. If unchecked, read across table is ignored, default is 'Use read across info'. E.g. for pineapple no measurements are found but by specifying that pineapple is converted to FruitMix (with a default proportion of 100%), the TDS sample concentration value of FruitMix will be used for pineapple (as-eaten or as ingredient).
Censored values replacement	NonDetectsHandlingMethod	How to replace censored values (when not co-modelled, as in censored models).
Default concentration model	<i>ConcentrationModelType</i>	The concentration model type that will be used as default for al food/substance combinations. If this model type cannot be fitte e.g., due to a lack of data, a simpler model will be chosen automatically as a fall-back.
Apply reduction-to-limit scenario	Boolean	Total diet study: specify reduction-to-limit scenario. If unchecked, all residue values are taken as such (base scenario: apply no reduction factors). If checked, reduction factors are applied for selected foods. Select foods where a reduction is assumed (only foods with Percentile > Limit are shown). Only foods with reduction factors > 1 (percentile / limit) are shown.
Cutoff MCR	Numeric	For selection of individual(day) exposures with maximum cumulative ratio (MCR = total exposure/maximum) above the cutoff.
Cutoff percentage in population ranked on total exposure	Numeric	For selection of individual(day) exposures above the cutoff percentage in the set of individual(day)s ranked on total exposu
Display ratio total exposure/ maximum (in MCR plot)	Numeric	For MCR plot: specify ratio total exposure/ maximum for individual(day) exposures.
Show tail percentiles (MCR	Numeric	Give specific percentiles of exposure distribution (%), e.g. 97.5
plot) for:		99 (space separated).
Set minimum percentage contribution per substance to the tail exposure (MCR plot)	Numeric	Set minimum percentage contribution per substance to the tail exposure.
Target level	TargetLevelType	Select whether the assessment is at the level of external dose, internal (target) concentration or internal (systemic) dose. For a aggregate assessment, combining dietary and nondietary exposures, the target level is always internal concentration or internal (systemic) dose. When only dietary exposures are available, the target level is optional and may be an external (administered) dose, an internal (tissue) concentration or an internal (absorbed) dose.
Seed for pseudo-random number generator	Numeric	A value of 0 will use a pseudo-random seed in each run, a value 0 will provide the same results in a repeated run.
DIET: Population alignment method diet exposures	PopulationAlignmentMethod	Population aligment methods for dietary exposure sources.

Table	3.193 -	continued	from	previous	page
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Output settings

Name	Туре	Description
Exclude privacy sensitive data from outputs	Boolean	Use this setting to not report the parts of the results (i.e., figure tables, or sections) that are marked as (potentially) privacy sensitive.
Include drill-down on 9 individuals around specified percentile	Boolean	Specifies whether drilldown on 9 individuals is to be included in the output.
Show percentiles for	Numeric	Give specific percentiles of exposure distribution (%), e.g. 50 9 95 97.5 99 (space separated).
Percentage for drilldown	Numeric	Gives detailed output for nine individuals near this percentile of the exposure distribution.
Percentage for upper tail	Numeric	Gives detailed output for this upper percentage of the exposure distribution.
Show % of population below level(s)	ExposureMethod	This setting is used for reporting the percentages of individuals (chronic) or individual days (acute) exceeding certain exposure levels. These exposure levels can be generated automatically based on the observed exposures (Automatic, default) or specific explicitly (Manual).
Exposure levels	Numeric	Specify exposure levels for which to give the percentage of exposure below these levels, e.g. 1 10 50 100 200 500.
Number of levels of covariable to predict exposure	Numeric	Specify the number of levels, e.g. 20. The range of the covaria is divided by the number of levels: range = (max - min)/levels. For these covariable levels exposures are predicted.
Predict exposure at extra covariable levels	Numeric	Specify specific prediction levels in addition to the automaticall generated prediction levels (space separated).
Lower percentage for variability (%)	Numeric	The default value of 25% may be overruled.
Upper percentage for variability (%)	Numeric	The default value of 75% may be overruled.
Report consumptions and exposures per individual instead of per kg body weight	Boolean	Specifies whether body weights should be ignored and consumptions and exposures should be expressed per individual Otherwise, the consumptions and exposures are per kg body weight.

Table 3.194: Output settings for module Dietary exposures.

Uncertainty settings

Name	Туре	Description
Resample imputation exposure distributions	Boolean	Specifies whether to resample the imputated exposure distributions.
Resample portion sizes	Boolean	Specifies whether portion sizes should be resampled based on food consumption quantification data, see (Souverein et al. 201
Perform uncertainty analysis	Boolean	In probabilistic risk assessment of dietary exposure, distributio describe the variability in consumption within a given population of individuals and the variability of the occurrence and level of substances in the consumed foods. However, these calculations not consider the amount of uncertainty that is due to the limited size of the underlying datasets.
Iterations uncertainty analysis	Numeric	Specifies the number of uncertainty cycles (default 100).
Monte Carlo iterations per uncertainty run	Numeric	Specifies the number of Monte Carlo iterations in each uncertainty cycle (default 10,000).
Lower uncertainty limit (%)	Numeric	Percentage lower bound, e.g. 2.5%.
Upper uncertainty limit (%)	Numeric	Percentage upper bound, e.g. 97.5%.

Table 3.195: Uncertainty settings for module Dietary exposures.

Dietary exposures tiers

Overview

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tie
C su ti or th sa co fr th sa sa p	false	false	false	false	false	true	true	false	false	fals
U uu va al it	true		true		false	true		true		true
M th ac	false	false	false	false	false		false	false	false	fals

Table 3.196: Tier overview for module Dietary exposures

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes-	EFSA 2012 Chronic Pes-	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
						simistic	simistic			
C va at m el	false	false	false	false	false	false	false	false	false	false
It ei at su	false	false	false	false	false	false	false	false	false	false
R p c c s u ti a c p s u ti d i r d i r d i r d i r d i r d i r s u s v s u s v s u s v s v s v s v s v	false	false	false	false	false	false	false	false	false	false
Ig ne sa pl w	true	true	true	true				true	true	true
E cl in di vi u al w le th N di	false	true	false	true				false	true	false

Table 3.196 - continued from previous page

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N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
T ta d st co co tr ti d	false	false	false	false	false		false	false	false	false
F tee sa p c c c c irr th c c c c t r th t ti t ti t ti t t t t t e : c c c c c c c c c c c c c c c c c c	false	false	false	false		false	false	false	false	false
U su st co vo si ru	true	true	true	true				true	true	true
S st co vo si n	UseMost- Toxic	UseMost- Toxic	DrawRan- dom	DrawRan- dom				UseMost- Toxic	UseMost- Toxic	Drav dom

Table 3.196 - continued from previous page

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
R ta a lc c s l c s l c s l c s l c s l c s l c s l c s l c c s l c s l c c s l c c s l c c s l c c s l c c s l c c s l c c s l c c s l c c s l c c s l c c s l c c s l c c s l c c s l c c s l c c s l c c s l c s l c c s l c s l c c s l c s l c c s l c s s l c s s l c s s l c s s l c s s s l c s s s s	true	true	true	true				true	true	true
A construction for start for start for start start start start v v start statttatt	false	false	true	true				false	false	true
F d p c s t s t a l c c t t i t i t c c s t s t s t t c c s t s t s t s t	false	false	false	false				false	false	false

_						Tabl	e 3.196 – co	ntinued from	previous pag	ge
	N EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202: Acu Tier
	U true ez tr o- la ti ru	true	true	true				false	false	false
	T 10 ol fc e: tr o- la ti	10	10	10						
	R true st c: tr o- la ti tc ec N	true	true	true						
	R true st c: tr o- la ti tc au tr tr i u	true	true	true						
	Ir true p w te co co tr ti	true	true	true				true	true	true

	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
tt cc tt tt	Ø 0.1 e co r i i	0.1	0.05	0.05				0.1	0.1	0.05
l s t t i i l t t t t t t t s s s	R true st v e r n a i i c t t t t t	true	true	true				true	true	true
] s t i i t t t t t	t false st e r i a i t t t t	false	false	false				false	false	false
1ssvtiiiittttssss	R false	false	false	false				true	true	true

						Tubi	0.100 00		provious pu	ge
N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202: Acu Tier
A pl pr ce in fa	true	true	true	true	true	true	true	true	true	true
U di tr bi ti	false	false	false	false	false	false	false	false	false	false
Ig no pi ce in fa to le th 1	false	false	false	false	false	true	true	false	false	false
D fa co tr ti	Empirical	Empirical	Empirical	Empirical	Empirical	NonDe- tect- SpikeLog- Normal	NonDe- tect- SpikeLog- Normal	Empirical	Empirical	Emp
Ir cl M fa ba m	false	false	false	false	false	true	true	false	false	false
R st L in pu ta tiv to au th ri	false	false	false	false		false	false	false	false	false

Table 3.196 - continued from previous page

Ν	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
C S ⁱ V U r ⁱ P n	Replace- ByLOR	Replace- ByLOR	Replace- ByLOR	Replace- ByLOR	Replace- ByZero	Replace- ByLOR	Replace- ByLOR	Replace- ByLOR	Replace- ByLOR	Rep ByL
F tu f f (() x x L o f f x L L t f r tu c f f f tu f f f f f f f f f f f f f f f	0.5	0.5	0.5	0.5		1	1	0.5	0.5	0.5
S p b	true	true	true	true	true	true	true	true	true	true

N EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
Ir true p r r va u fr av al va u (i u (i u r va u u o av av av av u u o av av u u o av av av av av av av av av av	true	true	true	false	true	true	true	true	true
C true re la ir p va u w sz pl p te	true	false	false	false	true	true	true	true	false
U true oo ci re fr qi ci fc ir pi ta ti	true	true	true	false			true	true	true

Table 3.196 - continued from previous page

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N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
P. m ri un ce ta	false	false	false	false	false	true	false	false	false	false
A pl oc cu re pa te po ce ag	false	false	true	true				false	false	true
T ge le	External	External	External	External	External	External	External	External	External	Exte
U va al it	BetaDis- tribution		BetaDis- tribution			BetaDis- tribution		BetaDis- tribution		Beta tribu
E ti m na tu	Realistic		Realistic			Realistic		Realistic		Real
U va al it pa ra e- te	Variabili- tyFactor		Variabili- tyFactor			Variabili- tyFactor		Variabili- tyFactor		Vari tyFa
M ty		OIM		OIM	OIM		OIM		OIM	
N (r b ot da in su ve		2		2					2	

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N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
S uj us fr qi tc 1			true	true						true
R st us p c c a g uj sc tc au tc au th ri			true	true						true
M F tc (f x N						1	1			
C c tr ti lii fii t e e c e fa tc										
Ir cl fc ca ca m it ca ca tr ti										

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
F ca co m it su su su cu re p ca aş										
A ju m fa tc fc tc fc cc tc tc tr										
U du te m is ti st cu vu st cu fc fc cu cu m it										

					Table	e 3.196 - co	ntinued from	previous pa	ige
N EC 201 Acute Tier I	8 EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
Ir cl pi cc di ti of tf fc cc fc ir r st m r ę pi n									

Retrospective dietary CRA (EC 2018) - Acute / Tier I

Table 3.197: Tier definition for Retrospective dietary CRA (EC 2018) - Acute / Tier I.

Name	Setting	From input tier	In module
Consumptions on the same day come from the same sample	false		
Use unit variability	true		
Model-then-add	false		
Covariate modelling	false		
Iterate survey	false		
Report consumptions and exposures per individual instead of per kg body weight	false		
Ignore sampling weights	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Consump- tions

Name	Setting	From input tier	ln module
Exclude individuals with less than N days	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Account for substance authorisations in substance conversions	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations

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Name	Setting	From input tier	In module
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Use extrapolation rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Threshold for extrapolation	10	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Restrict extrapolations to equal MRLs	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Restrict extrapolations to authorised uses	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Impute water concentrations	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Water concentration value (µg/kg)	0.1	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations

Table 3.197 - continued from previous page

Name	Setting	From input tier	ln module
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Restrict water imputation to approved substances	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Processing factors
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Processing factors
Default concentration model	Empirical	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models

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Name	Setting	From input tier	ln module
Include MRL fallback model	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Sample based	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Correlate imputed values with sample potency	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models

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Name	Setting	From input tier	ln module
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Parametric uncertainty	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Apply occurrence pattern percentages	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Occur- rence patterns
Target level	External	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Hazard character- isations
Unit variability model	BetaDis- tribution	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Unit variability factors
Estimates nature	Realistic	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Unit variability factors
Unit variability parameter	Variabili- tyFactor	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Unit variability factors

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Retrospective dietary CRA (EC 2018) - Chronic / Tier I

Name	Setting	From input tier	In module
Consumptions on the same day come from the same sample	false		
Model type	OIM		
Model-then-add	false		
Covariate modelling	false		
Iterate survey	false		
Report consumptions and exposures per individual instead of per kg body weight	false		
Ignore sampling weights	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Consump- tions
Exclude individuals with less than N days	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Consump- tions
N (number of days in survey)	2	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations

Table 3.198: Tier definition for Retrospective dietary CRA (EC 2018) - Chronic / Tier I.

Name	Setting	From input tier	ln module
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Account for substance authorisations in substance conversions	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Use extrapolation rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Threshold for extrapolation	10	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations

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Name	Setting	From input tier	ln module
Restrict extrapolations to equal MRLs	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Restrict extrapolations to authorised uses	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Impute water concentrations	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Water concentration value (µg/kg)	0.1	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Restrict water imputation to approved substances	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations

Table	3.198 -	continued	from	previous	page
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Name	Setting	From input tier	In module
Apply processing factors	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Processing factors
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Processing factors
Default concentration model	Empirical	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models

	Table	3.198 -	continued	from	previous	page
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Name	Setting	From input tier	ln module
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Sample based	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Correlate imputed values with sample potency	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Parametric uncertainty	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Apply occurrence pattern percentages	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Occur- rence patterns

Table 3.198 - continued from previous page

Name	Setting	From input tier	ln module
Target level	External	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Hazard character- isations

Table 3.198 - continued from previous page

Retrospective dietary CRA (EC 2018) - Acute / Tier II

Table 3.199: Tier definition for Retrospective dietary CRA (EC 2018) - Acute / Tier II.

Name	Setting	From input tier	In module
Consumptions on the same day come from the same sample	false	ſ	
Use unit variability	true		
Model-then-add	false		
Covariate modelling	false		
Iterate survey	false		
Report consumptions and exposures per individual instead of per kg body weight	false		
Ignore sampling weights	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Consump- tions
Exclude individuals with less than N days	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations

Name	Setting	From input tier	In module
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Use extrapolation rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Threshold for extrapolation	10	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations

Table 3.199 - continued from previous page

Name	Setting	From	In
		input tier	module
Restrict extrapolations to equal MRLs	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Restrict extrapolations to authorised uses	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Impute water concentrations	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Restrict water imputation to approved substances	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Apply processing factors	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Processing factors
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Processing factors
Default concentration model	Empirical	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models

Table 3.199 - continued from previous page
Name	Setting	From input tier	In module
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Sample based	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Correlate imputed values with sample potency	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Parametric uncertainty	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Apply occurrence pattern percentages	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Occur- rence patterns

Table	3.199 -	continued from	previous	page

Name	Setting	From input tier	In module
Scale up use frequency to 100%	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Occur- rence patterns
Target level	External	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Hazard character- isations
Unit variability model	BetaDis- tribution	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Unit variability factors
Estimates nature	Realistic	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Unit variability factors
Unit variability parameter	Variabili- tyFactor	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Unit variability factors

Table 3.199 - continued from previous page

Retrospective dietary CRA (EC 2018) - Chronic / Tier II

Table 3.200: Tier definition for Retrospective dietary CRA (EC 2018) - Chronic / Tier II.

Name	Setting	From input tier	ln module
Consumptions on the same day come from the same sample	false		

Name	Setting	From input tier	In module
Model type	OIM		
Model-then-add	false		
Covariate modelling	false		
Iterate survey	false		
Report consumptions and exposures per individual instead of per kg body weight	false		
Ignore sampling weights	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Consump- tions
Exclude individuals with less than N days	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Consump- tions
N (number of days in survey)	2	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Use extrapolation rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Threshold for extrapolation	10	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Restrict extrapolations to equal MRLs	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations

Table	3.200 -	continued	from	previous	page

Name	Setting	From input tier	In module
Restrict extrapolations to authorised uses	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Impute water concentrations	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Restrict water imputation to approved substances	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Processing factors

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Name	Setting	From input tier	In module
Use distribution	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Processing factors
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Processing factors
Default concentration model	Empirical	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models

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Name	Setting	From input tier	ln module
Sample based	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Correlate imputed values with sample potency	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Parametric uncertainty	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Apply occurrence pattern percentages	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Occur- rence patterns

Table	3.200 -	continued	from	previous	page

Name	Setting	From input tier	ln module
Restrict use percentage up-scaling to authorised uses	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Occur- rence patterns
Target level	External	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Hazard character- isations

Table 3.200 – continued from previous page

Retrospective dietary CRA (EFSA 2012) - Optimistic

Use the optimistic model settings according to the EFSA Guidance 2012. Concentration values are sampled using a sample-based empirical distribution. Available processing factors are applied. No unit variability model should be applied.

317

Ontimistic

N	ame	Setting	From input tier	In module
C	onsumptions on the same day come om the same sample	false		
	se unit variability	false		
N	Iodel type	OIM		
N	Iodel-then-add	false		
	ovariate modelling	false		
It		false		
P	eport consumptions and exposures	false		
po w	er individual instead of per kg body eight	laise		
Т	otal diet study concentration data	false	Retrospec- tive dietary CRA (EFSA 2012) - Optimistic	Food con- versions
A	pply processing factors	true	Retrospec- tive dietary CRA (EFSA 2012) - Optimistic	Processing factors
U	se distribution	false	Retrospec- tive dietary CRA (EFSA 2012) - Optimistic	Processing factors
Ig	more processing factors less than 1	false	Retrospec- tive dietary CRA (EFSA 2012) - Optimistic	Processing factors
D	efault concentration model	Empirical	Retrospec- tive dietary CRA (EFSA 2012) - Optimistic	Concen- tration models
Ir	nclude MRL fallback model	false	Retrospec- tive dietary CRA (EFSA 2012) - Optimistic	Concen- tration models
С	ensored values replacement	Replace- ByZero	Retrospec- tive	Concen- tration
Exposure n	nodules		dietary CRA (EFSA	models

Table 3.201: Tier definition for Retrospective dietary CRA (EFSA 2012) - Optimistic.

Retrospective dietary CRA (EFSA 2012) - Acute / Pessimistic

Acute probabilistic exposure assessment using the pessimistic model settings according to the EFSA Guidance 2012. Only processing factors > 1 are applied. For unit variability, the Beta distribution is applied.

319

Name	Setting	From input tier	In module
Consumptions on the same day come from the same sample	true		
Use unit variability	true		
Covariate modelling	false		
Iterate survey	false		
Report consumptions and exposures per individual instead of per kg body weight	false		
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Processing factors
Ignore processing factors less than 1	true	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Processing factors
Default concentration model	NonDe- tect- SpikeLog- Normal	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Concen- tration models
Include MRL fallback model	true	Retrospec- tive dietary CRA (EFSA 2012) -	Concen- tration models
posure modules		Passimistic	
Destrict I OD immediate	false	Detre -	Contract
authorised uses	Taise	Ketrospec- tive	tration

Table 3.202: Tier definition for Retrospective dietary CRA (EFSA 2012) - Acute / Pessimistic.

Retrospective dietary CRA (EFSA 2012) - Chronic / Pessimistic

Chronic probabilistic exposure assessment using the pessimistic model settings according to the EFSA Guidance 2012. Only processing factors > 1 are applied.

321

	Name	Setting	From input tier	In module
Ī	Consumptions on the same day come from the same sample	true		
	Model type	OIM		
	Model-then-add	false		
	Covariate modelling	false		
	Iterate survey	false		
	Report consumptions and exposures	false		
	per individual instead of per kg body weight	laise		
	Total diet study concentration data	false	Retrospec- tive dietary CRA (EFSA 2012) - Chronic / Pessimistic	Food con- versions
	Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2012) - Chronic / Pessimistic	Concen- trations
	Apply processing factors	true	Retrospec- tive dietary CRA (EFSA 2012) - Chronic / Pessimistic	<i>Frocessing</i> <i>factors</i>
	Use distribution	false	Retrospec- tive dietary CRA (EFSA 2012) - Chronic / Pessimistic	<i>Processing</i> <i>factors</i>
	Ignore processing factors less than 1	true	Retrospec- tive dietary CRA (EFSA 2012) - Chronic / Pessimistic	Processing factors
	Default concentration model	NonDe- tect- SpikeLog- Normal	Retrospec- tive dietary CRA (EFSA 2012)	Concen- tration models
Exposure	e modules		Chronic / Pessimistic	
	Include MRL fallback model	true	Retrospec-	Concen-

Table 3.203: Tier definition for Retrospective dietary CRA (EFSA 2012) - Chronic / Pessimistic.

Retrospective dietary CRA (EFSA 2022) - Acute / Tier I

Name	Setting	From input tier	In module
Consumptions on the same day come from the same sample	false		
Use unit variability	true		
Model-then-add	false		
Covariate modelling	false		
Iterate survey	false		
Report consumptions and exposures per individual instead of per kg body weight	false		
Ignore sampling weights	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Consump- tions
Exclude individuals with less than N days	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations

Table 3.204: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier I.

Name	Setting	From input tier	ln module
Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Account for substance authorisations in substance conversions	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Use extrapolation rules	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Impute water concentrations	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations

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Name	Setting	From input tier	ln module
Water concentration value (µg/kg)	0.1	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Restrict water imputation to approved substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Processing factors

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Name	Setting	From input tier	ln module
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Processing factors
Default concentration model	Empirical	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models

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Name	Setting	From input tier	ln module
Sample based	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Correlate imputed values with sample potency	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Parametric uncertainty	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Apply occurrence pattern percentages	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Occur- rence patterns

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Name	Setting	From input tier	ln module
Target level	External	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Hazard character- isations
Unit variability model	BetaDis- tribution	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Unit variability factors
Estimates nature	Realistic	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Unit variability factors
Unit variability parameter	Variabili- tyFactor	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Unit variability factors

Table 3.204 – continued from previous page
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Retrospective dietary CRA (EFSA 2022) - Chronic / Tier I

Table 3.205: Tier definition for Retrospective dietary CRA (EFSA 2022)
- Chronic / Tier I.	

Name	Setting	From input tier	In module
Consumptions on the same day come	false		
from the same sample			
Model type	OIM		
Model-then-add	false		
Covariate modelling	false		
Iterate survey	false		
Report consumptions and exposures	false		
per individual instead of per kg body			
weight			
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Name	Setting	From input tier	ln module
Ignore sampling weights	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Consump- tions
Exclude individuals with less than N days	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Consump- tions
N (number of days in survey)	2	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations

Table 3.205 - continued from previous page

Name	Setting	From input tier	ln module
Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Account for substance authorisations in substance conversions	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Use extrapolation rules	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Impute water concentrations	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations

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Name	Setting	From input tier	ln module
Water concentration value (µg/kg)	0.1	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Restrict water imputation to approved substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Processing factors

Table 3.205 - continued from previous page

Name	Setting	From input tier	ln module
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Processing factors
Default concentration model	Empirical	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models

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Name	Setting	From input tier	In module
Sample based	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Correlate imputed values with sample potency	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Parametric uncertainty	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Apply occurrence pattern percentages	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Occur- rence patterns

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Name	Setting	From input tier	In module
Target level	External	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Hazard character- isations

Retrospective dietary CRA (EFSA 2022) - Acute / Tier II

Name	Setting	From input tier	In module
Consumptions on the same day come from the same sample	false		
Use unit variability	true		
Model-then-add	false		
Covariate modelling	false		
Iterate survey	false		
Report consumptions and exposures per individual instead of per kg body weight	false		
Ignore sampling weights	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Consump- tions
Exclude individuals with less than N days	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Food con- versions

Table 3.206: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier II.

Name	Setting	From input tier	ln module
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations

Table 3.206 - continued from previous page

Name	Setting	From input tier	In module
Use extrapolation rules	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Impute water concentrations	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Restrict water imputation to approved substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Apply processing factors	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Processing factors
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Processing factors
Default concentration model	Empirical	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models

Table 3.206 - continued from previous page

Name	Setting	From input tier	ln module
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Sample based	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Correlate imputed values with sample potency	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models

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Name	Setting	From input tier	In module
Parametric uncertainty	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Apply occurrence pattern percentages	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Occur- rence patterns
Target level	External	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Hazard character- isations
Unit variability model	BetaDis- tribution	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Unit variability factors

Table 3.206 - continued from previous page

Name	Setting	From input tier	ln module
Estimates nature	Realistic	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Unit variability factors
Unit variability parameter	Variabili- tyFactor	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Unit variability factors

Table 3.206 - continued from previous page

Retrospective dietary CRA (EFSA 2022) - Chronic / Tier II

Table 3.207: Tier definition for Retrospective dietary CRA (EFSA 2022) - Chronic / Tier II.

Consumptions on the same day come from the same samplefModel typeCModel-then-addffCovariate modellingffIterate surveyffReport consumptions and exposuresff	false OIM false false false		
Model typeCModel-then-addfrCovariate modellingfrIterate surveyfrReport consumptions and exposuresfr	OIM false false false		
Model-then-addfCovariate modellingfIterate surveyfReport consumptions and exposuresf	false false false		
Covariate modellingf.Iterate surveyf.Report consumptions and exposuresf.	false false		
Iterate surveyf.Report consumptions and exposuresf.	false		
Report consumptions and exposures f			
per individual instead of per kg body weight	false		
Ignore sampling weights t	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Consump- tions
Exclude individuals with less than N takes that the second	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Consump- tions

Name	Setting	From input tier	In module
N (number of days in survey)	2	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations

Table 3.207 - continued from previous page

Name	Setting	From input tier	In module
Account for substance authorisations in substance conversions	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Use extrapolation rules	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Impute water concentrations	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations

Table 3.207 - continued from previous page

Name	Setting	From input tier	ln module
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Restrict water imputation to approved substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Processing factors
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Processing factors
Default concentration model	Empirical	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models

Table 3.207 - continued from previous page

Name	Setting	From input tier	ln module
Include MRL fallback model	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Sample based	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models

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Name	Setting	From input tier	ln module
Correlate imputed values with sample potency	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Parametric uncertainty	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Apply occurrence pattern percentages	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Occur- rence patterns

Table 3.207 - continued from previous page
Name	Setting	From input tier	In module
Target level	External	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Hazard character- isations

Table	3.207 -	- continued fr	om previous	page

Prospective dietary CRA (EFSA 2023) - Acute / Tier II

Table 3.208:	Tier definition for	Prospective	dietary	CRA (EFSA	2023) -
Acute / Tier I	I.					

Name	Setting	From input tier	In module
Consumptions on the same day come from the same sample	false		
Use unit variability	true		
Model-then-add	false		
Covariate modelling	false		
Iterate survey	false		
Report consumptions and exposures per individual instead of per kg body weight	false		
Ignore sampling weights	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Consump- tions
Exclude individuals with less than N days	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Consump- tions
Total diet study concentration data	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Food con- versions

Name	Setting	From input tier	In module
Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations

Table 3.208 - continued from previous page

Name	Setting	From input tier	In module
Fix duplicate substance allocation inconsistencies	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Use extrapolation rules	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Impute water concentrations	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations

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Name	Setting	From input tier	ln module
Restrict water imputation to approved substances	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Include focal commodity concentrations	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Focal commodity substance occurrence percentage	20	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Adjustment factor for the focal food/substance concentration	1	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Use deterministic substance conversions for focal commodity	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Apply processing factors	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Processing factors

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Name	Setting	From input tier	ln module
Use distribution	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Processing factors
Ignore processing factors less than 1	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Processing factors
Default concentration model	Empirical	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Include MRL fallback model	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models

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Name	Setting	From input tier	In module
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Sample based	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Correlate imputed values with sample potency	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Parametric uncertainty	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models

Table 3.208 - continued from previous page

Name	Setting	From input tier	ln module
Apply occurrence pattern percentages	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Occur- rence patterns
Target level	External	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Hazard character- isations
Unit variability model	BetaDis- tribution	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Unit variability factors
Estimates nature	Realistic	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Unit variability factors

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Name	Setting	From input tier	ln module
Unit variability parameter	Variabili- tyFactor	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Unit variability factors

Table 3.208 - continued from previous page

Prospective dietary CRA (EFSA 2023) - Chronic / Tier II

Table 3.209: Tier definition for Prospective dietary CRA (EFSA 2023) - Chronic / Tier II.

Name	Setting	From input tier	In module
Consumptions on the same day come from the same sample	false		
Model type	OIM		
Model-then-add	false		
Covariate modelling	false		
Iterate survey	false		
Report consumptions and exposures per individual instead of per kg body weight	false		
Ignore sampling weights	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Consump- tions
Exclude individuals with less than N days	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Consump- tions
N (number of days in survey)	2	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Consump- tions

Name	Setting	From input tier	ln module
Total diet study concentration data	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Food con- versions
Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations

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Name	Setting	From input tier	ln module
Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Use extrapolation rules	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Impute water concentrations	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations

Table 3.209 - continued from previous page

Name	Setting	From input tier	ln module
Restrict water imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Restrict water imputation to approved substances	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Include focal commodity concentrations	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Focal commodity substance occurrence percentage	20	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Adjustment factor for the focal food/substance concentration	1	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Use deterministic substance conversions for focal commodity	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Apply processing factors	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Processing factors
Use distribution	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Processing factors
Ignore processing factors less than 1	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Processing factors
Default concentration model	Empirical	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Include MRL fallback model	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models

Table 3.209 - continued from previous page

Name	Setting	From input tier	ln module
Censored values replacement	Replace- ByLOR	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Sample based	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Correlate imputed values with sample potency	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models

Table	3.209 -	continued	from	previous	page

Name	Setting	From input tier	ln module
Parametric uncertainty	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Apply occurrence pattern percentages	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Occur- rence patterns
Target level	External	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Hazard character- isations

Table 3.209 - continued from previous page

Prospective dietary CRA (EFSA 2024) - Acute / Tier II

Table 3.210: Tier definition for Prospective dietary CRA (EFSA 2024) - Acute / Tier II.

Name	Setting	From input tier	In module
Consumptions on the same day come from the same sample	false		
Use unit variability	true		
Model-then-add	false		

Name	Setting	From input tier	In module
Covariate modelling	false		
Iterate survey	false		
Report consumptions and exposures per individual instead of per kg body weight	false		
Ignore sampling weights	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Consump- tions
Exclude individuals with less than N days	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Consump- tions
Total diet study concentration data	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Food con- versions
Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations

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Name	Setting	From input tier	ln module
Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Use extrapolation rules	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Impute water concentrations	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations

Table 3.210 - continued from previous page

Name	Setting	From input tier	ln module
Water concentration value (µg/kg)	0.05	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Restrict water imputation to approved substances	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Include focal commodity concentrations	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Focal commodity substance occurrence percentage	20	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Adjustment factor for the focal food/substance concentration	1	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Use deterministic substance conversions for focal commodity	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Include processed derivatives of the focal food in measurement replacement	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Apply processing factors	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Processing factors
Use distribution	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Processing factors
Ignore processing factors less than 1	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Processing factors

Table 3.210 - continued from previous page

Name	Setting	From input tier	ln module
Default concentration model	Empirical	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Include MRL fallback model	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Sample based	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models

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Name	Setting	From input tier	In module
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Correlate imputed values with sample potency	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Parametric uncertainty	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Apply occurrence pattern percentages	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Occur- rence patterns

Table 3.210 - continued from previous page

Name	Setting	From input tier	In module
Restrict use percentage up-scaling to authorised uses	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Occur- rence patterns
Target level	External	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Hazard character- isations
Unit variability model	BetaDis- tribution	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Unit variability factors
Estimates nature	Realistic	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Unit variability factors
Unit variability parameter	Variabili- tyFactor	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Unit variability factors

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Prospective dietary CRA (EFSA 2024) - Chronic / Tier II

Table 3.211: Tier definition for Prospective dietary CRA (EFSA 2024) - Chronic / Tier II.

Name	Setting	From input tier	ln module
Consumptions on the same day come from the same sample	false		
Model type	OIM		
Model-then-add	false		

Name	Setting	From input tier	ln module
Covariate modelling	false		
Iterate survey	false		
Report consumptions and exposures per individual instead of per kg body weight	false		
Ignore sampling weights	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Consump- tions
Exclude individuals with less than N days	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Consump- tions
N (number of days in survey)	2	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Consump- tions
Total diet study concentration data	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Food con- versions
Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations

Table 3.211 - continued from previous page

Name	Setting	From input tier	ln module
Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Use extrapolation rules	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Impute water concentrations	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Restrict water imputation to approved substances	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Include focal commodity concentrations	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations

Table 3.211 - continued from previous page

Name	Setting	From input tier	In module
Focal commodity substance occurrence percentage	20	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Adjustment factor for the focal food/substance concentration	1	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Use deterministic substance conversions for focal commodity	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Include processed derivatives of the focal food in measurement replacement	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Apply processing factors	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Processing factors
Use distribution	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Processing factors

Table 3.211 – continued from previous pa	age
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Name	Setting	From input tier	In module
Ignore processing factors less than 1	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Processing factors
Default concentration model	Empirical	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Include MRL fallback model	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models

Table 3.211 - continued from previous page

Name	Setting	From input tier	ln module
Sample based	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Correlate imputed values with sample potency	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Parametric uncertainty	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Apply occurrence pattern percentages	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Occur- rence patterns

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Name	Setting	From input tier	In module
Scale up use frequency to 100%	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Occur- rence patterns
Target level	External	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Hazard character- isations

Table 3.211 – continued from previous page

Calculation of dietary exposures

Dietary exposures are calculated from consumptions per modelled food and concentration models. Optionally, also processing factors and unit variability models are applied.

• Dietary exposures calculation

Inputs used: Consumptions by modelled food Concentration models Processing factors Unit variability factors High exposure food-substance combinations Active substances Occurrence patterns Relative potency factors Food conversions Concentration distributions

Settings used

• Calculation Settings

3.4.3 Dust exposures

Dust exposures are the amounts of substances, expressed per kg bodyweight, to which individuals in a population are exposed from dust per day. Dust exposures are long-term/chronic exposures, they represent the average exposure per day over an unspecified longer time period.

This module has as primary entities: Populations Substances

Output of this module is used by: Internal exposures

Dust exposures calculation

Long-term (chronic) exposure to substances via dust can be estimated as a single value calculated from single value inputs. MCRA implements the calculation method that is also available in the RSExpo tool.



Figure 3.38: Modular design for dust exposures module when simulated individuals are used to generate exposures for source dust

Dust exposures calculation details

The long term (chronic) dust exposure assessment can be split in dermal and oral exposure related to exposure to substances in dust. MCRA applies in principle the equations as used in RSExpo.

Dust exposures are generated using properties age, sex and *BSA* of a *simulated set of individuals* or, as an alternative, an existing set of individuals (currently, the dietary individual population, see *module dietary exposures*). Specify the reference population with option *Specify individuals for generating dust exposures* with settings *Simulate individuals* or *Use dietary individuals*.

Dermal exposure

The calculation for dermal exposure is:

$$\mathbf{E}_{\text{dust-derm}} = \frac{1}{\text{BW}} \cdot \mathbf{f}_{\text{d}} \cdot \mathbf{M}_{\text{adh,d}} \cdot \frac{\mathbf{t}_{\text{dust}}}{24} \cdot \mathbf{S} \cdot \mathbf{f}_{\text{s,d}} \cdot \mathbf{c}_{\text{d}}$$

Parameters used in the equation

 $E_{air-inhal}$: average daily external exposure of an individual to substances by dust dermal contact $(\mu g/kg bw/day)$

- BW: body weight of the individual (default in kg)
- f_d : fraction of substance in dust available for dermal contact (%)
- $M_{adh,d}$: amount of dust adhering to the skin (default in $g/m^2/day$)
- $\mathtt{t}_{\mathtt{dust}}$: time of home dust exposure
- S: body surface (default in m^2)
- $f_{s,d}$: fraction of the body surface exposed to dust (%)
- c_d : substance concentration in dust (default in $\mu g/g$)

Ingestion exposure

The calculation for exposure through oral ingestion is:

$$\mathbf{E}_{\text{dust-ing}} = \frac{1}{\mathbf{BW}} \cdot \mathbf{q}_{\mathrm{d}} \cdot \mathbf{c}_{\mathrm{d}}$$

Parameters used in the equation

 ${\rm E}_{\rm dust-ing}$: average daily external exposure of an individual to substances by dust ingestion $(\mu g/kg\,bw/day)$

BW: body weight of the individual (default in kg)

 q_d : average daily amount of dust ingested by the individual (default in g/day)

 c_d : substance concentration in dust (default in $\mu g/g$)

Dust exposures settings

Calculation settings

Name	Туре	Description
Method for generating individual set for dust exposures	DustExposuresIndividualGener- ationMethod	Method for generating individual set for generation of dust exposures.
Seed for pseudo-random number generator	Numeric	A value of 0 will use a pseudo-random seed in each run, a valu 0 will provide the same results in a repeated run.
Selected exposure routes	SelectedExposureRoutes	Exposure routes included in dust exposure computation.
Time exposed to home dust	Numeric	Hours per day exposed to dust.
DUST: Population alignment method (RSExpo) dust exposures	PopulationAlignmentMethod	Population aligment methods for (RSExpo) dust exposure source

Table 3.212: Calculation settings for module Dust exposures.

Output settings

Table 3.213: C	output settings	for module	e Dust ex	posures.
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Name	Туре	Description
Lower percentage for variability (%)	Numeric	The default value of 25% may be overruled.
Upper percentage for variability (%)	Numeric	The default value of 75% may be overruled.
Report consumptions and exposures per individual instead of per kg body weight	Boolean	Specifies whether body weights should be ignored and consumptions and exposures should be expressed per individua Otherwise, the consumptions and exposures are per kg body weight.

Uncertainty settings

Name	Туре	Description
Lower uncertainty limit (%)	Numeric	Percentage lower bound, e.g. 2.5%.
Upper uncertainty limit (%)	Numeric	Percentage upper bound, e.g. 97.5%.

Table 3.214: Uncertainty settings for module Dust exposures.

Calculation of dust exposures

Dust exposures are calculated from dust concentration models.

• Dust exposures calculation

Inputs used: Dietary exposures Dust concentration distributions Dust exposure determinants Individuals

Settings used

• Calculation Settings

3.4.4 Soil exposures

Soil exposures are the amounts of substances, expressed per kg bodyweight, to which individuals in a population are exposed from soil per day. Soil exposures are long-term/chronic exposures, they represent the average exposure per day over an unspecified longer time period.

This module has as primary entities: Populations Substances

Output of this module is used by: Internal exposures

Soil exposures calculation

Long-term (chronic) exposure to substances via dust can be estimated as a single value calculated from single value inputs. MCRA implements the calculation method that is also available in the RSExpo tool.

Soil exposures calculation details

The long term (chronic) soil exposure assessment is assessed as oral exposure related to exposure to substances in soil. MCRA applies in principle the equations as used in RSExpo.

Soil exposures are generated using properties age and sex of a *simulated set of individuals* or, as an alternative, an existing set of individuals (currently, the dietary individual population, see *module dietary exposures*). Specify the reference population with option *Specify individuals for generating soil exposures* with settings *Simulate individuals* or *Use dietary individuals*.

Oral exposure

The calculation for oral exposure is:

$$\mathbf{E}_{\texttt{soil-ing}} = \frac{1}{\mathtt{BW}} \cdot \mathbf{q}_{\texttt{s}} \cdot \mathbf{c}_{\texttt{s}}$$

Parameters used in the equation

 ${\rm E_{soil-ing}}$: average daily external exposure of an individual to substances by soil ingestion $(\mu g/kg\,bw/day)$

BW: body weight of the individual (default in kg)

- q_s : average daily amount of soil ingested by the individual (default in g/day)
- c_s : substance concentration in soil (default in $\mu g/g$)



Figure 3.39: Modular design for soil exposures module when simulated individuals are used to generate exposures for source soil

Soil exposures settings

Calculation settings

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Name	Туре	Description
Method for generating individual set for soil exposures	SoilExposuresIndividualGener- ationMethod	Method for generating individual set for generation of soil exposures.
Seed for pseudo-random number generator	Numeric	A value of 0 will use a pseudo-random seed in each run, a valu 0 will provide the same results in a repeated run.
SOIL: Population alignment method (RSExpo) soil exposures	PopulationAlignmentMethod	Population aligment methods for (RSExpo) soil exposure source

Table 3.215: Calculation settings for module Soil exposures.

Output settings

Table 3.216:	Output	settings	for	module	Soil	exposures.

Name	Туре	Description
Lower percentage for variability (%)	Numeric	The default value of 25% may be overruled.
Upper percentage for variability (%)	Numeric	The default value of 75% may be overruled.
Report consumptions and exposures per individual instead of per kg body weight	Boolean	Specifies whether body weights should be ignored and consumptions and exposures should be expressed per individua Otherwise, the consumptions and exposures are per kg body weight.

Uncertainty settings

Table 3.217:	Uncertainty	settings	for modu	le Soil	exposures.
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Name	Туре	Description
Lower uncertainty limit (%)	Numeric	Percentage lower bound, e.g. 2.5%.
Upper uncertainty limit (%)	Numeric	Percentage upper bound, e.g. 97.5%.

Calculation of soil exposures

Soil exposures are calculated from soil concentration models.

• Soil exposures calculation

Inputs used: Dietary exposures Soil concentration distributions Soil exposure determinants Individuals

Settings used

• Calculation Settings

3.4.5 Air exposures

Air exposures are the amounts of substances, expressed per kg bodyweight, to which individuals in a population are exposed from air per m3. Air exposures are long-term/chronic exposures, they represent the average exposure per m3 over an unspecified longer time period.

This module has as primary entities: Populations Substances

Output of this module is used by: Internal exposures

Air exposures calculation

Long-term (chronic) exposure to substances via air can be estimated as a single value calculated from single value inputs. MCRA implements the calculation method that is also available in the RSExpo tool.



Figure 3.40: Modular design for air exposures module when simulated individuals are used to generate exposures for source air

Air exposures calculation details

The long term (chronic) air exposure assessment through inhalation is related to exposure to substances in air. MCRA applies in principle the equations as used in RSExpo.

Air exposures are generated using properties age and sex of a *simulated set of individuals* or, as an alternative, an existing set of individuals (currently, the dietary individual population, see *module dietary exposures*). Specify the reference population with option *Specify individuals for generating air exposures* with settings *Simulate individuals* or *Use dietary individuals*.



Figure 3.41: Modular design for air exposures module when dietary individuals are used to generate exposures for source air

Inhalation exposure

The calculation for inhalation exposure is:

$$\mathbf{E}_{\texttt{air-inhal}} = \frac{1}{\texttt{BW}} \cdot \mathbf{F}_{\texttt{a}} \cdot \left[\texttt{c}_{\texttt{in,a}} \cdot \texttt{t}_{\texttt{in}} + \texttt{c}_{\texttt{out,a}} \cdot (1 - \texttt{t}_{\texttt{in}}) \right]$$

Parameters used in the equation

 $E_{air-inhal}$: average daily external exposure of an individual to substances by air inhalation (default in $\mu g/kg bw/day$)

BW: body weight of the individual (default in kg)

 F_a : daily ventilatory flow (m^3/day)

 t_{in} : fraction of daily time spend indoors

 $c_{in,a}$: substance concentration in indoor air (default in $\mu g/m^3$)

 ${\rm c_{out.a}}$: substance concentration in outdoor air (default in $\mu g/m^3)$

Air exposures settings

Calculation settings

Name	Туре	Description
Method for generating	AirExposuresIndividualGenera-	Method for generating individual set for generation of air
individual set for air exposures	tionMethod	exposures.
Seed for pseudo-random	Numeric	A value of 0 will use a pseudo-random seed in each run, a value
number generator		0 will provide the same results in a repeated run.
Selected exposure routes	SelectedExposureRoutes	Exposure routes included in dust exposure computation.

Table 3.218: Calculation settings for module Air exposures.

Output settings

Name	Туре	Description
Lower percentage for variability (%)	Numeric	The default value of 25% may be overruled.
Upper percentage for variability (%)	Numeric	The default value of 75% may be overruled.
Report consumptions and exposures per individual instead of per kg body weight	Boolean	Specifies whether body weights should be ignored and consumptions and exposures should be expressed per individua Otherwise, the consumptions and exposures are per kg body weight.

Table 3.219: Output settings for module Air exposures.

Uncertainty settings

Table 3.220:	Uncertainty	settings	for module	Air exposures.
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Name	Туре	Description
Lower uncertainty limit (%)	Numeric	Percentage lower bound, e.g. 2.5%.
Upper uncertainty limit (%)	Numeric	Percentage upper bound, e.g. 97.5%.

Calculation of air exposures

Air exposures are calculated from air concentration models.

• Air exposures calculation

Inputs used: Dietary exposures Indoor air concentrations Outdoor air concentrations Air exposure determinants Individuals

Settings used

• Calculation Settings

3.4.6 Consumer product exposures

Consumer product exposures are the amounts of substances, expressed per kg bodyweight, to which individuals in a population are exposed from average consumer product usage per individual. Consumer product exposures are long-term/chronic exposures, they represent the average exposure per day over an unspecified longer time period.

This module has as primary entities: Populations Substances Consumer products

Output of this module is used by: Internal exposures

Consumer product exposures calculation

Long-term (chronic) exposure to substances via consumer products is based on realistic product usage information obtained from surveys. Currently, information on the usage (frequency and amount) of personal care products and household cleaning products in various European countries is included. The methods implemented are following the approach in *PACEM* and *RSExpo*.

See also, PACEM website.
Consumer product exposures calculation details

The long term (chronic) consumer product exposure assessment through oral ingestion, dermal contact and inhalation via personal care products, household cleaning products and do-it-yourself products is following the methodology as developed in *PACEM* and *RSExpo*. MCRA applies in principle the equations as used in PACEM and RSExpo.

Consumer product exposures are generated using an existing set of individuals, see *module consumer product use frequencies*. Together with information on the application amounts the exposure of personal care products, household cleaning products and do-it-yourself products in various European countries is estimated.

Consumer product exposure

The calculation for exposure to consumer products is:

 $\mathbf{E}_{\text{consumer-product}} = \texttt{frequency} \cdot \texttt{amount} \cdot \texttt{concentration} \cdot \frac{\sum_{route=1}^{p} EF_{route}}{BW} \cdot \texttt{occurrence}\%/100$

Parameters used in the equation

 $E_{consumer-product}$: average daily external exposure of an individual to substances in consumer products (default in $\mu g/kg \, bw/day$)

BW: body weight of the individual (default in kg)

amount: application amount (mg)

concentration: substance concentration in consumer product (mg/kg)

frequency: use frequency of een individual

occurrence%: occurence percentage, % of products with substance (default 100%)

EF_{route}: exposure fraction of a specific route (substance specific)

Consumer product exposures settings

Calculation settings

Name	Туре	Description
Selected exposure routes	SelectedExposureRoutes	Exposure routes included in dust exposure computation.
Seed for pseudo-random	Numeric	A value of 0 will use a pseudo-random seed in each run, a value
number generator		0 will provide the same results in a repeated run.

Output settings

Name	Туре	Description
Lower percentage for variability (%)	Numeric	The default value of 25% may be overruled.
Upper percentage for variability (%)	Numeric	The default value of 75% may be overruled.
Report consumptions and exposures per individual instead of per kg body weight	Boolean	Specifies whether body weights should be ignored and consumptions and exposures should be expressed per individua Otherwise, the consumptions and exposures are per kg body weight.

Uncertainty settings

Table 3.223:	Uncertainty	settings for	module Co	nsumer p	product exp	posures.
		<u> </u>				

Name	Туре	Description
Lower uncertainty limit (%)	Numeric	Percentage lower bound, e.g. 2.5%.
Upper uncertainty limit (%)	Numeric	Percentage upper bound, e.g. 97.5%.

Calculation of consumer product exposures

Consumer product exposures are calculated from consumer product concentration, use frequencies and amount models.

Consumer product exposures calculation

Inputs used: Consumer product concentrations Consumer product use frequencies Consumer product exposure determinants

Settings used

• Calculation Settings

3.4.7 Individuals

Simulates individuals with properties like age, body weight, sex, and body surface area.

This module has as primary entities: Populations

Output of this module is used by: Dust exposures Soil exposures Air exposures

Individuals calculation

This module is used to generate a set of simulated individuals with properties age, bodyweight, sex and body surface area (BSA). The simulated individuals are used for generating exposures in the modules for sources like dust, air and soil. After selecting *Simulate individuals* (see option 'Specify individuals for generating *dust, air* and *soil* exposures') in the *dust exposures, air exposures* and *soil exposures* modules the individuals module becomes visible. Also specify the number of individuals.

Ages are randomly drawn in a range from 0 to 100. For sex, available levels are *male* and *female*. Bodyweights are calculated as

 $\texttt{BW} = \texttt{BW}_{\texttt{birth}} + 4.47 \cdot \texttt{age} - 0.093 \cdot \texttt{age}^2 + 0.00061 \cdot \texttt{age}^3$

(see https://efsa.onlinelibrary.wiley.com/doi/10.2903/j.efsa.2020.6223, appendix, sensitivity analysis)

BW: body weight of the individual (default in kg)

BW_{birth}: body weight at birth = 3.68 (in kg)

The *BSA* is based on an allometric scaling using a standard bodyweight of 70 kg and standard BSA of 2.70 m^2 . Then the applied scaling is:

scaling =
$$BW_{standard} / BW^{(1-0.7)}$$

See also dust exposures, soil exposures and air exposures module.

Individuals settings

Calculation settings

Name	Туре	Description	
Number of simulated individuals	Numeric	Number of individuals that is simulated for dust exposures.	
Seed for pseudo-random number generator	Numeric	A value of 0 will use a pseudo-random seed in each run, a valu 0 will provide the same results in a repeated run.	

Table 3.224: Calculation settings for module Individuals.

Calculation of individuals

Simulates individuals.

• Individuals calculation

Settings used

• Calculation Settings

3.4.8 High exposure food substance combinations

Identification of food-as-eaten/modelled food/substance combinations that have the highest expected contribution to exposure based on a simple screening model.

This module has as primary entities: Foods Substances Effects

Output of this module is used by: Dietary exposures

High exposure food substance combinations calculation

A full Monte Carlo analysis can be unwieldy for large cumulative assessment groups (CAGs) and/or large number of foods or concentration data. An algorithmic approach was developed to handle large CAGs. Two unique features of MCRA are:

- contributions to the exposure results can be seen both in terms of foods as eaten (e.g. white bread) and modelled foods (e.g. wheat), and
- a drill-down can be made into the exact foods and substances contributing for simulated individuals or individual-days in the upper tail.

The number of combinations of simulation, substance, modelled food and food as eaten can be very large. To avoid memory problems with very large datasets, an additional optional modelling step, named *screening*, was added to MCRA. *Screening* should be used if the data dimensions are too large for a direct analysis. Screening identifies risk drivers. A full analysis based on screened risk drivers will still retain all food/substance combinations in the exposure calculation, and will therefore produce exactly the same cumulative exposure distribution, and allow to see contributions of all substances and all modelled foods. Details with respect to foods as eaten are however restricted to the risk drivers selected in the screening step. For more details see *screening calculation for large Cumulative Assessment Groups*.

The two-step approach consists of:

•Step 1: Data screening and selection of risk drivers

Run a simple analysis for each potential source/substance combination (SCC). Here source means the combination of food as eaten and modelled food, for example apple in apple pie. The screening is based on this combination, and not just modelled foods, to avoid problems with potentially multi-modal consumption distributions as much as possible (see van der Voet et al. 2014). SCCs are also referred to as risk driver components. The screening step in MCRA implements a simple model that is applied to

each SCC. The model calculates a percentile of interest in a distribution, consisting of a spike of zeroes (non-consumptions), and a mixture of two lognormal distributions for the exposure related to censored and positive concentrations, respectively. SCCs (risk driver components) can be combined to measured source/substance combinations (MSCCs, risk drivers). For example APPLE/apple juice/captan and AP-PLE/apple pie/captan combine to APPLE/captan. MCRA has an interface which identifies the Top-N SCCs (based on a chosen exposure percentile, e.g. p95) with an option to select N based on cumulative importance according to some criterion. Remark: Screening is performed before concentration modelling. Therefore there is no correction for processing at the screening stage. Note, originally SCC stands for Source Compound Combination, MSCC for Measured Source Compound Combination.

•Step 2: Full MC analysis

Perform the standard MC to all combinations of substances and foods, but restrict the stored information regarding foods as eaten to the SCCs selected in step 1.

The screening method requires to specify:

- Which percentile to consider for each single source/substance combination (SCC, potential risk driver component) (default p95)
- Which percentage of all exposures (according to the screening model) should be covered by the selected set of SCCs (default 95%)
- How to impute censored (non-detect or non-quantified) concentrations (c < LOR) in the screening step. The choice of a factor 0 (default) represents optimistic imputation, the choice of a factor 1 represents a pessimistic imputation. It may be noted that a factor 1 (pessimistic imputation) may select many SCCs (risk driver components) with relatively high LORs and high RPFs, but with only censored measurements. Choosing a lower fraction, e.g. 0.25 can be useful if a more realistic method is sought.

Based on limited experience with the EFSA test data, useful settings of these three screening parameters were found to be (95, 95, 0) in preparation for an EFSA optimistic run, and (50, 95, 0.25) in preparation for an EFSA pessimistic run. See also screening calculation *acute exposure* and *chronic exposure*.

Screening calculation for large Cumulative Assessment Groups

Statistical model for the screening step (acute exposure)

The screening step implements a simple model that is applied to each SCC. Assume independent *NonDetectSpike-LogNormal* (NDS-LN) models for both the consumptions of modelled food in source S and the concentrations of substance C in source S. A 'non-detect' consumption is assumed to be a zero consumption. A censored concentration will be imputed by a user-specified fraction f of the Limit of Reporting. Then the model for consumption has 3 parameters and the model for concentration has four parameters, as specified in Table 3.225. Note that the parameters of the consumption distribution are estimated from the consumption data using sampling weights if these have been provided in the consumption data set.

parameter	consumptions	concentrations
probability of a positive	π_x	π_c
mean positives (ln scale)	μ_x	μ_c
standard deviation positives (ln scale)	σ_x	σ_c
value to use for censored values (ln scale)		$f \cdot L_c$

Exposure is consumption times concentration, so on logarithmic scale they can be added:

e = x + c.

The assessment will focus on a chosen percentile of exposure, e.g. p95. The relevant fraction will be denoted by p, for example p = 0.95 for the 95th percentile. The two NDS-LN models combine to three possibilities, depending on whether there is consumption and if so, whether the concentration is censored or positive. In the screening model the two possibilities that lead to potential exposure are modelled with a mixture of two lognormal distribution. For the censored case the positive exposure distribution equals the positive consumption distribution modified by the

multiplication of a user-chosen factor times an estimate of the average worst-case limit value for concentration (LOR):

$$\pi_1 = \pi_x (1 - \pi_c); \mu_1 = \mu_x + f \cdot L_c; \sigma_1 = \sigma_x$$

where L_c is the logarithm of the LOR, or, if there are multiple analytical methods with different LOR, a weighted average of these different LORs.

For the detect case the positive exposure distribution is easily combined from the positive consumption distribution and the positive concentration distribution:

$$\pi_{2} = \pi_{x}\pi_{c}; \mu_{2} = \mu_{x} + \mu_{c}; \sigma_{12} = \sqrt{\sigma_{x}^{2} + \sigma_{c}^{2}}$$

p can be corrected for the non-consumptions to the appropriate fraction needed in the mixture of the two positive distributions:

$$p' = \frac{p - (1 - \pi_x)}{\pi_x}$$

If $p' \leq 0$ then all positive exposures are beyond the requested fraction, and the estimated exposure is just 0.

If p' > 0 then the relevant log exposure e_p satisfies

$$(1-\pi_c)\cdot\Phi\left(\frac{e_p-\mu_1}{\sigma_1}\right)+\pi_c\cdot\Phi\left(\frac{e_p-\mu_{12}}{\sigma_2}\right)=p'$$

where $\Phi(\cdot)$ represents the cumulative standard normal distribution function. The value of e_p can easily be found in a bisection search within the interval

$$[\mu_{min}-4\sigma_{max},\mu_{max}+max(0,z_{p'}\sigma_{max})].$$

The final exposure percentile estimate then is $\exp(e_n)$.

Denote by $e_{(p,max)}$ the highest estimate (for the SCC denoted by $SSC_{highest}$). Then evaluate for each SCC the probability to exceed $e_{(p,max)}$.

$$P_i = Pr(e > e_{p,max}) = \pi_x \cdot \left[(1 - \pi_c) * \Phi\left(\frac{e_{p,max} - \mu_1}{\sigma_1}\right) + \pi_c \cdot \Phi\left(\frac{e_{p,max} - \mu_2}{\sigma_1}\right) \right]$$

 P_i is a tentative measure for the 'probability of a high exposure'. For $SSC_{highest}$ $P_i = 1 - p$, for all other SCCs it will be lower. The sum of all these probabilities is not a meaningful probability in itself. However, this sum is used to scale the individual P_i values to measures of relative importance for the SCCs

$$Imp_i = P_i / \sum P_i$$

Rank all SCCs according to Imp_i and calculate cumulative importance. The relative importance of the two mixture components at e_p can be estimated as

$$w_{1,2} = \frac{\pi_{1,2} \cdot \phi\left(\frac{e_p - \mu_{1,2}}{\sigma_{1,2}}\right) / \sigma_{1,2}}{\pi_1 \cdot \phi\left(\frac{e_p - \mu_1}{\sigma_1}\right) / \sigma_1 + \pi_2 \cdot \phi\left(\frac{e_p - \mu_2}{\sigma_2}\right) / \sigma_2}$$

where $\phi(.)$ represent the standard normal probability density function. The user interface should allow to select the top-*N* SCCs from the list, based on a chosen percentage (e.g. 95%) of cumulative importance included. The full analysis will calculate exactly the same exposure distribution as a full analysis without screening. However, less information is retained in the output. This concerns tables with information on foods-as-eaten, which is only shown for the selected risk driver components (SCCs). Risk drivers are groupings of SCCs (risk driver components) at the level of measured-source-substance combinations (MSCCs). Note that output for an MSSC (e.g. APPLE/captan) only covers the selected SCCs (e.g. APPLE from apple juice/captan and APPLE from apple pie/captan), but not unselected SCCs (e.g. APPLE from fruit yoghurt/captan).

Statistical model for the screening step (chronic exposure)

In chronic exposure assessments, the mean concentration of chemicals is calculated first, and combined with the consumption distribution. For this reason a chronic calculation uses less memory, and therefore larger datasets can be handled.

The model described under *acute exposure* can be simplified for a chronic screening. The concentration distribution is only used to estimate a mean exposure, incorporating any effect from the imputation of censored values. The exposure distribution is therefore only a scaled version of the consumption distribution.

$$\pi_2=\pi_x\pi_c; \mu_2=\mu_x+\mu_c; \sigma_2=\sigma_x$$

The parameters of the consumption distribution (π_x, μ_x, σ_x) are calculated from the observed individual means (*OIM*), i.e. the mean daily consumptions over the survey days of each person in the data (allowing for sampling weights). The percentiles are calculated as $e_p = \mu_2 + z_p$ where z is a percentile of the standard normal distribution. The exceedances of the maximum percentile are calculated as

$$P_i = Pr(e > e_{p,max}) = \pi_x \cdot \Phi\left(\frac{e_{p,max} - \mu_2}{\sigma_2}\right)$$

High exposure food substance combinations settings

Calculation settings

Table 3.226: Calculation settings for module High exposure food-substance
combinations.

Name	Туре	Description
Exposure type	ExposureType	The type of exposure considered in the assessment; acute (short term) or chronic (long-term).
Compute cumulative exposures	Boolean	Specifies whether the assessment involves multiple substances a results should be cumulated over all substances.
Percentage defining the exposure percentile of interest per food-as-eaten/food-as- measured/substance combination	Numeric	Percentage defining the exposure percentile of interest per food-as-eaten/food-as-measured/substance combination.
Include risk drivers to include minimally a percentage	Numeric	The selection criterion for the risk drivers. The cumulative contribution percentage of the selected risk drivers will be this percentage.
Censored value replacement: factor x LOR	Numeric	A constant between 0 and 1. A value 0 can be used for an optimistic screening (LOR not used). Note that a factor 0.5 (pessimistic) may result in many and often high contributions from food-substance combinations with only censored values.
Report consumptions and exposures per individual instead of per kg body weight	Boolean	Specifies whether body weights should be ignored and consumptions and exposures should be expressed per individual Otherwise, the consumptions and exposures are per kg body weight.

Calculation of high exposure food-substance combinations

Screening results are computed for each combination of source (being a specific combination of food-aseaten/modelled food) and substance by combining simple approximations of the consumption and the concentration distribution.

• High exposure food-substance combinations calculation

Inputs used: Consumptions by modelled food Concentration models Active substances Relative potency factors

Settings used

• Calculation Settings

3.4.9 Internal exposures

Internal doses and concentrations are amounts of substances, typically expressed per mass unit and per day, to which individuals in a population are exposed at the chosen target level. This target level is an internal dose or an internal concentration (expressed per unit [organ] weight). Internal exposures may be aggregated from dietary and non-dietary exposures using either absorption factors for systemic doses or kinetic conversion factors or PBK models to translate the external exposures to internal concentrations at the specific target or tissue. Internal exposures can be short-term/acute exposures and then contain exposures for individual-days, or they can be long-term/chronic exposures, in which case they represent the average exposure per day over an unspecified longer time period.

This module has as primary entities: Populations Foods Consumer products Substances

Output of this module is used by: *Exposure mixtures Biological matrix concentration comparisons Risks Environmental burden of disease*

Internal exposures calculation

The main goal of the internal exposures module is to calculate the exposure as an internal concentration or internal dose. The internal concentration refers to the amount of chemical present in a specific target or tissue using kinetic conversion factors or PBK models. The internal dose is the amount of chemical that is systemically available using absorption factors and is the fraction of the external dose that is absorbed and enters the general circulation, see also WHO, Chapter 5 (2008). External exposures from dietary and non-dietary routes are aggregated at a specified internal level in two steps:

- 1. Linking dietary and non-dietary individual/individual-day exposures.
- 2. Computing the (aggregated) internal exposures at the specified internal level concentration or dose.

Absorption factors are just simple multiplication factors and aggregate the exposure from multiple routes into an internal dose. *Kinetic conversion factors* or more advanced *PBK models* aggregate the exposures from multiple routes into an internal concentration at a specified target (*biological matrix*). Currently, only dietary exposures or dietary exposures combined with non-dietary exposures are aggregated to an internal level. Internal exposure calculation of non-dietary sources only, is yet not available but will be implemented in the future.

In cumulative internal exposure calculations two simple approaches are used to identify and select mixtures contributing to the exposure of a target population:

- 1. qualitative approach: *counting of co-exposure*. To which combinations of substances are individuals exposed? Just the co-occurence of substances is observed.
- 2. quantitative approach: *maximum cumulative ratio (MCR)*. To what degree are mixtures more important than single substances? The relative exposure levels of the substances present in a measurement, e.g. an individual (chronic) or individual day (acute) are taken into account.

In the *exposures mixtures module*, two more advanced approaches are available to analyse the co-occurence of substances, the *SNMU approach* and a *network analysis*.

Computation of internal exposures (e.g. an internal doses or internal concentrations) requires *absorption factors*, *kinetic conversion factors* or *PBK models* to translate external doses, possibly from multiple routes, to internal doses (absorption) or concentrations at the target matrix/organ of interest (kinetic conversion).

Calculation of internal doses using absorption factors

In the simplest form, internal doses are derived from external doses using absorption factors specified for the combination substance and route. That is, for a given substance, the internal dose $dose_{int}$ is computed as

$$\textit{dose}_{int} = \sum_{r \in \textit{routes}} f_{abs,r} \cdot \textit{exp}_{ext,r}$$

Here, *routes* denotes the set of external exposure routes, $exp_{ext,r}$ denotes the external exposure for route r and $f_{abs,r}$ denotes the absorption factor of route r. Note that this model assumes that both external and internal doses refer to

amounts or doses depending on the *dietary exposures* setting (External exposure: substance amount per individual, or substance amount divided by body weight; internal exposure: substance amount per organ, or substance amount divided by organ weight). Also, both external and internal doses are expressed per day.

Calculation of internal concentrations using kinetic conversion factors

Kinetic conversion factors are used to convert an external (administered) dose to a concentration in a target or *biological matrix*. That is, for a given substance, the internal concentration $conc_{int}$ is computed as

$$conc_{int} = \sum_{r \in routes} f_{kc,r} \cdot exp_{ext,r}$$

Here, *routes* denotes the set external exposure routes, $exp_{ext,r}$ denotes the external exposure for route r and $f_{kc,r}$ denotes the kinetic conversion factor of route r.

When biological matrix urine is selected, concentrations are interpreted as such or interpreted as standardised by creatinine or normalised by specific gravity. Check option *Standardised (creatinine) or normalised (specific gravity) urine concentrations* and make a choice between expression type *creatinine* or *specific gravity*

When biological matrix blood is selected, concentrations are interpreted as such or interpreted as standardised by lipids. Check option *Standardised blood concentrations (lipids)*.

Calculation of internal concentrations using PBK models

A more detailed alternative to using kinetic conversion factors is to use one of the *advanced kinetic models* available in MCRA. In this approach, for each substance independently, the external exposures of an individual (chronic) or individual-day (acute) are presented for a number of simulated days to a PBK model of the individual. This yields a time course of the internal substance amount at the specified target or *biological matrix* from which a long term average substance amount (chronic) or peak substance amount (acute) can be obtained. An example of such a time course is given in Figure 3.42 for acute exposure assessments, and in Figure 3.43 for chronic exposure assessments. By dividing this substance amount by the weight of the biological matrix, an internal concentration is obtained. Notice that this procedure also changes the unit of the exposures from exposure per day to long term exposure.

Mathematically, the calculation of the peak substance amount (d_{peak}) for deriving acute internal concentration is as follows:

$$d_{\rm peak} = \max_{i=0,\dots,n_{\rm stop}} \left\{ d(t_{\rm start} + i\Delta t) \right\}. \label{eq:dpeak}$$

Here, d(t) denotes the substance amount at time t, t_{start} denotes the starting time of the evaluation window (defined by the *non-stationary period*), Δt denotes the time resolution of the kinetic model (e.g., hours or minutes), and n_{stop} denotes the total number of time-points, marking the end of the evaluation window (defined by the specified number of simulation days), which is computed as

$$n_{\rm stop} = \left\lfloor \frac{t_{\rm stop} - t_{\rm start}}{\Delta t} \right\rfloor.$$

Likewise, chronic long term average substance amounts (d_{ava}) are computed as:

$$d_{\rm avg} = \frac{\sum_{i=0}^{n_{\rm stop}} d(t_{\rm start} + i\Delta t)}{n_{\rm stop}}$$

Dosing patterns

In MCRA, the dietary and non-dietary exposures are computed at the level of exposures per day. However, when applying advanced PBK models, dosing patterns may be specified at a much finer resolution (e.g., hours or minutes). For this, a method is needed to translate external exposures provided per day to dosing patterns of substance amounts during the day. The simplest, yet not very realistic model is to apply, per route, the full exposure amount in one single dose at the beginning of the day. Alternatively, MCRA offers the possibility to specify, per route, the *number of exposure events per day*. If it is specified to use multiple doses per day, then the total substance amount of each day is divided into equal portions which are applied at regular time-intervals during the day.



Figure 3.42: Time course of the internal substance amount when applying the same single dose on each day. The acute internal concentration is derived as the peak substance amount (the green line in the figure) divided by the weight of the biological matrix. The vertical line at 50 indicates the selected end of an assumed non-stationary period, defining a burn-in period that is to be ignored for computing the peak substance amount.



Figure 3.43: Time course of the internal substance amount when randomly applying one of the individual-day doses for a number days. The chronic internal concentration is derived as the average substance amount (the blue line in the figure), divided by the weight of the biological matrix. The vertical line at 50 indicates the selected end of an assumed non-stationary period, defining a burn-in period that is to be ignored for computing the average substance amount.

Non-stationary period

Especially in the case of chronic exposure assessments, where a long term average exposure is computed based on the simulated time-course, it is important to realise that at time zero, the substance is commonly considered to be completely absent in the simulated system. However, this is not a realistic assumption. It is much more likely that the substance was already present in the system, and that the level is equal to the level obtained from applying the same chronic exposures to the system. For this, a specification of the *number of days skipped* (or burn-in period) is required in order to come to these initial concentration levels. This period is not used for computing the long term average or peak exposures, but just to determine initial (background) concentration levels.

Combining exposure sources

Individuals in a population are exposed to substances in sources like *food (diet)*, *plant protection products (PPPs)* (in MCRA also referred as non-dietary exposures), *dust*, *air* and *soil*.

In a dietary exposure assessment, individuals generally have known values for properties such as age, sex, and body weight. The same applies to non-dietary exposures generated by *BROWSE* when individual IDs are linked to a known set of individuals—such as those from a dietary exposure assessment—or through the specification of *survey properties* related to age and/or sex. For exposures generated by the dust, air and soil modules, no explicit individuals or population are available. However, the exposure determinants used in these modules require information about age and/or sex. For example, dust exposure determinants such as *ingestions* rates, *body exposure fractions*, *adherence amounts* or *availibility fractions* are age- and/or sex specific. Therefore, in order to generate exposures, a population of individuals is required.

There are two options available for generating this set of individuals. Generate exposures based on:

- individuals of the dietary survey e.g. see option Use dietary exposure,
- a set of Simulated individuals (see e.g. generation method individuals).

Navigate to the air, dust and soil modules to specify which set of individuals for generating exposures should be used. This choice is independently for each exposure source.

Before aggregating exposures from different sources at the internal level, all exposures must first be combined at the external level for each individual. To begin, specify one exposure source, its corresponding set of individuals (simulated or dietary) will serve as the reference set. All other exposure sources will then be aligned with this reference.

It is clear that combining exposures at the external level becomes complex when different sets of individuals are used by the exposure generator in the air, dust and/or soil modules.

After specifying the reference source, there are three options available for matching an individual in the reference set with an individual from the other sources:

- completely at random,
- based on individual properties like age, sex (or other properties)
- based on their individual ID,

For the first option, no additional information is required. Option 3, on the other hand, is the most restrictive, see also *population alignment methods*. This option is only applicable when each exposure source uses the same set of individuals. If the assessment includes a dietary source, all exposures from other source modules should be generated based on dietary individuals. If no dietary source is present, the appropriate choice is to use simulated individuals.

For the second option, individuals are matched based on specific properties. You can choose whether to use age and/or sex (male, female) or both as stratifying covariate. For age, matching can be done by specifying bins (e.g. 18 - 45 years) or by selecting individuals whose ages are closest to the age of the individual in the reference set. From the resulting subset, one individual is randomly sampled.

Non-dietary exposures

Originally, the *non-dietary exposure* module was developed to implement Bystanders, Residents, Operators and WorkerS Exposure models for plant protection products (*BROWSE*) software. Because this module has a different data format compared to the format of the *dust*, *air* and *soil* modules the method of matching individuals on the reference set is further explained.

In general, dietary and non-dietary surveys are independently taken (separated in space, location and time etc). Each survey contains a unique set of individuals and the only way to match individuals is *completely at random* or *use stratifying covariates* like age and or sex e.g. males, females between 18-45 years, children in the age of 0 to 5 years, etc.

Matching completely at random

For each individual in the dietary survey a random nondietary individual(day) from the nondietary surveys is sampled or vice versa. This option is also possible when surveys are identical.

Matching based on individual ID

When both dietary and non-dietary surveys contain identical individuals, individuals are matched based on their ID.

Matching based on individual properties

When dietary and non-dietary surveys are identical or partly identical, dietary and non-dietary exposures of individuals are matched based on the individual ID, e.g. for all *Individuals* of the food survey also a nondietary exposure is available or vice versa (method 3: *individuals are matched based on the individual Id*). Any nondietary exposure that does not correspond to an individual in the food survey will be ignored and vice versa (see *Example 2*). Occasionally, this is too restrictive and the assumption can be relaxed by specifying the individual ID in the nondietary survey as *Id* = *General*. Then, individual properties of the dietary individual should match the non-dietary survey properties.

When multiple non-dietary surveys are available, option *Match individuals of multiple non-dietary surveys* is relevant. When *matching* (correlation) is chosen, the exposure contributions of non-dietary individuals with identical *Id* in different surveys are combined and allocated to a dietary individual. When no correlation is chosen, the non-dietary exposures of randomly selected individuals from different surveys are combined and allocated to a dietary individual.

The user may also define demographic criteria for the assignment (for each source of non-dietary exposure) to indicate that those exposures are relevant only to a defined sub-population. Only those individuals in the food survey who meet the criteria of the non-dietary survey will be assigned non-dietary exposures from that source e.g. only males aged 18 to 65 (see *Example 1*). The simplest assessment consists of a single (deterministic) non-dietary exposure estimate which is assigned to all individuals in the food survey (Id = General). This case, and more complex possibilities are illustrated below using hypothetical examples (method 2: *individuals are match based on individual properties*).

Example 1

Deterministic cumulative (multi-substance) non-dietary exposure input, adult male sub-population. Unmatched case.

		7 1			
idIndividual	idNonDietarySurvey	idSubstance	Dermal	Oral	Inhalation
General	1	011003001	10	5	17
General	1	011003002	34	20	18
General	1	011003003	56	43	19

Table 3.227:	NonDietar	yExposures
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Figure 3.44: Exposure distributions for routes.

idNonDietary- Survey	Description	Location	Date	NonDietary- IntakeUnit
1	BROWSE, acute, cumulative, operators	York	09/10/2012	$\mu g/day$

Table 3.228: NonDietarySurveys

idNonDietary- Survey	Property- IndividualName	Individual- PropertyText- Value	Individual- Property- DoubleMin- Value	Individual- Property- DoubleMax- Value
1	Age		18	65
1	Gender	Male		

In this example, there are exposure values for multiple substances in Table 3.227 and the user has provided Table 3.229 which specifies that the non-dietary exposures given in survey number 1 relate to males aged 18 to 65.

When this assessment is performed, only those individuals whose property values fit the criteria in Table 3.229 will receive the non-dietary exposures in survey 1. The use of *idIndividual* = *General* indicates that this is the default exposure. All individuals in the dietary survey meeting the criteria receive all exposures given in the 3 rows, corresponding to 3 substances. The following should be noted:

- Specify only one *General* entry in the non-dietary exposures table per substance x survey combination.
- Property names and text properties must precisely match the values in the **IndividualProperties** and **IndividualPropertyValues** tables.
- The minimum and maximum values for numeric properties are inclusive boundaries.

In the next example, all males aged 18 to 65 receive the given exposures of all three substances and the other members of the population receive no non-dietary exposure. Note that example 1 describes the unmatched case.

Example 2

Variability (but no uncertainty) in cumulative non-dietary exposure input (matched to dietary survey individuals).

idIndividual	idNonDietarySurvey	idSubstance	Dermal	Oral
5432	1	011003001	10	5
5432	1	011003002	33	22
5433	1	011003001	12	7
5433	1	011003002	34	23
5434	1	011003001	18	9
5434	1	011003002	35	25
5435	1	011003001	10	5
5435	1	011003002	33	21

Table	3 230.	NonDietary	Exposures
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Table 3.231: NonDietarySurveys

idNonDietary- Survey	Description	Loca- tion	Date	NonDietaryIntakeU- nit
1	BROWSE, acute, cumulative, opera- tors	York	09/10/2012	$\mu g/day$

In this example, the non-dietary exposures are specified explicitly for individuals in the dietary population. Switch 'matching' on to allow exposure variability to be specified at the individual level. Only four individuals are used. The values in the *idIndividual* column of Table 3.230 match those in the *Individuals* table (dietary population).

It is not mandatory to specify exposures for every individual in the population. Those not included will simply receive a zero non-dietary exposure, unless there is also a default exposure value (*General* row(s) in Table 3.230) and the individual matches the specified demographic criteria for the survey, as specified in Table 3.229. (In this example, a default exposure would apply to all individuals not listed in Table 3.230 because Table 3.229 has not been used).

There is variability between individuals in this example, but no uncertainty in exposure. Note that these data could also be used with matching switched off. This would be the same as treating the *idIndividual* values as generic individuals, so that each pair of exposure lines would be assigned at random to individuals meeting the criteria.

Example 3

Variability (no uncertainty) in cumulative non-dietary exposure input (unmatched individuals).

idIndividual	idNonDietarySurvey	idSubstance	Dermal	Oral	Inhalation
ND1	1	011003001	10	5	17
ND1	1	011003002	33	22	45
ND2	1	011003001	12	7	18
ND2	1	011003002	34	23	47
ND3	1	011003001	18	9	19
ND3	1	011003002	35	25	49
ND4	1	011003001	10	5	17
ND4	1	011003002	33	21	45

Table 3.232: NonDietaryExposures

idNonDietary- Survey	Description	Loca- tion	Date	NonDietaryIntakeU- nit
1	BROWSE, acute, cumulative, opera- tors	York	09/10/2012	$\mu g/day$

Table 3.234: NonDietarySurveyProperties

idNonDi- etarySurvey	PropertyIndi- vidualName	IndividualProper- tyTextValue	IndividualProperty- DoubleMinValue	IndividualProperty- DoubleMaxValue
1	Age		50	65
1	Gender	Female		

This example is similar to example 2, except that the values in the *idIndividual* column of Table 3.232 do not match those in the *Individuals* table. In this instance, 'matching' would not be an option, and the non-dietary exposures would be randomly assigned to individuals who meet the criteria in Table 3.234. (In fact for the same result rather than changing the values in the *idIndividual* column in Table 3.230 from the previous example may be used with matching switched off). Exposures in Table 3.232 will be recycled if the number of exposure rows is less than the number of dietary records with which to aggregate exposures.

Again, there is variability between individuals in this example, but no uncertainty in exposure.

By allowing generic *idIndividual* values in this way, correlations between different sources (within individual) can be accounted for even in the unmatched case. If the same *idIndividual* value is used in different surveys, then the corresponding exposure values will be kept together and assigned to an eligible individual as a combined exposure.

So for option matching switched of, it is relevant whether individuals are correlated or not. In the following example, two non-dietary surveys are available, per survey three individuals are specified.

idIndividual	idNonDietarySurvey	idSubstance	Dermal	Oral	Inhalation
ND0	1	011003001	10	5	17
ND1	1	011003001	23	22	45
ND2	1	011003001	12	7	18
ND0	2	011003001	34	23	47
ND3	2	011003001	18	9	19
ND4	2	011003001	33	16	35

Table 3.235: matching switched of, with correlation or without.

- When a correlation is applied, the non-dietary exposure for individual ND0 from survey 1 and 2 are combined and allocated to a dietary individual. For individual ND1, ND2, ND3 and ND4 just a single non-dietary exposure is found and allocated to a dietary individual.
- When no correlation is applied, the exposure for individual ND0 from survey 1 is combined with one of the exposures of ND0, ND3 or ND4 from survey 2; exposure of ND1 from survey 1 is combined with one of the exposures of ND0, ND3 or ND4 from survey 2, etc.

When the intention is to sample just one exposure for a dietary individual, specify per survey different codes, e.g. ND1, ND2, ND3 for survey 1, ND4, ND5, ND6 for survey 2 and apply correlation, or specify 6 different individual codes and just one *idNonDietarySurvey*. Then, options with or without correlation are irrelevant and sampling results are identical.

See non-dietary exposure settings.



Contribution to the total exposure distribution by route

Figure 3.45: Contributions by route of the external exposure distributions.

Counting of co-exposure

In this qualitative approach, the number of combinations of substances to which an individual is exposed are recorded, see Table 3.236. There is no cut-off level, the only criterion is the presence of a substance in the simulated daily diet (eventually aggregated withe exposure from non-dietary sources) or not. For an *acute* or short term exposure assessment, a simulated individual day is the smallest entity to determine co-exposure. For a *chronic* or long term exposure assessment, co-exposures are summarized at the individual level, e.g. co-exposure is determined combining all consumption days of an individual.

Table 3.236: Counting combinations of substances in the exposure matrix: for example, on day 1 there is coexposure to substances Tebuconazole, Bitertanol and Triadimefon

Substance	day 1	day 2	day 3	 day n
Tebuconazole	х	Х		
Bitertanol	Х		Х	 Х
Triadimefon	Х			 Х

In Table 3.237, the frequency and percentage for the number of substances occurring together are shown.

	-r	
Number of substances	Frequency	Percentage
0	337	3.4
1	959	9.6
2	1207	12.1
3	1275	12.8

Table 3.237: Co-exposure of substances

In Table 3.238, the mixtures containing the substance(s) including all other combinations with the specified combination of substance(s), (a maximum number of 15 records is shown).

Number of substances	Percentage	Substances
1	5.88	Tebuconazole
2	3.88	Imazalil (aka enilconazole), Tebuconazole
0	3.37	
3	2.20	Difenoconazole, Imazalil (aka enilconazole), Tebuconazole
1	1.78	Imazalil (aka enilconazole)
3	1.76	Imazalil (aka enilconazole), Tebuconazole, Triadimenol

Table 3.238: Mixtures containing substances

Maximum Cumulative Ratio

Price and Han (2011) propose the Maximum Cumulative Ratio (MCR) which is defined as the ratio of the cumulative exposure received by an individual on an intake day to the largest exposure received from a single substance:

 $MCR = \frac{Cumulative exposure}{Maximum exposure}$

This MCR statistic is also picked up as a practical device in a recent JRC report, Bopp et al. (2015), to investigate cumulative exposure. If MCR is large, it is important to consider cumulative effects. If MCR is close to 1, the individual exposure will not be much different from a single-substance assessment. The MCR can therefore be interpreted as the degree to which the risk of being exposed is underestimated by not performing a cumulative risk assessment.

The MCR statistic is implemented in MCRA for both the *acute* risk and the *chronic* risk cases. In the acute risk case the short-term (single-day) exposures are used. For the chronic case long-term individual exposures (estimated by aggregating over the available survey days of each individual) are used.

Risk based, standardised or unweighted exposures

Before calculating the MCR statistics, three optional choices are available, see settings, MCR exposure approach type.

- Risk based exposures: exposures are multiplied by the *relative potency factor* (RPF) of each substance and thus exposures for different substances are on the same and comparable scale.
- Standardised exposures: all exposures are standardised to equal variance (unit variance).
- Unweighted exposures: exposures are taken as such, this is equivalent to RPF s equal to 1 for each substance.

Table 3.239 shows an artificial example how the MCR is calculated in the acute risk case. First the cumulative exposure per day is calculated by cumulating the exposure of each substance multiplied by the RPF. Then, for each day, the cumulative exposure (in equivalents of the reference substance) is divided by the maximum exposure of a single substance on that day. The last column shows the MCR values, with the substance with the highest exposure in parenthesis. The MCR has a value of 1 or close to 1 for mixtures where the exposure is dominated by one substance (e.g. day 1, substance B). When all substances have approximately equal exposure (e.g. day 3) the MCR value is equal or close to the number of substances, here 4. Day 2 represents an intermediate case. The MCR suggest that for exposure days (or persons) with MCR values close to 1, the need for a cumulative risk assessment is low.

	Substance A	Substance B	Substance C	Substance D	total exposure	ratio
day 1	0.01	0.99	0	0	1	1.01 (B)
day 2	0.1	0.2	0.3	0.4	1	2.50 (D)
day 3	0.25	0.25	0.24	0.26	1	3.99 (D)

Table 3.239: Maximum Cumulative Ratios

In this artificial example, all days have equal values for total exposure (= 1). For real data, total exposure will vary. It is obviously of interest to know if the MCR is high or low at those days (or individuals) where the total exposure is highest.

In Figure 3.46, French steatosis data (39 substances, 4079 persons) are used to calculate the chronic exposure matrix. For each individual the MCR is calculated and plotted against the total exposure. The different colours are used to identify the single substances with maximum exposure. From the original 39 substances, 10 different substances have the largest exposures. For the total exposure and MCR, the p_5 , p_{50} and p_{95} percentiles are indicated with the black line segments. The red line indicates the ratio with value 5. The dashed green lines indicate the p_{95} percentiles for the MCR value for different ranges of the total exposure.



Using MCR to identify substances that drive cumulative exposures

Figure 3.46: Maximum Cumulative Ratios vs total exposure

The plot shows that MCR values with Imazalil as risk driving substance (purple) are predominantly found in the lower part of the plot for relatively high values of the total exposure. A second finding is that MCR values decline when total exposure increases. This implies that cumulative exposure for most individuals is driven by multiple substances. At the right site of the plot, individuals are found with high exposure. Because MCR values tend to be lower here, higher exposures are received from one predominant substance and not because many substances are above the average level. For those individuals a cumulative risk assessment has less value.

Because Figure 3.46 can be very dense, in Figure 3.47, 95% confidence regions representing bivariate lognormal distributions of MCR and total exposure are plotted. The latter figure facilitates interpretation of the first figure. Note that substances with just one or two observations cannot be plotted in this display (substances with 2 observations are represented by a line).

In Figure 3.48 and Figure 3.49 scattered MCR distributions for the total and upper tail (here 37%) that drive the cumulative exposure are shown. The red line indicates the MCR threshold, 1.5. The black lines represent the regression lines MCR vs ln(Cumulative exposure) for each tail. Substances with an exposure contribution less than 15% are not displayed.

In Table 3.240 contributions to tail exposures at various percentile are shown. Column MCR = 1 shows the percent-



Bivariate distributions

Figure 3.47: Bivariate distributions MCR vs total exposure



Figure 3.48: Using MCR to identify substances that drive cumulative exposures, scatter distributions (total).



Figure 3.49: Using MCR to identify substances that drive cumulative exposures, scatter distributions (upper tail 37%).

age of tail exposure due to individual(day)s with a single substance. Column $1 < MCR \le 2$ shows the percentage of tail exposure due to individual(day)s with multiple substances, but the MCR ≤ 2 . Column MCR > 2 shows the percentage of tail exposure due to individual(day)s with multiple substances with MCR > 2.

Tail %	% with MCR = 1	Sub- stances	% with 1 < MCR<=2	Substances	% with MCR > 2	Sub- stances
37	20.6	Difeno, Tebu	73.7	Difeno, Tebu	5.7	Difeno, Tebu
50	19.2	Difeno, Tebu	75.6	Difeno, Tebu	5.2	Difeno, Tebu
90	16.3	Difeno, Tebu	78.8	Difeno, Tebu	5.0	Difeno, Tebu
95	15.0	Difeno, Tebu	82.5	Difeno, Tebu	2.5	Difeno, Tebu
99	25.0	Difeno	75.0	Difeno, Tebu Propi	0.0	

Table 3.240:	Maximum	Cumulative	Ratio	summary
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To configure the MCR plot, see *dietary exposures settings, human monitoring analysis settings* and *internal exposures settings* with options to display the ratio total exposure/ maximum for individual(day) exposures (MCR plot), to specify tail percentiles of the exposure distribution, e.g. 95, 97.5 and 99% (MCR plot) or to set the minimum percentage contribution per substance to the tail exposure (MCR plot).

Internal exposures settings

Calculation settings

Name	Туре	Description
Exposure type	ExposureType	The type of exposure considered in the assessment; acute (shor term) or chronic (long-term).
Target level	TargetLevelType	Select whether the assessment is at the level of external dose, internal (target) concentration or internal (systemic) dose. For aggregate assessment, combining dietary and nondietary exposures, the target level is always internal concentration or internal (systemic) dose. When only dietary exposures are available, the target level is optional and may be an external (administered) dose, an internal (tissue) concentration or an internal (absorbed) dose.
Multiple substances analysis	Boolean	Specifies whether the assessment involves multiple substances.
Compute cumulative exposures	Boolean	Specifies whether the assessment involves multiple substances a results should be cumulated over all substances.
Specify the type of kinetic model	InternalModelType	Specify the type of model to convert external exposure to the internal level.
Biological matrix	BiologicalMatrix	The target biological matrix (internal compartment) for which results are computed. Biological matrices from kinetic conversions will become available after selecting a data source the kinetic models module.
Exposure sources	ExposureSource	Specifies the exposure sources involved in the exposure assessment.
Exposure routes	ExposureRoutes	Exposure routes included in assessment.
Individual reference set	ExposureSource	Specifies the exposure sources used as reference set for matchi individuals.

T_11_ 2 241.	Calardation	a	£	- 11-	Lot a man al	
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Name	Туре	Description
Match individuals of multiple	Boolean	If checked, then non-dietary exposures of individuals from
non-dietary surveys		different surveys with the same identifier (ID) are combined.
Substance weighting in	ExposureApproachType	Risk based: exposures in equivalents of the reference substance
mixtures		standardised: standardised exposures per substance have varian
		1; or unweighted exposures: RPFs are equal to 1.
Perform MCR analysis	Boolean	Perform a Maximum Cumulative Ratio (MCR) analysis to
		determine co-exposure between substances.
Display ratio total exposure/	Numeric	For MCR plot: specify ratio total exposure/ maximum for
maximum (in MCR plot)		individual(day) exposures .
Show tail percentiles (MCR	Numeric	Give specific percentiles of exposure distribution (%), e.g. 97.5
piot) for:	Numaria	99 (space separated).
contribution per substance to	Numeric	set minimum percentage contribution per substance to the tail
the tail exposure (MCR plot)		exposure.
Cutoff MCR	Numeric	For selection of individual(day) exposures with maximum
	1 (unione	cumulative ratio (MCR = total exposure/maximum) above the
		cutoff.
Cutoff percentage in	Numeric	For selection of individual(day) exposures above the cutoff
population ranked on total		percentage in the set of individual(day)s ranked on total exposu
exposure		
Seed for pseudo-random	Numeric	A value of 0 will use a pseudo-random seed in each run, a value
number generator		0 will provide the same results in a repeated run.
Standardised blood	Boolean	Standardised blood concentrations (lipids).
concentrations (lipids)		
Standardised (creatinine) or	Boolean	Standardised (creatinine) or normalised (specific grafity) urine
normalised (specific grafity)		concentrations.
Selected expression type	ExpressionTune	Standardised uring (graatining) or normalised uring (specific
Selected expression type	Expression1 ype	grafity)
Exposure events and dose	ExposureEventsGeneration-	Generates doses and events for PBK model
generation method	Method	Constants, doses and events for f bit model
Number of simulated days	Numeric	The total duration (in days) of the PBK model simulations.
Number of initial days skipped	Numeric	The initial period (or burn-in period, specified in days) that nee
		to be skipped when calculating the steady-state or peak internal
		exposures from the simulated time series.
Use PBK model parameter	Boolean	When specified, use parameter variability.
variability		
Number of events per day for	Numeric	The daily dose is administered in equal portions (dose / number
the DERMAL dose	NT .	events) per event.
Number of events per day for	Numeric	The daily dose is administered in equal portions (dose / number
Number of quests per day for	Numaria	The dealy does is administered in equal particips (does / number
the ORAL dose	Numeric	events) per event
Specify the hours (events)	Numeric	Specify the hours on which exposure events are simulated
speeny the notats (crems).	1 (unitorite	Allowed numbers are 1, 2, 3, 4,, 24 separated by spaces.
Specify event timings	Boolean	If checked, a specific sequence of hours on a day can be specifi
		on which exposure events are simulated. Otherwise the events a
		derived based on the number of exposure events per day.
PBK simulation method	PbkSimulationMethod	Specifies in which mode to run the PBK model
Expected life time in years	Numeric	The total expected lifetime (in years) of the PBK model.
Use bodyweight adjusted	Boolean	Use bodyweight adjusted exposures in PBK model simulations.
exposures		
DIET: Population alignment	PopulationAlignmentMethod	Population aligment methods for dietary exposure sources.
method diet exposures		

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Name	Туре	Description
Diet: match at random based on sex	Boolean	Match at random based on stratifying cofactor sex.
Diet: match at random based	Boolean	Dietary exposures are randomly matched on stratifying covarial
Diet: matching method for age	AgeAlignmentMethod	Match at random based on age using user defined bins (in years or fit to closest available set of ages.
Diet: specify bins:	Numeric	Specify bins, e.g. 10 25 50 (space separated) in years.
AIR: Population alignment method (RSExpo) air exposures	PopulationAlignmentMethod	Population aligment methods for (RSExpo) air exposure source
Air: match at random based on sex	Boolean	Match at random based on stratifying cofactor sex.
Air: match at random based on age	Boolean	Air exposures are randomly matched on stratifying covariable a
Air: matching method for age	AgeAlignmentMethod	Match at random based on age using user defined bins (in years or fit to closest available set of ages.
Air: specify bins:	Numeric	Specify bins, e.g. 10 25 50 (space separated) in years.
SOIL: Population alignment method (RSExpo) soil exposures	PopulationAlignmentMethod	Population aligment methods for (RSExpo) soil exposure source
Soil: match at random based on sex	Boolean	Match at random based on stratifying cofactor sex.
Soil: match at random based on age	Boolean	Soil exposures are randomly matched on stratifying covariable age.
Soil: matching method for age	AgeAlignmentMethod	Match at random based on age using user defined bins (in years or fit to closest available set of ages.
Soil: specify bins:	Numeric	Specify bins, e.g. 10 25 50 (space separated) in years.
DUST: Population alignment method (RSExpo) dust exposures	PopulationAlignmentMethod	Population aligment methods for (RSExpo) dust exposure source
Dust: match at random based on sex	Boolean	Match at random based on stratifying cofactor sex.
Dust: match at random based on age	Boolean	Dust exposures are randomly matched on stratifying covariable age.
Dust: matching method for age	AgeAlignmentMethod	Match at random based on age using user defined bins (in years or fit to closest available set of ages.
Dust: specify bins:	Numeric	Specify bins, e.g. 10 25 50 (space separated) in years.
NON-DIETARY: Population alignment method other non-dietary exposures	PopulationAlignmentMethod	Population aligment methods for nondietary exposure sources.
Nondietary: match at random based on sex	Boolean	Match at random based on stratifying cofactor sex.
Nondietary: match at random based on age	Boolean	Nondietary exposures are randomly matched on stratifying covariable age.
CONSUMER-PRODUCT: Population alignment method consumer product exposures	PopulationAlignmentMethod	Population aligment methods for consumer product exposure sources.
Consumer product: match at random based on sex	Boolean	Match at random based on stratifying cofactor sex.
ConsumerProduct: match at	Boolean	Consumer product exposures are randomly matched on stratify
random based on age		covariable age.
ConsumerProduct: matching method for age	AgeAlignmentMethod	Match at random based on age using user defined bins (in years or fit to closest available set of ages.
Consumer product: specify bins:	Numeric	Specify bins, e.g. 10 25 50 (space separated) in years.

continues on next pa

NameTypeDescriptionNumber of initial years skippedNumericThe initial period (or burn-in period, specified in years) that ne to be skipped when calculating the steady-state or peak internal exposures from the simulated time series.	Table 0.241 Continued from previous page			
Number of initial years skippedNumericThe initial period (or burn-in period, specified in years) that ne to be skipped when calculating the steady-state or peak internal exposures from the simulated time series.	Name	Туре	Description	
	Number of initial years skipped	Numeric	The initial period (or burn-in period, specified in years) that ne to be skipped when calculating the steady-state or peak internal exposures from the simulated time series.	

Table 3.241 – continued from previous page
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Output settings

Table 3.242: Output settings for module	Internal exposures.
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Name	Туре	Description
Exclude privacy sensitive data from outputs	Boolean	Use this setting to not report the parts of the results (i.e., figure tables, or sections) that are marked as (potentially) privacy sensitive.
Include drill-down on 9 individuals around specified percentile	Boolean	Specifies whether drilldown on 9 individuals is to be included in the output.
Store simulated individual day exposures	Boolean	Store the simulated individual day exposures. If unchecked, no additional output will be generated. If checked, the output will contain an additional section with the simulated individual day exposures.
Show percentiles for	Numeric	Give specific percentiles of exposure distribution (%), e.g. 50 9 95 97.5 99 (space separated).
Percentage for drilldown	Numeric	Gives detailed output for nine individuals near this percentile of the exposure distribution.
Percentage for upper tail	Numeric	Gives detailed output for this upper percentage of the exposure distribution.
Show % of population below level(s)	ExposureMethod	This setting is used for reporting the percentages of individuals (chronic) or individual days (acute) exceeding certain exposure levels. These exposure levels can be generated automatically based on the observed exposures (Automatic, default) or specific explicitly (Manual).
Exposure levels	Numeric	Specify exposure levels for which to give the percentage of exposure below these levels, e.g. 1 10 50 100 200 500.
Number of levels of covariable to predict exposure	Numeric	Specify the number of levels, e.g. 20. The range of the covaria is divided by the number of levels: range = (max - min)/levels. For these covariable levels exposures are predicted.
Predict exposure at extra covariable levels	Numeric	Specify specific prediction levels in addition to the automaticall generated prediction levels (space separated).
Lower percentage for variability (%)	Numeric	The default value of 25% may be overruled.
Upper percentage for variability (%)	Numeric	The default value of 75% may be overruled.
Report consumptions and exposures per individual instead of per kg body weight	Boolean	Specifies whether body weights should be ignored and consumptions and exposures should be expressed per individua Otherwise, the consumptions and exposures are per kg body weight.

Uncertainty settings

	, ,	1
Name	Туре	Description
Resample kinetic conversion factors	Boolean	Specifies whether kinetic conversion factors are resampled.
Resample PBK model parameter values	Boolean	Specifies whether PBK model parameter values are resampled.
Lower uncertainty limit (%)	Numeric	Percentage lower bound, e.g. 2.5%.
Upper uncertainty limit (%)	Numeric	Percentage upper bound, e.g. 97.5%.

Table 3.243: Uncertainty settings for module Internal exposures.

Calculation of internal exposures

Internal exposures are computed by linking dietary and (if available) non-dietary individual/individual-day exposures. The (aggregated) internal exposure is computed as an internal concentration at the specified target compartment using kinetic conversion factors or PBK models or as an internal dose that is systemically available in the general circulation using absorption factors.

• Internal exposures calculation

Inputs used: Dietary exposures Non-dietary exposures Dust exposures Soil exposures Active substances Relative potency factors Absorption factors Kinetic conversion factors PBK model parametrisations Air exposures Consumer product exposures

Settings used

• Calculation Settings

3.4.10 Exposure mixtures

Exposure mixtures will select sets of co-occurring substances (one or more) that contribute most to the overall exposure patterns.

This module has as primary entities: Substances Effects

Exposure mixtures calculation

The most common model of cumulative risk assessment is to focus on substances that belong to the same common assessment groups (CAG). *Substances* in such a group belong to the same chemical family and may or may not have a similar mode of action. In assessing the risk, possible interactions between substances are often ignored and, moreover, little information is available about synergistic effects at low doses. More information is needed about the combined effects of such substances, but it is impractical to investigate all possible mixtures, and therefore instruments are needed to select the most relevant substances for further investigation. This selection should not only be based on the hazard (highest relative potencies) but also on the exposure of the population to these substances. The potential risk of being exposed to chemicals in a mixture depends on the food *consumption* patterns of *individuals* in a population. A regular diet can contain hundreds of substances, so the number of combinations of substances to which an individual in a population is exposed can be numerous. The exposures mixtures module is used to identify the most relevant components to which a population is exposed.

Risk based, standardised or unweighted exposures

Before performing the mixture exposure assessment, the data are preprocessed. Three optional choices are available, *see settings, exposure approach type*.

• Risk based exposures: exposures are multiplied by the *relative potency factor* (RPF) of each substance and thus exposures for different substances are on the same and comparable scale.

- Standardised exposures: all exposures are standardised to equal variance (unit variance).
- Unweighted exposures: exposures are taken as such, this is equivalent to RPF s equal to 1 for each substance.

Exposure mixtures are identified using a quantitative approach: *sparse non-negative matrix underapproximation* (*SNMU*) (Gillis and Plemmons (2013)) and answers the question what combination of substances predominantly determine the underlying patterns in the exposure matrix (substance x person (day)).

After identifying the components, a cluster analysis is applied to group individuals with similar profiles of exposure to the obtained component (Crépet et al. (2022)).



Transformed dietary exposure distribution (96.6% positives)

Figure 3.50: Example of co-exposure distribution (from >1 substance per individual-day, red) super-imposed on the total exposure distribution (blue).

Sparse nonnegative matrix underapproximation

Starting point to identify major components of combinations of substances using exposure data was to use Nonnegative Matrix Factorization (NMF). This algorithm was proposed by Lee and Seung (1999) and Saul and Lee (2002) and deals specifically with non-negative data that have excess zeros such as exposure data. Zetlaoui et al. (2011), introduced this method in food risk assessment to define diet clusters.

The NMF method was then implemented by Béchaux et al. (2013) in order to identify food consumption patterns and main components of combinations of pesticides to which the French population was exposed using *TDS* exposure to 26 priority pesticides.

At the start of the Euromix project ideas had evolved: to obtain less substances per component experiments were made using Sparse Nonnegative Matrix Factorization (SNMF) (Hoyer (2004)). This method was found not to give stable solutions. Better results were obtained with Sparse Nonnegative Matrix Underapproximation (SNMU) (Gillis and Plemmons (2013)). This model also fits better to the problem situation because also the residual matrix after extracting some components should be nonnegative. The SNMU method has been implemented in MCRA.

Indeed, NMF may be used to identify patterns or associations between substances in exposure data. NMF can be described as a method that finds a description of the data in a lower dimension. There are standard techniques available

such as principal components analysis or factor analysis that do the same, but their lower rank representation is less suited because they contain negative values which makes interpretation hard and because of the modelling with a Gaussian random vectors which do not correctly deal with the excess of 0 and non-negative data. The NMF solution approximates a non-negative input matrix (i.c. exposure data) by two constrained non-negative matrices in a lower dimension such that the product of the two is as close as possible to the original input matrix. In Figure 3.51, the exposure matrix M with dimensions m (number of substances) and n (number of intake days or persons) is approximated by matrix U and V with dimensions $(m \times k)$ and $(k \times n)$ respectively, where k represents the number of components. Matrix U contains weights of the substances per component, matrix V contains the coefficients of presence of components in exposure per intake day or person. M is non-negative (zero or positive) and U and V are constraint to be non-negative. The minimization criterion is: $||M-UV||^2$ such that $U \ge 0$ and $V \ge 0$.



Figure 3.51: NMF approximation of exposure data

The solution found by NMF contains zeros, but components still contain many substances which complicates interpretability. Therefore, the Sparse Nonnegative Matrix Underapproximation (SNMU) (Gillis and Plemmons (2013)) which also provide sparse results was investigated. The SNMU has also some nice features well adapted to the objective of the Euromix project: the solution is independent of the initialization and the algorithm is recursive. Consequently, the SNMU method which is based on the same decomposition process as the NMF was the one implemented in MCRA.

SNMU is initialized using an optimal nonnegative rank-one approximation using the power method (https://en. wikipedia.org/wiki/Power_iteration). This initialization is based on a singular value decomposition and results in general in a unique solution that is sparse. The SNMU algorithm is called recursive because after identifying the first optimal rank-one underapproximation u_1v_1 , the next rank-one factor is recovered by subtracting u_1v_1 from M and applying the same factorization algorithm to the remainder $M - u_1v_1$. The solution u_1v_1 is called a rank-one underapproximation because an upper bound constraint is added to ensure that the remainder $M - u_1v_1$ is non-negative. For Matlab code see: https://sites.google.com/site/nicolasgillis/code.

For each component, a percentage of explained variance is calculated. M is the exposure matrix with m rows (substances) and n columns (individuals or individual days) S_t is sum of squared elements of M:

$$S_t = ||M||^2 = \sum_{i,j}^{m,n} e_{i,j}^2$$

Apply SNMU on M, then for the first component:

- u is $m \times 1$ vector, contains weights of the component.
- v is $1 \times n$ vector, contains presence of component in exposure per individual.

Calculate residual matrix R:

$$R = M - uv$$

1

Calculate S_r , residual sum of squared elements of R:

$$S_r = ||R||^2 = \sum_{i,j}^{m,n} e_{i,j}^2$$

Percentage explained variance first component (k = 1) is:

$$V_k = (1 - S_r)/S_t) \cdot 100$$

For the second component:

- 1. continue with residual matrix R (replace M by R),
- 2. estimate u and v,
- 3. calculate new residual matrix R
- 4. calculate S_r , residual sum of squared elements of R

Percentage explained variance second component (k = 2) is:

$$V_k = (1-S_r)/S_t) \cdot 100 - \sum_{l=1}^{k-1} V_l$$

The last term is de percentage explained variance of the first component. Continue with the third component etc....

The given factorization by SNMU is not unique. A matrix D and its inverse can be used to transform the two lower rank matrices U and V by, e.g.

 $M = UV = UDD^{-1}V$

where matrix D is non-negative and corresponds to a scaling of matrix U and V to matrix $\tilde{U} = UD$ and $\tilde{V} = D^{-1}V$.

In Figure 3.52, SNMU is applied on the exposure matrix. The SNMU solution after scaling results in a matrix that represents the mixture composition and a matrix representing the individual scores. The first matrix is interpreted as the set of contributions of the substances to a component., the second matrix, as the set of exposures of the individuals to the mixtures.



Figure 3.52: SNMU approximation of exposure data after scaling.

Exposure matrix

Basically, exposure is calculated as consumption x concentration. By summing the exposures from the different foods for each substance per person day separately, the exposure matrix for component selection is estimated:

$$y_{ijc} = \frac{\sum_{k=1}^{p} x_{ijk} c_{ijkc}}{bw_i}$$

where y_{ijc} is the exposure to substance c by individual i on day j (in microgram substance per kg body weight), x_{ijk} is the consumption by individual i on day j of food k (in g), c_{ijkc} is the concentration of substance c in food k eaten by individual i on day j (in mg/kg), and bw_i is the body weight of individual i (in kg). Finally, p is the number of foods accounted for in the model. More precisely, for an *acute* or short term risk assessment, the exposure matrix summarises the y_{ijc} with columns denoting the individual-days (ij) and rows denoting the substances (c). Each cell represents the sum of the exposures for a substance on that particular day. For a *chronic* or long term risk assessment, the exposure matrix is constructed as the sum of all exposures for a particular substance averaged over the total number of intake days of an individual (substances x persons). So each row represents a substance and a column an individual. Each cell represents the observed individual mean exposure for a substance for that individual. Note that in the exposure calculation, the concentration is the average of all residue values of a substance.

When *relative potency factors* (RPF) are available then exposures are multiplied by the RPF and thus exposures to the different substances are on the same and comparable scale (toxicological scale). In this case, the selection of the components is risk-based. In some cases, RPFs may not be available. In this case exposure to different substances, even in the same unit, may lead to very different effects. To give all substances an equal weight a priori and avoid scaling effects, a standardisation of the data can be applied as done in Béchaux et al. (2013). In this case, all substances are assigned equal mean and variance, and the selection of the components will work on patterns of correlation only. Then component selection is not risk-based anymore but, what could be called, co-exposure-based.

Finally, in the mixture selection module of MCRA, the SNMU expects RPF data for a risk-based selection. If not available, the user should provide alternative RPF data, e.g. values 1 for a purely exposure-based selection, or lower-tier estimates (e.g. a maximum value on RPF thought possible).

Mechanisms to influence sparsity

Two mechanisms to *influence sparsity* are available. The SNMU algorithm incorporates a sparsity parameter and by increasing the value, the final components will be more sparse (sparsity close to 0: not sparse, many substances; sparsity close to 1: sparse, few substances). The other approach is by using a subset of the exposure matrix based on a cut-off value for the *MCR*. High ratios correspond to high co-exposure, so it is reasonable to focus on these (person) days and not include days where exposure is received from a single substance (ratio close to 1). To illustrate the combined use of MCR and the sparsity parameter, the French steatosis data (39 substances, 4079 persons) are used and person days with a ratio > 5 (see Figure 3.46) are selected yielding a subset of 139 records.

In Figure 3.53, the effect of using a cut-off level is immediately clear. The number of substances of the first component is 17 whereas in the unselected case (not shown) only 4 substances were found The three plots show the influence of increasing the sparsity parameter from 0 to 1 on the number of substances in the component. The first plot shows the estimated sparsity versus the specified sparsity. The second and third plot the the number of substances in a component. For values close to 0, the component contains 17 substances. For values > 0.4 the number of substances in the component. The first plot shows in the component drops to 3 and for a sparsity close to 1, only one substance is found in a component.



Sparseness for 39 compounds



Number of compounds 39





Figure 3.53: Influence of the specified sparsity parameter on the realized sparsity, n = 139.

Substance contributions to components

The SNMU solution of matrix U can be displayed in a heatmap. The heatmap shows the relative contribution of each substance to a component.





In Figure 3.55 and Figure 3.56 the sparsity parameter is set to 0.1 (not sparse) and 0.8 (sparse), respectively. This leads to components containing different number of substances.

In Figure 3.57, the relative contributions of the substances to the first component are displayed in a piechart.

As mentioned before, one of the nice features of the SNMU algorithm is its recursive character which results in identical components. In Figure 3.58, the U matrix is visualized using three components. Compare this solution with Figure 3.56, the first three components are identical. Because of the ordering the plots look slightly different, but a closer inspection of the first 3 components of each solution shows that they are the same. In both figures, component 1 contains mbzp, A, B, C and D; component 2, mibp, E, F and ohmehp; and component 3 mnbp and G.

In paragraph network analysis, an alternative to the SNMU approach is proposed.

For selection of individual(day) exposures with a *maximum cumulative ratio* above a cutoff and/or above a cutoff percentage in the set of individual(day)s ranked on total exposure, see 'Cutoff MCR' and 'Cutoff percentage' settings.

Component exposures in population and subgroups

The SNMU solution of matrix V is used to group individuals with similar mixture exposure profiles. In Figure 3.59, the idea of clustering is shown.

Crépet et al. (2022) propose to identify components by coupling statistical criteria with the relevance of combined exposure profiles and component composition. First, the optimal choice for k, the number of components, is determined using a trade off between the decrease of the residual sum of squares and number of components. Then, hierarchical clustering was applied to the matrix of individual scores V to group individual(day)s with similar exposure profiles to the k components. This identification of components is repeated for different values of k where inspection of components not relevant to characterize a cluster, or concerned with only a small part of the population leads to rejection of the mixture.

In MCRA, two clustering methods are availabe. The first, hierarchical clustering, is implemented as described in Crépet et al. (2022). Ward's clustering criterion is implemented using Euclidean distances (Ward.D2, Murtagh and



Figure 3.55: Co-exposure of substances. Heatmap for a solution with 6 components. The sparsity = 0.1. Each component, especially the first, contains many substances (see also Figure 3.56).



Figure 3.56: Co-exposure of substances. Heatmap for a solution with 6 components. The sparsity = 0.8. Components contain less substances compared to Figure 3.55.



Figure 3.57: Relative contributions substances to component 1. The sparsity is set to 0.8 (sparse), estimated sparsity = 0.62.



Figure 3.58: Heatmap for solution with 3 components. The first 3 components in Figure 3.58 and Figure 3.56 contain the same substances.


Figure 3.59: SNMU: matrix V, individual scores to components.

Legendre (2014)). Specification of the optimal number of clusters is not needed. Results of the clustering are displayed in a dendrogram, Figure 3.60. The second one, based on K-means, requires specification of the number of clusters. The results of the clustering are represented in a scatter plot using principal components and convex envelopes to identify the clusters, Figure 3.62.

In Figure 3.61, the relative exposure to componaents in the total populations are shown. These plots are also available for the subgroups resulting from the clustering.

Advantages of K-means clustering is that it is simple and fast and large datasets can be handeled. Visualisation for large data sets is straightforward but for hierarchical clustering dendrograms maybe very dense. Disadvantage of K-means is that it requires the number of clusters set before. For large datasets, hierarchical clustering maybe slow, $O(n^2)$, but for small datasets, the dendrogram helps in interpreting the results and in selecting the optimal number of clusters.

Network analysis

As an alternative to *SNMU*, a network analysis is proposed. The outcome of a network analysis is graphically displayed by a network. This is a collection of nodes, that may have pairwise relationships. Each node represents a substance, and an edge represents pairwise dependence between substances (e.g. correlation or partial correlation). In MCRA, the network is estimated using a Gaussian graphical model (GLASSO) based on partial correlation and a sparseness penalty to control the number of nonzero edges (Friedman et al. (2008)). Parameters are automatically tuned. The communities are detected using a Walkman algorithm. In Figure 3.63, using HBM data, a network is displayed with 6 communities. The largest community contains 5 substances: mbzp, A, B, C and D. Compared to the results of the *SNMU mixture analysis*, the communities are almost identical to the components found in the SNMU approach.

The exposure data may be log transformed before the network analysis. Zeros are replaced by the logarithm of the minimum of the non-zero exposure values per substance multiplied by a factor 0.01.

Exposure mixtures settings



Figure 3.60: Hierarchical clustering of human monitoring data, 3 clusters, largest and smallest cluster contain 152 and 37 individuals, respectively



Figure 3.61: Relative exposure to components in the population

Calculation settings

Name	Туре	Description
Exposure type	ExposureType	The type of exposure considered in the assessment; acute (short term) or chronic (long-term).
Target level	TargetLevelType	Select whether the assessment is at the level of external dose, internal (target) concentration or internal (systemic) dose. For a aggregate assessment, combining dietary and nondietary exposures, the target level is always internal concentration or internal (systemic) dose. When only dietary exposures are available, the target level is optional and may be an external (administered) dose, an internal (tissue) concentration or an internal (absorbed) dose.
Exposure calculation method	<i>ExposureCalculationMethod</i>	Method for obtaining exposure estimates. These can be modell exposures (e.g., external (dietary) exposures or internal exposure estimates obtained by aggregating dietary and non-dietary exposures) or exposure estimates derived from human (bio)monitoring data.
Report consumptions and exposures per individual instead of per kg body weight	Boolean	Specifies whether body weights should be ignored and consumptions and exposures should be expressed per individual Otherwise, the consumptions and exposures are per kg body weight.
Sampling method	AlphaNumeric	The sampling method that should be included in the action.
Substance weighting in mixtures	ExposureApproachType	Risk based: exposures in equivalents of the reference substance standardised: standardised exposures per substance have varian 1; or unweighted exposures: RPFs are equal to 1.
Number of components	Numeric	The number of components to select in SNMU.
Sparseness parameter 3.4. Exposure modules	Numeric	Sparseness parameter. Value between 0 (not sparse, many substances) and 1 (sparse, few substances).
Iterations	Numeric	Maximum number of iterations, e.g. 1000.
Seed for pseudo-random number generator.	Numeric	Random seed for initialising matrix W and H.

Table 3.244: Calculation settings for module Exposure mixtures.



Figure 3.62: K-means clustering of human monitoring data, 3 clusters, largest and smallest cluster contain 204 and 21 individuals, respectively



Figure 3.63: Network analysis.



Figure 3.64: SNMU solution with 6 components. Sparsity = 0.8.

Calculation of exposure mixtures

Exposure mixtures or components can be computed from (external) dietary exposures, or from (internal) exposures (possibly from combined dietary- and non-dietary sources) or human monitoring concentrations. A multivariate decomposition method, sparse non-negative matrix underapproximation (SNMU), is applied to the matrix of exposures per substance and per individual (chronic) or individual-day (acute) to find component containing substances that contribute most to the cumulative exposure. Exposures per substance are preprocessed either by multiplication with relative potency factors (RPFs) to make the analysis risk-based, or by standardisation to variance 1 to make the analysis correlation-based. An alternative to SNMU is network analysis. This method estimates communities of substances that have pairwise relationships.

• Exposure mixtures calculation

Inputs used: Dietary exposures Internal exposures Relative potency factors HBM analysis

Settings used

• Calculation Settings

3.4.11 Food conversions

Food conversions relate foods-as-eaten, as found in the consumption data, to modelled foods (foods-as-measured), which are the foods for which concentration data are available. A food-as-eaten can be linked to one, or multiple modelled foods using various conversion steps (e.g., using food recipes to translate a composite food into its ingredients). There are several types of conversion steps, and a conversion path may comprise multiple conversion steps between a food-as-eaten and a modelled food.

This module has as primary entities: Foods Substances

Output of this module is used by: Consumptions by modelled food Dietary exposures

Food conversions calculation

Food conversions are computed using a recursive search algorithm to link foods-as-eaten to modelled foods, possibly through intermediate conversion steps. For instance, if (unpeeled) apple and grapes are the modelled foods, the foodas-eaten apple pie contains peeled apple and raisins, peeled apple is linked to unpeeled apple, and raisins are dried grapes. Hence, for apple pie, there are two conversions, one to apple (with processing type 'peeled') and one to grapes (with processing type 'dried'), each with its own conversion path of intermediate conversion steps.

Substance independent conversion

The current implementation of the food conversion algorithm can be run substance independent. The *Find processing link (deprecated)* is skipped from the algorithm (default = false) and is only retained for backwards compatibility reasons only (see Advanced, set to true). Processed foods are easily recognized in the food translation step and retrieving the processing factor that belongs to a certain processing type is done outside the algorithm. In fact, finding processing types with corresponding processing factors is not a task of the conversion algorithm: conversion is about converting food codes to other food codes.

When the processing step is skipped, there is no need to run the conversion algorithm on a substance basis. The only information that is needed is whether a food code is a modelled food or not (i.c. is there a concentration available or not). This information can be computed beforehand: for each substance all modelled foods are collected and supplied to the conversion algorithm in a common dictionary containing all modelled food codes. As soon as a food code is found in the dictionary, the conversion ends and the next food code is converted.

For each food-as-eaten, the food conversion algorithm recursively builds up the conversion paths using the following procedure:

1. *Substance independent conversion:* the conversion algorithm is substance independent. Check whether the current food is a modelled food. If successful, the food has been found, and the current search stops.

- 2. Check whether the current food translates to one or more foods through composition or read-across. Identify any processing types of facets.
- a. *Food recipe link:* try to find food translations for the current food (i.e., the ingredients of a composite food). This may result in one or more food codes for ingredients, and the iterative algorithm will proceed with each of the ingredient food codes in turn. Simultaneously check, whether the current food is a processed food or not. If so, determine the processing type or facets.
- b. *TDS food sample composition link:* try to find the code in the *TDS food sample compositions* table (column idFood), a default translation proportion of 100% is assumed. The iterative algorithm will proceed with a TDS food (column idTDSFood) sample.
- c. *Read-across link:* try to find a food extrapolation rule for the current food, a default translation proportion of 100% for 'idToFood' is assumed.

Note that in the *food recipe link* processed foods are recognized and that the translation proportion to correct for a weight reduction or increase is stored.

If successful, restart at the first step with each of the new codes of the ingredient foods, TDS foods or Read Across foods.

- 3. *Marketshares link:* try to find subtype codes, e.g. 'xxx\$*' in the *Market shares* table. In general, marketshares should sum to 100%. Foods with marketshares not summing to 100% are ignored in the analysis unless the checkbox *Allow marketshares not summing to 100%* is checked. This step is optional, see advanced settings. If successful, restart at step 1 with each of the new codes of the subtype foods.
- 4. *Supertype link:* try to find supertypes, e.g. 'xxx\$yyy' is converted to 'xxx'. This step is optional, see advanced settings if you want to use this. If successful, restart at step 1 with the new code of the supertype food.
- 5. *Default processing factor:* remove processing part (-xxx) of the code. If successful, restart at step 1 with the new code without processing part.
- 6. Check whether the food is a FoodEx 2 code. If so, strip all food facets of the code. The code that remains i.e the food base code of FoodEx 2 classification is returned and the algorithm restarts at step 1 with the base code (without facets).

Substance dependent conversion

The original conversion algorithm contains two steps which are substance dependent. For each substance all food codes are supplied to the conversion algorithm and for each food code it is checked whether there is:

- a concentration,
- a processing type.

When a concentration is available for the food, this food is a modelled food (formerly known as food as measured). The food may be a food as eaten as such, like apple, or an ingredient of a food as eaten like tomato sauce on pizza which is converted to tomato. If concentrations are available, the food code is found and the conversion algorithm starts with converting the next food code. Otherwise, the conversion proceeds to the *processing link (deprecated)*. Here, basically, processed foods are converted to an unprocessed food and processing type with corresponding processing factor. This processing step may be substance specific and, occasionally, this results in different conversion paths for different substances. This is undesirable behaviour and normally not the case (dependent on the supplied data in the *Processing factors* table). However, on rare occasions this might happen.

Find processing link (deprecated): Check whether the current food can be considered to be a processed variant (e.g., cooked or peeled) of another food.

Match processing factor: try to find the code in the Processing factors table.

If successful, try to find the corresponding food translation proportion in the food recipes data to correct for a weight reduction or increase. Then, restart at the first step with the new code of the unprocessed food.

Warning: the *find processing link (deprecated)* step is not recommended and is currently maintained for backwards compatibility reasons only. Finding different conversions paths depending on the substance is undesirable behaviour.





Food conversion settings

Calculation settings

Ī	Name	Туре	Description
	Selected tier	SettingsTemplateType	Specifies all module settings should be set according to a pre-defined tier or using custom settings.
l	Multiple substances analysis	Boolean	Specifies whether the assessment involves multiple substances.
	Allow conversion using processing info	Boolean	Try to find the code in the processing table. When a match is found, then also a food translation record is searched for to also include information on weight reduction/increase (translation proportion). Warning: the processing step is deprecated and is currently only maintained for backwards compatibility reasons.
	Allow conversion using food translations	Boolean	Try to find food translations for the current food (i.e. the ingredients of a composite food). This may result in one or most food codes for ingredients, and the iterative algorithm will proceed with each of the ingredient food codes in turn.
	Total diet study concentration data	Boolean	Specifies whether exposure is based on sampling data from tota diet studies.
	Allow conversion using food extrapolations	Boolean	Try to find read across codes. If unchecked, read across table is ignored, default is 'Use read across info'. E.g. for pineapple no measurements are found but by specifying that pineapple is converted to FruitMix (with a default proportion of 100%), the TDS sample concentration value of FruitMix will be used for pineapple (as-eaten or as ingredient).
	Allow conversion using market shares	Boolean	Try to find subtype codes, e.g. 'xxx\$*' in the market shares tabl
	Allow market shares not summing to 100%	Boolean	Specify whether to rescale market share percentages that do no sum to 100%. If checked, then foods with marketshares not summing to 100% are allowed. If not, then these foods are ignored in the analysis.
	Allow conversion to supertypes	Boolean	Try to find supertypes, e.g. 'xxx\$yyy' is converted to 'xxx'. If checked, allows for linkage of consumed foods coded at a lower hierarchical level to foods with measured concentrations at a higher hierarchical level e.g. consumed is Apple (code PF\$App -> measured is Pome Fruit (code PF). Note: food codes are sp on '\$'. Measurements of substances on food are available at a le detailed food coding level than consumption data.
	Allow conversion using default processing	Boolean	Allow conversion from processed to unprocessed foods. Either matching to processing types specified as substrings separated using a '-' character or by matching FoodEx2 facet strings (separated by a '#' character) to processing types.
	Conversion is substance independent	Boolean	Conversion of foods is independent of the substance.
	Derive modelled foods from concentration limits	Boolean	Derive modelled foods from concentration limits.
	Include foods with only censored value measurements	Boolean	Specifies whether foods with only censored value measurement are part of the exposure assessment (default yes).
	Include substances with only censored value measurements	Boolean	Specifies whether substances with only censored value measurements are part of the exposure assessment (default yes)
	Include substances without measurements	Boolean	Specifies whether substances without any measurements should included.

Table 3.245: Calculation settings for module Food conversions.

Food conversions tiers

Overview

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFSA 2022 Acute Tier II	EFS 2022 Chro Tier
T ta di st co co tr ti di	false	false	false	false	false	false	false	false	false	false
Ig ne sa pl	true	true	true	true			true	true	true	true
E cl ir di vi u al w le th N d	false	true	false	true			false	true	false	true
F te sa pl es ca in th ca ca tr ti lii it	false	false	false	false		false	false	false	false	false
U st co vo si rt	true	true	true	true			true	true	true	true
S st	UseMost- Toxic	UseMost- Toxic	DrawRan- dom	DrawRan- dom			UseMost- Toxic	UseMost- Toxic	DrawRan- dom	Drav dom
428 si m			3011				Chapte	r 3. Module	som S	com
R	true	true	true	true			true	true	true	true

Table 3.246: Tier overview for module Food conversions.

430

Retrospective dietary CRA (EC 2018) - Acute / Tier I

Name	Setting	From input tier	In module	
Total diet study concentration data	false			
Ignore sampling weights	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Consump- tions	
Exclude individuals with less than N days	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Consump- tions	
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations	
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations	
Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations	
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations	
Account for substance authorisations in substance conversions	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations	
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive	Concen- trations	
		CRA (EC 2018) - Acute / Tier I	Chapte	r 3. Mo

Table 3.247: Tier definition for Retrospective dietary CRA (EC 2018) - Acute / Tier I.

Retrospective dietary CRA (EC 2018) - Chronic / Tier I

Name	Setting	From input tier	In module	
Total diet study concentration data	false			
Ignore sampling weights	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Consump- tions	
Exclude individuals with less than N days	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Consump- tions	
N (number of days in survey)	2	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Consump- tions	
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations	
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations	
Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations	
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations	
Account for substance authorisations in substance conversions	false	Retrospec- tive	Concen- trations	
		atetary CRA (EC 2018) - Chronic /	Chapter	3. Modules

Table 3.248: Tier definition for Retrospective dietary CRA (EC 2018) - Chronic / Tier I.

Retrospective dietary CRA (EC 2018) - Acute / Tier II

Name	Setting	From input tier	In module
Total diet study concentration data	false		
Ignore sampling weights	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Consump- tions
Exclude individuals with less than N days	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Consump- tions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive	Concen- trations
		CRA (EC 2018) - Acute / Tier II	Chapte

Table 3.249: Tier definition for Retrospective dietary CRA (EC 2018) - Acute / Tier II.

Retrospective dietary CRA (EC 2018) - Chronic / Tier II

Name	Setting	From input tier	In module	
Total diet study concentration data	false			
Ignore sampling weights	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Consump- tions	
Exclude individuals with less than N days	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Consump- tions	
N (number of days in survey)	2	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Consump- tions	
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations	
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations	
Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations	
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations	
Account for substance authorisations in substance conversions	true	Retrospec- tive	Concen- trations	
		atetary CRA (EC 2018) - Chronic /	Chapter	3. Module

Table 3.250: Tier definition for Retrospective dietary CRA (EC 2018) - Chronic / Tier II.

Retrospective dietary CRA (EFSA 2012) - Optimistic

Use the optimistic model settings according to the EFSA Guidance 2012. Concentration values are sampled using a sample-based empirical distribution. Available processing factors are applied. No unit variability model should be applied.

Name	Setting	From input tier	In module
Total diet study concentration data	false		
Apply processing factors	true	Retrospec- tive dietary CRA (EFSA 2012) - Optimistic	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EFSA 2012) - Optimistic	Processing factors
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EFSA 2012) - Optimistic	Processing factors

Table 3.251:	Tier definition for	Retrospective	dietary	CRA	(EFSA	2012)
- Optimistic.						

Retrospective dietary CRA (EFSA 2012) - Chronic / Pessimistic

Chronic probabilistic exposure assessment using the pessimistic model settings according to the EFSA Guidance 2012. Only processing factors > 1 are applied.

Name	Setting	From input tier	ln module
Total diet study concentration data	false		
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2012) - Chronic / Pessimistic	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EFSA 2012) - Chronic / Pessimistic	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EFSA 2012) - Chronic / Pessimistic	Processing factors
Ignore processing factors less than 1	true	Retrospec- tive dietary CRA (EFSA 2012) - Chronic / Pessimistic	Processing factors

Table 3.252: Tier definition for Retrospective dietary CRA (EFSA 2012) - Chronic / Pessimistic.

Retrospective dietary CRA (EFSA 2022) - Acute / Tier I

	Name	Setting	From input tier	In module
	Total diet study concentration data	false		
	Ignore sampling weights	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Consump- tions
	Exclude individuals with less than N days	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Consump- tions
	Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
	Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
	Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
	Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
	Account for substance authorisations in substance conversions	false	Retrospec- tive dietary	Concen- trations
3.4. Exposu	re modules		CRA (FES A	
-			2022) - Acute /	

Tier 1

Table 3.253: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier I.

Retrospective dietary CRA (EFSA 2022) - Chronic / Tier I

	Name	Setting	From input tier	ln module
	Total diet study concentration data	false		
	Ignore sampling weights	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Consump- tions
	Exclude individuals with less than N days	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Consump- tions
	N (number of days in survey)	2	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Consump- tions
	Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
	Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
	Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
	Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary	Concen- trations
3.4. Exposu	re modules		CKA (EFSA 2022) - Chronic /	

Tier 1

Table 3.254: Tier definition for Retrospective dietary CRA (EFSA 2022) - Chronic / Tier I.

Retrospective dietary CRA (EFSA 2022) - Acute / Tier II

	Name	Setting	From input tier	ln module
	Total diet study concentration data	false	[
	Ignore sampling weights	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Consump- tions
	Exclude individuals with less than N days	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Consump- tions
	Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
	Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
	Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
	Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
	Account for substance authorisations in substance conversions	true	Retrospec- tive dietary	Concen- trations
3.4. Exposu	re modules		CRA (EFSA 2022) -	

Table 3.255: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier II.

Retrospective dietary CRA (EFSA 2022) - Chronic / Tier II

Name	Setting	From input tier	In module
Total diet study concentration data	false		
Ignore sampling weights	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Consump- tions
Exclude individuals with less than N days	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Consump- tions
N (number of days in survey)	2	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Consump- tions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary	Concen- trations
e modules		CKA (EFSA 2022) -	

Table 3.256: Tier definition for Retrospective dietary CRA (EFSA 2022) - Chronic / Tier II.

Prospective dietary CRA (EFSA 2023) - Acute / Tier II

	Name	Setting	From input tier	In module
	Total diet study concentration data	false		
	Ignore sampling weights	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Consump- tions
	Exclude individuals with less than N days	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Consump- tions
	Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
	Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
	Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
	Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
	Retain all allocated substances after active substance allocation	true	Prospec- tive dietary	Concen- trations
3.4. Exposur	e modules		CRA (FES A	
			(EFSA 2023) - Acute /	

Table 3.257: Tier definition for Prospective dietary CRA (EFSA 2023) - Acute / Tier II.

Prospective dietary CRA (EFSA 2023) - Chronic / Tier II

Name	Setting	From input tier	In module
Total diet study concentration data	false		
Ignore sampling weights	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Consump- tions
Exclude individuals with less than N days	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Consump- tions
N (number of days in survey)	2	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Consump- tions
Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Prospec- tive dietary	Concen- trations
e modules		CKA (EFSA 2023) - Chronic / Tier II	

Table 3.258: Tier definition for Prospective dietary CRA (EFSA 2023) - Chronic / Tier II.

Prospective dietary CRA (EFSA 2024) - Acute / Tier II

	Name	Setting	From input tier	In module
	Total diet study concentration data	false		
	Ignore sampling weights	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Consump- tions
	Exclude individuals with less than N days	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Consump- tions
	Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
	Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
	Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
	Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
	Retain all allocated substances after active substance allocation	true	Prospec- tive dietary	Concen- trations
3.4. Exposu	re modules		CRA (FESA	
			2024) - Acute /	

Table 3.259: Tier definition for Prospective dietary CRA (EFSA 2024) - Acute / Tier II.
Prospective dietary CRA (EFSA 2024) - Chronic / Tier II

	Name	Setting	From input tier	In module
	Total diet study concentration data	false		
	Ignore sampling weights	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Consump- tions
	Exclude individuals with less than N days	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Consump- tions
	N (number of days in survey)	2	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Consump- tions
	Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
	Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
	Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
	Substance conversion method	DrawRan- dom	Prospec- tive dietary	Concen- trations
3.4. Exposur	e modules		(EFSA 2024) - Chronic / Tier II	

Table 3.260: Tier definition for Prospective dietary CRA (EFSA 2024) - Chronic / Tier II.

Calculation of food conversions

Food conversions are computed recursively, starting with a food-as-eaten and following a path to ingredients (food recipes), super/sup-type foods, etc. until either arriving at a modelled food (commonly the raw primary commodity) or concluding that the path does not lead to a modelled food.

• Food conversions calculation

Inputs used: Consumptions Modelled foods Processing factors Food recipes Market shares Food extrapolations Total diet study sample compositions Active substances

Settings used

• Calculation Settings

3.4.12 Biological matrix concentration comparisons

Substances in the human body are absorbed, excreted without transformation, excreted after metabolization or stored in various tissues, bones or body fluids. The term biological matrix refers to all human specimens where concentratrions of a chemical can be measured like bodily fluids, such as blood, urine, saliva, breast milk, sweat, and other specimens, such as faeces, hair, teeth, and nails. Biological matrix concentration comparisons compares observed human monitoring data with predictions made for the same population of individuals from dietary survey data, concentration data and (optionally) non-dietary exposure data.

This module has as primary entities: Populations Substances

Biological matrix concentration comparisons calculation

In this module, concentration estimates from measurements in biological (human) matrices are compared with modelled or predicted concentrations obtained from *dietary* and/or *non-dietary* surveys via *exposure* assessments. The comparison provides insight in the coherence between modelled exposures and the measured reality. It is required that monitoring data and dietary/non-dietary exposure data are available for the same individual or individual-day. In Figure 3.66, summary statistics are visualised for the monitoring and modelled concentrations for bisphenols BPA, BPS and BPF.



Figure 3.66: Boxplots for monitoring and modelled concentrations for bisphenols BPA, BPS and BPF. Lower whiskers indicate the p5 and p10 percentiles, upper whiskers the p90 and p95. The edges of the box indicate the p25 and p75 percentiles with the median in the centre of the box.

When both human monitoring data and exposure data are available for the same individuals in a population, a direct comparison can be made between the monitoring and modelled concentrations. In Figure 3.67, an example is shown.



Monitoring versus modelled (p25, p50, p75) exposures BPA

Figure 3.67: Monitoring versus modelled concentrations for bisphenol BPA

Biological matrix concentration comparisons settings

Calculation settings

Table 3.261: Calculation settings for module Biological matrix concentration comparisons.

Name	Туре	Description
Exposure type	ExposureType	The type of exposure considered in the assessment; acute (shor term) or chronic (long-term).
Correlate monitoring with modelled concentrations	Boolean	Correlate monitoring with modelled concentrations. This is opt prevents that monitoring and modelled individuals that are code unintentionally with identical id's or codes are correlated.

Output settings

Table 3.262: Output settings for module Biological matrix concentration comparisons.

Name	Туре	Description
Lower percentage for variability (%)	Numeric	The default value of 25% may be overruled.
Upper percentage for variability (%)	Numeric	The default value of 75% may be overruled.
Store simulated individual day exposures	Boolean	Store the simulated individual day exposures. If unchecked, no additional output will be generated. If checked, the output will contain an additional section with the simulated individual day exposures.

Calculation of biological matrix concentration comparisons

Biological matrix concentration comparisons calculations comprise two parts. The first part is to compute estimates of the human monitoring concentrations based on the human monitoring data. The second part is to relate the human monitoring concentrations to modelled concentrations from exposure assessments.

• Biological matrix concentration comparisons calculation

Inputs used: HBM analysis Internal exposures

Settings used

• Calculation Settings

3.4.13 Exposure biomarker conversions

Occasionally, the biomarker of interest (substance) is not measured. Exposure biomarker conversions specify how measured biomarkers are converted to the relevant biomarkers (substances). This type of conversion is within biological matrices.

This module has as primary entities: Substances

Output of this module is used by: HBM analysis

Exposure biomarker conversions from data

In table *Exposure biomarker conversions*, conversion factors are specified to convert measured biomarkers to the biomarkers of interest. Conversion factors may be dependent on individual properties like age and gender. Specify in table *Exposure biomarker conversion subgroups* age and/or *gender* specific conversion factors.

For age, specify the lower bound of the age interval (in years) of the exposure biomarker conversion subgroup. Individuals belong to a subgroup when the age of the individual is equal or greater than the specified lower bound and smaller than the specified lower age of the next subgroup.

Check option Use exposure biomarker conversion factors subgroup to use age and/or gender specific conversion factors in your assessment.

Column *VariabilityDistributionType* in table *Exposure biomarker conversions* is used to specify the variability distribution to sample from. When the column is left empty, the conversion factor itself is used. Currently, four distributions are available.

See also uniform, lognormal, beta and inverse uniform distribution.

Exposure biomarker conversions data formats

Describes the exposure biomarker conversions (i.c. conversions within a biological matrix).

Download empty dataset template: Zipped CSV Excel

Exposure biomarker conversions

Exposure biomarker conversions main table. Biomarker conversion is within a biological matrix.

Name	Туре	Description	Aliases	Required
idExposure- Biomarker- Conversion	AlphaNumeric (50)	Id of the exposure biomarker conversion	idExposure- Biomarker- Conversion, idExposure- Biomarker- Factor, idEBCFactor	Yes
idSubstance- From	AlphaNumeric (50)	Identification code of substance of the 'from' substance. That is, the code of the (measured) substance for which concentrations need to be translated.	idSubstance- From, SubstanceId- From, SubstanceCode- From, SubstanceFrom	Yes
Biological matrix	BiologicalMatrix	The biological matrix (e.g., blood, urine) for which this conversion rule is defined.	Biological- Matrix, MatrixSource, SourceMatrix	Yes
Expression- TypeFrom	ExpressionType	The expression type (or adjustment method) of the 'from'-part. This field specifies how the dose unit (source) is expressed, e.g., for blood lipids for fat soluble biomarkers ('mg/g lipids') or the dilution level of the urine ('mg/g creatinine').	ExpressionType- From, Adjustment- MethodFrom	No
UnitFrom	DoseUnit	The unit of the 'from'-part of the conversion rule.	UnitSource- From, SourceUnitFrom	Yes
idSubstanceTo	AlphaNumeric (50)	Identification code of substance of the 'to' substance.	idSubstanceTo, SubstanceIdTo, SubstanceCode- To, SubstanceTo	Yes
Expression- TypeTo	ExpressionType	TODO: to be removed. The expression type or adjustment method of 'to'-part of the conversion rule. This field specifies how the dose unit (target) is adjusted, e.g., for blood lipids for fat soluble biomarkers ('mg/g lipids') or the dilution level of the urine ('mg/g creatinine').	ExpressionType- To, Adjustment- MethodTo	No
UnitTo	DoseUnit	TODO: to be removed. The unit of the 'to'-part of the conversion rule.	UnitSourceTo, SourceUnitTo	Yes
Conversion- Factor	Numeric	The conversion factor value. If the conversion is provided as a distribution, then this factor is the nominal value of the distribution.	Conversion- Factor, Factor	Yes
Variability- Distribution- Type	Biomarker- Conversion- Distribution	Distribution type (Uniform, LogNormal or Beta). If not specified, the conversion is assumed to be a constant	Variability- Distribution- Type, Variability-	No
4. Exposure m	odules	factor.	Distribution, Distribution- Type, Distribution	4
Variability-	Numeric	The upper value of the	Variability-	No

Table 3.263: Table definition for Exposure biomarker conversions.

Accepted table names: ExposureBiomarkerConversions, ExposureBiomarkerConversion, BiomarkerConversions, BiomarkerConversion.

Exposure biomarker conversion subgroups

Exposure biomarker conversion subgroups.

Name	Туре	Description	Aliases	Required
idExposure- Biomarker- Conversion	AlphaNumeric (50)	Id of the exposure biomarker conversion	idExposure- Biomarker- Conversion, idExposure- Biomarker- Factor, idEBCFactor	Yes
Conversion- Factor	Numeric	Conversion factor value	Conversion- Factor, Factor	Yes
AgeLower	Numeric (50)	Specifies the lower bound of the age interval (in years) of the exposure biomarker conversion subgroup. Individuals belong to a subgroup when the age of the individual is equal or greater than the specified lower bound and smaller than the specified lower age of the next subgroup.	AgeLower, LowerAge	No
Gender	GenderType	The gender of the exposure biomarker conversion subgroup.	Gender, Sex	No
Variability- Upper	Numeric	The upper value of the distribution. If the distribution is uniform, then it is the upper bound of the uniform distribution. If the distribution is log-normal, then the upper value is assumed to correspond with the p95 percentile of the distribution. When a distribution is assumed, this value should be greater than the conversion factor.	Variability- Upper, Upper	No

Table 3.264: Table definition for Exposure biomarker conversion subgroups.

Accepted table names: ExposureBiomarkerConversionSGs, BiomarkerConversionSGs, ExposureBiomarkerSGs, EBCFactorSubGroups.

Exposure biomarker conversions

Calculation settings

Table 3.265: Calculation settings for module Exposure biomarker conver-

sions.

Name	Туре	Description
Use exposure biomarker conversion factors subgroup	Boolean	Exposure biomarker conversion factors are dependent on subgroups (e.g. based on age or gender, see dataformats ExposuerBiomarkerConversionsSgs).

Exposure biomarker conversions as data

Exposure biomarkers conversions are provided as data in the form of a conversion factor for a substances (from) and substance (to).

- Exposure biomarker conversions data formats
- Exposure biomarker conversions from data

Settings used

• Calculation Settings

3.4.14 HBM analysis

Human biomonitoring concentrations are substance concentration estimates for a biological matrix (e.g., urine or blood) derived from data obtained from human monitoring studies.

This module has as primary entities: Populations Substances

Output of this module is used by: *Exposure mixtures Biological matrix concentration comparisons Risks Environmental burden of disease*

HBM analysis calculation

Human monitoring analysis computes substance concentration estimates for a biological matrix (e.g., urine or blood) based on *human monitoring data*. These estimates are specified at the level of long term average concentrations for individuals in case of *chronic assessments*, or concentrations for individual-days in case of *acute assessments*. The concentrations are computed independently for each substance and biological matrix.

The main steps, and also performed in this order, for computing human monitoring concentration estimates are:

- 1. Imputation of censored values.
- 2. Imputation of missing values.
- 3. Standardise blood for lipid content (only available when data allows).
- 4. Standardise/normalise urine for creatinine or specific gravity (only available when data allows).
- 5. Filter for complete cases.
- 6. Combine urine from different sampling methods.
- 7. Apply exposure biomarker conversion of substance concentrations (within a biological matrix).
- 8. Apply *kinetic conversion* of substance concentrations from other biological matrices (only available when data allows).
- 9. Calculation of individual concentrations (chronic) or individual day concentrations (acute).

Imputation of censored values

Similar to *concentrations measurements in food*, human monitoring measurements contain measurements below the limit of reporting and similar to *concentrations modelling in foods*, human monitoring analysis addresses these censored values and replaces them with imputed concentration values, see also *imputation methods*. Two approaches are available:

- 1. Impute using a non-detects handling method.
- 2. Impute using a draw from the left tail of the *censored lognormal distribution*. See also *concentration models* and *concentration model types*.

The available non-detects handling methods for deterministic imputation are:

- Replace censored values by zero.
- Replace censored values by a factor * *LOR*, the factor is set between zero and one.
- Replace non-detects by a factor * *LOD* and non-quantifications by *LOD* + factor * (*LOQ* LOD), the factor is set between zero and one.
- Replace non-detects by zero and non-quantifications by factor * *LOQ*, the factor is set between zero and one.

For option 3, factor = 0, non-detects are replaced by zero, non-quantifications are replaced by LOD; for factor = 1, non-detects are replaced by LOD, non-quantifications are replaced by LOQ. See also PARC harmonised HBM data format: *Sample concentrations/measurements*.

Note, when LOD is not available then it is assumed to be 0. When LOQ is not available then it is assumed to be LOD (or zero if LOD is also not available).

For imputation based on the censored lognormal distribution, non-detect and non-quantification information is used. Non-detects are sampled from the entire left tail, e.g. the area below LOD. Non-quantifications are sampled from the intermediate segment, e.g. the area below LOQ and above LOD.

Note that for the option based on the censored lognormal distribution also a non-detects handling method has to be specified. Occasionally, fitting the censored lognormal model fails and the deterministic imputation method is used as fallback.

The *censored lognormal distribution*, with parameters μ and σ , is fitted using a likelihood function which is defined by contributions of the positive measurements (lognormal probability density function) and by contributions of the measurements below LOD and measurements above LOD and below LOQ separately, e.g. the (cumulative) probabilities for LODs and LOQs.

In Figure 3.68, the *human monitoring concentration data* in urine are imputed with a factor * LOR and the summary statistics are visualized.



Concentration (µg/kg bw/day)

Figure 3.68: Boxplots for imputed concentration data for five pesticides. Lower whiskers indicate the p5 and p10 percentiles, upper whiskers the p90 and p95. The edges of the box indicate the p25 and p75 percentiles with the median in the centre of the box.

Imputation of missing values

For missing concentration measurements, *three imputation methods* are available, see also *imputation methods*. The third opion ignores imputation and all missing values remain in the data set.

- 1. Replace missing values by zero. This is conditional on the specified minimum percentage of non-missing values.
- 2. For each substance and combination of biological matrix and sampling type, replace missing values by a random draw from the non-missing concentration values (samples). This is conditional on the specified minimum percentage of non-missing values.
- 3. Do not impute missing values.

Imputation of missing values by zeros or a random draw from the data is conditional on the specified percentage of non-missing values. When the percentage of non-missing values for a specific substance and combination of biological matrix and sampling type in the data is smaller than the specified percentage, imputation is ignored. This is to prevent that imputation takes place using a set of imputation values that is not representative or unrealistically small (e.g. one or a few values). Note that for the second imputation method, more refined methods could be used. E.g., when for a given day multiple samples are available and one is missing, then this sample might be neglected in the computation of an average exposure. Also, when samples have been taken at different time-slots, impute the missing records using samples from the same time-slot.

Note that imputation of missing values applies only to substances within the assessment scope, not to those used indirectly for standardisation and normalisation, such as creatinine, total cholesterol, and triglycerides.

Standardise blood for lipid content

Lipid-soluble substances measured in blood data are typically standardised by total lipid content, see also *standardi-sation methods*. Three methods are in available, namely:

- 1. Standardisation based on gravimatic analysis.
- 2. Standardisation based on enzymatic summation analysis.
- 3. Standardisation based on Bernert et al. (2007), total lipids (mg/dL) = 2.27 * total cholesterol + triglycerides + 62.3 mg/dL.

The standardisation is only applied to lipid-soluble substances, see *Substances data formats*. After standardisation, the amount of substance is expressed per g lipid e.g. $\mu g/g$ lipid. Note that substance concentrations in blood samples with unmeasured lipid concentrations are set to missing after specifying option blood standardisation.

This option is only available to the user when the data contains substances that are soluble in lipid and when total lipid content (or cholesterol and triglycerides) is measured.

Standardise/normalise urine for creatinine or specific gravity

Four methods are available for correcting spot urine measurements for creatinine of specific gravity, see also *stan- dardisation methods*.

- 1. Normalisation based on *specific gravity*.
- 2. Standardisation based on creatinine content.
- 3. Normalisation based on specific gravity derived from creatinine content, adults 18 68 years.
- 4. Normalisation based on *specific gravity derived from nonlinear modelling of creatinine content, children 6 14 years.*
- 5. Normalisation based on *specific gravity derived from nonlinear modelling of creatinine content, age and gender dependent, children 6 14 years.*

Urine's specific gravity is determined by the concentration of excreted molecules in the urine. In adult humans, normal specific gravity values range from 1.010 to 1.030. The specific gravity normalisation used here (1) is equal to (1.024 - 1)/(specific gravity - 1). The specific gravity value should be available in the HBM data, otherwise urine sample concentrations are set to missing.

After standardisation for creatinine content, the amount of substance is expressed per g creatinine e.g. $\mu g/g$ creatinine. Note that substance concentrations in urine samples where the creatinine content is not measured are set to missing values after specifying option urine standardisation.

Options 3 (Carrieri et al. (2000)) and 4 and 5 (Busgang et al. (2023)) are for specific subgroups in the population, e.g. adults (18 - 68 years) and children (6 - 14 years), respectively. For model 2 of Busgang et al. (2023), age and gender should be defined in the data, otherwise the sample concentration is set to missing.

The normalisation and standardisation methods are only available when the data contains creatinine or specific gravity values.

Combine urine from different sampling methods

Different urine sampling methods (morning, spot, or 24-hour) are considered a single biological matrix for further analysis in MCRA. If the selected human biomonitoring data contains concentrations from different urine sampling methods, the concentration values are combined by taking the average of the concentration values from the various urine sampling methods.

Filter for complete cases

After imputation of non-detects and missing values, standardisation and normalisation some individual day concentration records still contain missing values. The incomplete cases are removed from the dataset before the analysis continues.

Occasionally, removing all records with missing values results in empty datasets. Then a warning will be thrown 'All HBM individual day records were removed for having non-imputed missing substance concentrations'. To circumvent this warning, inspect your data and remove substances with too many missing values, lower the minimum percentage of non-missing values (*Impute from data*), or *Impute with zero*.

Filtering for complete cases is always applied before conversion. In general, filtering has more impact when many substances are in scope, resulting in smaller subsets of complete cases. It is good practice to set the scope to the relevant substances. In the interface of the primary entity *substances*, select the relevant substances for the assessment.

Apply exposure biomarker conversion of substance concentrations

Occasionally, the biomarkers of interest are not measured. *Exposure biomarker conversion* is used to convert measured biomarkers to the biomarker of interest and is applied within a *biological matrix*. Biomarker conversion is performed after imputation of censored and missing values and after normalisation/standardisation of urine/blood samples. For conversion factors that are dependent on age and/or *gender*, check option *Use exposure biomarker conversion factors subgroup* in the interface of the exposure biomarker conversion module.

Apply kinetic conversion of substance concentrations from other biological matrices

Kinetic conversion of substances from other biological matrices is used when the number of substances in the matrix of interest is limited or for reverse dosimetry when exposures at external level are required. As an example of an internal-to-internal conversion, consider a scenario where five substances are measured in *urine (spot)* samples from individuals. Additionally, blood (serum) concentrations are analysed for the same individuals, resulting in another set of concentrations for, let's say, five different substances. The five substances measured in blood are then converted to urine matrix concentrations by checking the Apply kinetic conversions option. Then, the analysis continues with ten substances in urine (spot). Selecting specific substances for conversion is possible through multiplication factors defined in the *kinetic conversion factors data* of the Kinetic models module.

Enable the *Convert to single exposure surface (biological matrix or external route) matrices* option to compute exposures at a single target biological matrix or to compute exposures at the external level. Specify *the target level* (Internal or External) and *biological matrix* accordingly. *Kinetic conversion of substance concentrations* is performed after imputation of censored and missing values, and after normalisation/standardisation of urine/blood samples.

Note that kinetic conversion is applied only to substances that are not measured in the target matrix. For instance, kinetic conversion cannot be used for imputing missing values of substances that have one or more missing values.

For conversion factors that are dependent on age and/or *gender*, check option *Use kinetic conversion factors subgroup* in the interface of the kinetic models module.

Calculation of acute human monitoring concentrations

For *acute* assessments, the monitoring concentrations are computed for each substance and biological matrix as average individual-day concentrations. The computation is done after imputation of censored and missing values, eventually followed by a conversion of biomarkers from other biological matrices. For a given substance and biological matrix, the acute individual-day concentration c_{ij} for individual *i* on day *j* is:

$$c_{ij} = \frac{\sum_{k=1}^{n_{\rm samples}} c_{ijk}}{n_{\rm samples}}$$

where n_{samples} is the number of samples available for individual i on day j, and c_{ijk} the concentration of the k-th sample of the individual day j.

After urine normalisation for specific gravity:

$$c_{ij}' = c_{ij} \cdot sg$$

where sg denotes the specific gravity correction factor for that individual day.

After standardisation for blood lipid content:

$$c_{ij}' = \frac{c_{ij}}{c_{ij} \text{ lipid}}$$

where c'_{ij} denotes the lipid concentration per g lipid. The standardisation is only performed for lipid soluble substances. After standardisation the concentration of the substance is expressed as substance amount, with a user specified unit, per g lipid.

The standardisation for creatinine is similar to the above equation replacing lipid by creatinine.

Calculation of chronic human monitoring concentrations

For *chronic* assessments, the monitoring concentrations are computed as the average monitoring concentrations of multiple individual-days for each substance and biological matrix. The computation is done after imputation of censored and missing values, eventually followed by a conversion of biomarkers from other biological matrices. The chronic concentration c_i for individual *i* is computed as:

$$c_i = \frac{\sum_{j=1}^{n_{\rm days}} c_{ij}}{n_{\rm days}},$$

where n_{days} is the number of days that individual *i* was monitored (after removing missing individual days), and c_{ij} denotes the average monitoring concentration of individual *i* on day *j*.

Standardisation and normalisation of blood and urine samples, respectively, are similar to the expressions for the calculation of individual day concentrations (acute).

For co-exposure of substances, see maximum cumulative ratio (MCR) and the exposure mixtures module.

HBM analysis settings

Calculation settings

Name	Туре	Description
Seed for pseudo-random number generator	Numeric	A value of 0 will use a pseudo-random seed in each run, a value 0 will provide the same results in a repeated run.
Exposure type	ExposureType	The type of exposure considered in the assessment; acute (shor term) or chronic (long-term).
Multiple substances analysis	Boolean	Specifies whether the assessment involves multiple substances.

Table 3.266: Calculation settings for module HBM analysis.

continues on next pa

Name Type Description	
Compute cumulative exposures Boolean Specifies whether the assessment involves multiple	ple substances a
results should be cumulated over all substances.	
Report consumptions and Boolean Specifies whether body weights should be ignore	d and
exposures per individual consumptions and exposures should be expressed of per kg body weight.	a per individua
weight weight	per kg body
Sampling method AlphaNumeric The sampling method that should be included in	the action.
Censored values handling <i>NonDetectsHandlingMethod</i> Method for dealing with censored value samples	s in human
method (also used as fallback monitoring data. Note that this method is also u	sed as a fallbac
for censored lognormal when fitting a censored lognormal model to the	concentration d
approach) fails.	
Fraction for censored value Numeric Factor used for replacing the censored value.	
replacement	1
Imputation method for non <i>NonDetectImputationMethod</i> Imputation method for non detect values: replace	e nondetects
detect values based on by I*LOD/LOQ) or from left tail cens	ored lognormal
Missing value imputation Missing ValueImputation Method Imputation method for missing values: 1) By ze	ro 2) Impute
method from data, 3) No missing value imputation	10, 2) impute
Apply kinetic conversions Boolean Convert substance concentrations from other bio	ological matrice
using kinetic conversion models. The substances	s for conversior
are designated within the kinetic models module	e. Substance
conversion proves valuable when a biomarker w	as not directly
measured for a matrix of interest.	
Convert to single exposure Boolean Convert all substance concentrations from other	biological
surface (biological matrix or matrices to the same target biological matrix. T	his conversion i
external route) applied when the number of substances measured	d on the target
biological matrix is influed. Substances measure matrices can be converted using kinetic convers	ion models
Target level Target evel Type Select whether the assessment is at the level of e	external dose
internal (target) concentration or internal (system	nic) dose. For
aggregate assessment, combining dietary and no	ndietary
exposures, the target level is always internal con	centration or
internal (systemic) dose. When only dietary exp	osures are
available, the target level is optional and may be	an external
(administered) dose, an internal (tissue) concent	tration or an
internal (absorbed) dose.	
Specify the minimum Percentage of non-missin	g values require
percentage of non-missing for imputation. No imputation is done when the values (q')	percentage of
values (%) non-missing values in the data is smaller than the	e specified
Standardise blood Boolean Standardise blood concentrations for lipid-solub	le substances
concentrations for lipid-soluble Standardise by total lipid measured via gravime	tric analysis, 2)
substances Standardise by total lipid measured via enzymat	ic summation,
Standardise by derived total lipid content of	
Triglycerides/Cholesterol (Bernert et al. 2007).	
Specify the standardisation <i>StandardiseBloodMethod</i> Specify the standardisation method of blood con	ncentrations for
method of blood lipid-soluble substances.	
concentrations for lipid-soluble	
substances	aluble autota
substances from linid	oluble substanc
standardisation	
Select substances to exclude AlphaNumeric The selected (lipid-soluble) substances will be e	xcluded from li
from lipid standardisation standardisation.	

Table 3.	266 – cor	ntinued f	from p	orevious	page
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Name	Туре	Description
Normalise or standardise urine concentrations for specific gravity or creatinine	Boolean	Normalise or standardise urine concentrations for specific gravit or creatinine: 1) Normalise by specific gravity, 2) Standardise by creatinine concentration.
Specify the normalisation/standardisation method of urine concentrations for specific gravity or creatinine	StandardiseUrineMethod	Specify the normalisation/standardisation method of urine concentrations for specific gravity or creatinine.
Subset selection: exclude substances from urine normalisation/standardisation	Boolean	Select this option to exclude one or more substances from normalization for specific gravity or creatinine standardisation.
Select the substances to exclude from urine normalisation/standardisation	AlphaNumeric	The selected substances will be excluded from urine normalization/standardisation.
Substance weighting in mixtures	ExposureApproachType	Risk based: exposures in equivalents of the reference substance standardised: standardised exposures per substance have varian 1; or unweighted exposures: RPFs are equal to 1.
Perform MCR analysis	Boolean	Perform a Maximum Cumulative Ratio (MCR) analysis to determine co-exposure between substances.
Display ratio total exposure/ maximum (in MCR plot)	Numeric	For MCR plot: specify ratio total exposure/ maximum for individual(day) exposures.
Show tail percentiles (MCR plot) for:	Numeric	Give specific percentiles of exposure distribution (%), e.g. 97.5 99 (space separated).
Set minimum percentage contribution per substance to the tail exposure (MCR plot)	Numeric	Set minimum percentage contribution per substance to the tail exposure.
Cutoff MCR	Numeric	For selection of individual(day) exposures with maximum cumulative ratio (MCR = total exposure/maximum) above the cutoff.
Cutoff percentage in population ranked on total exposure	Numeric	For selection of individual(day) exposures above the cutoff percentage in the set of individual(day)s ranked on total exposu
Apply exposure biomarker conversions	Boolean	Use this option to translate HBM concentrations derived from measured substances (biomarkers) to concentrations of other substances. This can be usefull when the measured substance is combination of multiple substances, e.g., to translate measured total arsenic (t-As) to toxicologically relevant arsenic (TRA). T option can also be used to translate between different expression types (e.g., from measured urine concentration to urine concentrations standardized for specific gravity), but not for translation between different biological matrices.
Target biological matrix	BiologicalMatrix	The target biological matrix (internal compartment) for which exposures are computed.
Specific gravity conversion factor	Numeric	A specific gravity adjustment is applied by multiplying a creatinine adjusted concentration with a factor (default 1.48 for adults 18 - 68 year).

Table 3.266 - continued from previous page

Output settings

Name	Туре	Description
Exclude privacy sensitive data from outputs	Boolean	Use this setting to not report the parts of the results (i.e., figure tables, or sections) that are marked as (potentially) privacy sensitive.
Lower percentage for variability (%)	Numeric	The default value of 25% may be overruled.
Upper percentage for variability (%)	Numeric	The default value of 75% may be overruled.
Percentage for upper tail	Numeric	Gives detailed output for this upper percentage of the exposure distribution.
Store simulated individual day exposures	Boolean	Store the simulated individual day exposures. If unchecked, no additional output will be generated. If checked, the output will contain an additional section with the simulated individual day exposures.

Table 3.267: Output settings for module HBM analysis.

Uncertainty settings

Table 3 268	Uncertainty	settings	for mod	ule	HBM	analysi	S
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Name	Туре	Description
Lower uncertainty limit (%)	Numeric	Percentage lower bound, e.g. 2.5%.
Upper uncertainty limit (%)	Numeric	Percentage upper bound, e.g. 97.5%.
Monte Carlo iterations per uncertainty run	Numeric	Specifies the number of Monte Carlo iterations in each uncertainty cycle (default 10,000).
Resample HBM individuals	Boolean	HBM individual data are resampled from the original database using the bootstrap methodology (Efron 1979, Efron & Tibshir 1993).

Calculation of hbm analysis

Human biomonitoring concentration estimates are computed from data collected in human monitoring studies. These concentration estimates are computed per substance for a selected (human) biological matrix. Modelling includes imputation of missing values and non-detects, as well as correction of concentrations for, e.g., specific gravity. Occasionally, the number of substances measured in the biological matrix of interest is limited or too low to perform a risk assessment or mixture analysis. In such cases, matrix conversion can be applied: substance concentrations measured in other biological matrices are translated to the matrix of interest using a kinetic conversion factor. Cumulative risks of multiple substances are calculated either by using RPFs weighted exposure or by using the sum of risk ratios method using hazard characterisations.

• HBM analysis calculation

Inputs used: *HBM concentrations Active substances Relative potency factors Kinetic conversion factors Exposure biomarker conversions*

Settings used

Calculation Settings

3.4.15 HBM concentrations

Human (bio)monitoring data quantify substance concentrations found in humans collected in human monitoring surveys.

This module has as primary entities: Populations Substances

Output of this module is used by: HBM analysis

HBM concentrations from data

Human monitoring data characterise human exposure to substances, their metabolites in body fluids or tissues. In Figure 3.69, the positive (> zero) human monitoring concentrations for a number of pesticides in urine (morning) are visualised through summary statistics.



Figure 3.69: Boxplots for positive/quantified HBM substance concentration measurements of 13 pesticides. Lower whiskers indicate the p5 and p10 percentiles, upper whiskers the p90 and p95. The edges of the box indicate the p25 and p75 percentiles with the median in the centre of the box. LORs are displayed by a vertical red bar and outliers outside range (Q1 - $3 \times IQR$, Q3 + $3 \times IQR$).

Find in Figure 3.70, boxplots for all human monitoring concentrations of 13 pesticides in urine (morning).

Note that all statistics are weighted using sampling weights.

Selection on sampling method

The dropdown menu displays the available sampling methods as defined in the data source. Select one or more sampling methods here to include them in the action.



Figure 3.70: Boxplots for all of HBM substance concentration measurements of 13 pesticides. Lower whiskers indicate the p5 and p10 percentiles, upper whiskers the p90 and p95. The edges of the box indicate the p25 and p75 percentiles with the median in the centre of the box. LORs are displayed by a vertical red bar and outliers outside range (Q1 - 3 * IQR, Q3 + 3 * IQR).

Filter on repeated measurements

This option is only available when the survey contains two or more measurements or time points. It allows for the restriction of the action to one measurement to prevent the combination of unrelated samples. If there is only one measurement in the survey or if this option is not selected, all measurements are included in the action.

Excluded substances from sampling method

Use this option to remove all samples of substances that are of no interest or have incomplete samples. This option can be combined with the "Use complete analysed samples" option. When "Use complete analysed samples" is selected, the resulting dataset might become very small due to some substances not being analysed. Excluding these non-analysed substances will result in a larger dataset.

Use complete analysed samples

Select this option to remove samples from individuals for whom one or more of the selected substances have not been analysed.

Use sampling weights

If checked, individual sampling weights are used. If unchecked, the individual sampling weights are not used in the calculations.

HBM concentrations data formats

Human biomonitoring data are analytical measurements of chemical substances or markers of health effects in body fluids or tissues. This data can be used for assessing (combined) exposure to chemicals and the risks associated with this exposure. MCRA supports upload of this data in two formats: a relational table structure that matches the internal representation of MCRA and the PARC harmonized HBM data format.

Relational human monitoring concentration data format

The relational human monitoring concentrations data format is the format that is used internally in MCRA. The data format consists of a number of tables for specification of the study, the individuals of the study (and their specific characteristics), and the human (bio)monitoring samples and sample analyses.

Download empty dataset template: Zipped CSV Excel

Human biomonitoring surveys

Contains the definitions of the human (bio)monitoring surveys/studies.

Name	Туре	Description	Aliases	Required
idSurvey	AlphaNumeric (50)	Unique identification code of the survey.	idSurvey, idStudy	Yes
Name	AlphaNumeric (100)	Name of the study/survey.	Name	No
Description	AlphaNumeric (200)	Description of the study/survey.	Description	No
Location	AlphaNumeric (50)	The location or country where survey is held. It is recommended to use ISO Alpha-2 country codes.	Location, Country	No
BodyWeight- Unit	AlphaNumeric (50)	The unit of bodyweight of the individuals of the survey: kg (default) or g.	BodyWeight- Unit, UnitBody- Weight, WeightIn	No
AgeUnit	AlphaNumeric (50)	The unit of age, i.e., year or month.	UnitAge, agein, AgeUnit	No
StartDate	DateTime	The starting date of the survey.	StartDate	No
EndDate	DateTime	The end date of the survey.	EndDate	No
NumberOf- SurveyDays	Integer	The number of days each individual participated in the survey.	NumberOf- SurveyDays, NDaysInSurvey	Yes
idPopulation	AlphaNumeric (50)	Unique identification code of the population.	IdPopulation, PopulationId	No
Lipid- Concentration- Unit	ConcentrationUnit	The unit of the lipid concentration (defaults mg/dL).	Lipid- Concentration- Unit, LipidUnit	No
Triglyc- Concentration- Unit	<i>ConcentrationUnit</i>	The unit of the triglycerides concentration (defaults mg/dL).	Triglyc- Concentration- Unit, Triglycerides- Concentration- Unit, Triglycerides- Unit	No
Cholest- Concentration- Unit	<i>ConcentrationUnit</i>	The unit of the cholesterol concentration (defaults mg/dL).	Cholest- Concentration- Unit, Cholesterol- Concentration- Unit, CholesterolUnit	No
Creat- Concentration- Unit	ConcentrationUnit	The unit of the creatinine concentration (defaults mg/dL).	Creat- Concentration- Unit, Creatinine- Concentration- Unit, CreatinineUnit	No

Table 3.269:	Table definition	for Human	biomonitoring	surveys.
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 $\label{eq:constraint} Accepted \ table \ names: \ HumanMonitoringSurveys, \ HumanMonitoringSurveys, \ HbmSurveys.$

Individuals

The individuals of a survey are recorded in the Individuals table. Add additional properties like Region, Breastfeeding to further describe an individual. In table IndividualProperties, each property in the Individuals table is described (recommended way). Note that only those properties that are available in the Individuals table are used in module Populations, table Populations or PopulationIndividualPropertyValues to subset the individuals. This is only relevant when the UseData option in the population module is used.

Name	Туре	Description	Aliases	Required
idIndividual	AlphaNumeric (50)	Unique identification code of the individual.	idIndividual, IndividualId, Individual, Id	Yes
idFoodSurvey	AlphaNumeric (50)	The identification code / short name of survey.	idSurvey, idFoodSurvey, Survey, FoodSurvey, SurveyId, FoodSurveyId, SurveyCode	Yes
BodyWeight	Numeric	The body weight of the individual.	BodyWeight, Weight	No
Sampling- Weight	Numeric	The sampling weight for an individual (default = 1).	SamplingWeight	No
NumberOf- SurveyDays	Integer	The number of days the individual participated in the survey.	NumberOf- SurveyDays, NumberOfDays- InSurvey, DaysInSurvey, NDaysInSurvey	No
Name	AlphaNumeric (100)	Name or label of the individual.	Name	No
Description	AlphaNumeric (200)	Additional description of the individual.	Description	No
Individual properties		Other individual properties can be added like the fields Age, Gender, Region etc. These properties are automatically parsed as co-factors or co-variables.		No

Table 3.270:	Table	definition	for	Individuals.
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Accepted table names: Individuals, SurveyIndividuals, ConsumptionSurveyIndividuals, FoodConsumptionSurveyIndividuals.

Individual properties

This table is used to describe the properties used in the Populations or PopulationIndividualPropertyValues table characterising the population (table Populations) and/or the properties used in the Individuals table characterising an individual. Properties like Age, Gender, Region are describing an individual (PropertyLevel = Individual). Properties like Period (for populations) or Month (sampling date for an individual day) are describing an individual day (PropertyLevel = IndividualDay).

Name	Туре	Description	Aliases	Required
idIndividual- Property	AlphaNumeric (50)	The code of the property.	idIndividual- Property, Individual- PropertyId, Individual- Property	Yes
Name	AlphaNumeric (100)	The name of the property.	Name	No
PropertyLevel	<i>PropertyLevelType</i>	The level of the property. This type follows a controlled terminology, with possible values: Individual or IndividualDay.	PropertyLevel, LevelProperty	No
Description	AlphaNumeric (200)	Description of the property.	Description	No
Туре	IndividualProperty- Type	This field specifies the type of the values of this individual property. This type follows a controlled terminology, with possible values: Boolean, Categorical (default), Numeric, Nonnegative, Integer, NonnegativeInteger, Month, Datetime, Gender.	Туре	No

Table 3.271: Table definition	for Individual	properties.
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Accepted table names: IndividualProperties, IndividualProperty.

Individual property values

Not recommended. This table describes individual property values. Property values are describing an individual for properties like e.g. Region, Breastfeeding. The recommended way is to add these columns as additional columns in the Individuals table. In table IndividualProperties, each property in the IndividualPropertyValues table is described.

Name	Туре	Description	Aliases	Required
idIndividual	AlphaNumeric (50)	The identification number of the Individual.	Id	Yes
PropertyName	AlphaNumeric (50)	The name of the property.	Name	Yes
TextValue	AlphaNumeric (50)	The value of the property as text value.		No
DoubleValue	Numeric	The value of the property as number.		No

Table 3.272: Table definition for Individual property values.

Accepted table names: IndividualPropertyValues, IndividualPropertyValue.

Analytical methods

The analytical methods used for analysing the samples are recorded in the analytical methods table. Each analytical method should have a unique identification code (idAnalyticalMethod). The description field may be used for a more detailed description of the analytical method. The records of this table should be linked to one or more analytical method substance properties table, which record the substances that are measured by this method (and their limits of reporting).

Name	Туре	Description	Aliases	Required
idAnalytical- Method	AlphaNumeric (50)	The code for the method of analysis.	idAnalytical- Method, Analytical- MethodId, Analytical- MethodName, Id	Yes
Name	AlphaNumeric (100)	Name of the analytical method.	Name	No
Description	AlphaNumeric (255)	Additional description of method of analysis.	Description	No

Table 3.273:	Table definition	for Analytical	methods.
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Accepted table names: AnalyticalMethod, AnalyticalMethods.

Analytical method properties for substances

This table describes the substances analysed by an analytical method. For each substance analysed by an analytical method a record should be included that describes the unit of measurement and the reporting limits (LOQ/LOD).

Table 3.274:	Table	definition	for	Analytical	method	properties	for	sub-
stances.								

Name	Туре	Description	Aliases	Required
idAnalytical- Method	AlphaNumeric (50)	The code of method of analysis.	idAnalytical- Method, Analytical- MethodName, Analytical- MethodId	Yes
idSubstance	AlphaNumeric (50)	The substance code.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
LOD	Numeric	The limit of detection (LOD) is the lowest concentration of an substance in a sample that can be consistently detected.	LOD	No
LOQ	Numeric	The limit of quantification (LOQ) is the lowest concentration of a substance that can be quantified. The LOQ should be larger than the LOD.	LOQ, LOR	No
Concentration- Unit	ConcentrationUnit	The unit used for reporting the LOD, LOQ, and the substance concentrations. When not specified, then a default unit of mg/kg is assumed.	Concentration- Unit, Units, Unit	No

 $\label{eq:accepted} Accepted\ table\ names:\ AnalyticalMethodSubstances,\ AnalyticalMethodSubstance,\ AnalyticalMethodCompounds,\ AnalyticalMethodCompound.$

Human biomonitoring samples

Describes the samples taken during the study. Each sample has a unique identifier/code.

Name	Туре	Description	Aliases	Required
idSample	AlphaNumeric (50)	Unique identification code of the monitoring sample.	idSample, Sample	Yes
idIndividual	AlphaNumeric (50)	Unique identification code of the individual.	idIndividual, IndividualId, Individual, Id	Yes
DateSampling	DateTime	Date of sampling.	DateSampling, DateOf- Sampling, SamplingDate	No
DayOfSurvey	AlphaNumeric (50)	Identification code of the day of measurement.	Day, idDay, DayId, DayOfSurvey	Yes
TimeOf- Sampling	AlphaNumeric (50)	Identification code of the time of sampling.	TimeOf- Sampling, SamplingTime, TimeSampling	No
SampleType	AlphaNumeric (50)	Type of sample (e.g., pooled, 24h urine, spot urine, serum from blood, etc.).	SampleType, SamplingType	No
Compartment	AlphaNumeric (50)	If applicable, the measured compartment of the human body (e.g., blood, urine). When specified, the measurements are considered at the level of internal doses.	Compartment	No
ExposureRoute	ExposureRoute	If applicable, the measured exposure route, e.g., dermal (in case of skin wipes). When specified, the measurements are considered at the level of external doses.	ExposureRoute	No
SpecificGravity	Numeric	Specific gravity of the measured person for this particular sample.	SpecificGrafity, SpecificGravity	No
SpecificGravity- Correction- Factor	Numeric	Correction factor for the concentration to account for the specific gravity of the measured person for this particular sample.	SpecificGravity- Correction- Factor	No
Name	AlphaNumeric (100)	Name of the human monitoring sample.	Name	No
Description	AlphaNumeric (200)	Additional description of the human monitoring sample.	Description	No
LipidGrav	Numeric	Lipid content based on gravimatic analysis of the measured person for this particular sample.	LipidGrav, LipidGravimatic	No
LipidEnz	Numeric	Lipid content based on enzymatric summation of the measured person for this particular sample.	LipidEnz, LipidEnzymatic	No
Triglycerides	Numeric	Triglycerides total of the measured person for this particular sample.	Triglyc, Triglycerides	No
Cholesterol	Numeric	Cholesterol total of the measured person for this	Cholest, Cholesterol	No
Exposure mo	odules Numeric	particular sample. Creatinine content of the measured person for this particular sample.	Creat, Creatinine	No
Ormatia	Numaria			NT.

Table 3.275: Table definition for Human	biomonitoring samples.
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Accepted table names: HumanMonitoringSamples, HumanMonitoringSample, HbmSamples.

Human biomonitoring sample analyses

Contains the measurements of the samples of human biomonitoring studies.

Name	Туре	Description	Aliases	Required
idSample- Analysis	AlphaNumeric (50)	Unique identification code of the sample analysis.	idSample- Analysis, SampleAnalysis	Yes
idSample	AlphaNumeric (50)	Code of the measured monitoring sample.	idSample, Sample	Yes
idAnalytical- Method	AlphaNumeric (50)	The code of method of analysis.	idAnalytical- Method, Analytical- MethodName, Analytical- MethodId	Yes
AnalysisDate	DateTime	Date of analysis.	AnalysisDate, DateAnalysis	No
Substance concentration(s)	AlphaNumeric (100)	Substance concentrations can be uploaded via the sample concentrations table or via additional columns of the sample analyses table. For the latter, one or more columns with the measured concentrations of the substances in the unit as specified by the analytical method should be included in the data table. The column name(s) should match the substances measured by the analytical methods. Empty fields for substances that should have been measured by the analytical method are considered to be censored with measurement values below LOQ or LOD.		No
Name	AlphaNumeric (100)	Name of the human monitoring sample analysis.	Name	No
Description	AlphaNumeric (200)	Additional description of the human monitoring sample analysis.	Description	No

Table 3 276.	Table	definition	for	Human	biomon	itoring	sample	analy	vses
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 $\label{eq:accepted} Accepted \ table \ names: \ HumanMonitoringSampleAnalyses, \ HumanMonitoringSampleAnalysis, \ HbmSampleAnalyses.$

Sample concentrations

The positive concentration values for substances from analysis in the unit specified in table human monitoring sample analyses. Censored values (i.e. results 'less than LOR') are not included, their existence can be inferred from the tables AnalysisSamples and AnalyticalMethodSubstances, and the LOR itself from the analytical method.

Name	Туре	Description	Aliases	Required
idAnalysis- Sample	AlphaNumeric (50)	The identification number of the analysed sample.	idAnalysis- Sample, AnalysisSample- Id	Yes
idSubstance	AlphaNumeric (50)	The substance code.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
Concentration	Numeric	The measured concentration.	Concentration	No
ResType	ResType	The type of residue. Should be VAL (= default), LOQ, LOD or MV.	ResType	No

Accepted table names: HumanMonitoringSampleConcentrations, HumanMonitoringSampleConcentration, HumanMonitoringConcentrations, HbmSampleConcentrations.

PARC HBM data format

The PARC HBM data format is the data format proposed in the EU Partnership for the Assessment of Risks from Chemicals (PARC) project. MCRA supports upload of files in this format, of which the data is mapped to the MCRA internal data format during the upload of the files.

PARC harmonised HBM data format

1 Note

The PARC harmonised data format as well as the MCRA data format are under development and may be changed in the future.

In PARC, a harmonised data format for human biomonitoring data is being developed by VITO. Data harmonization improves the comparability of data from different HBM studies and interoperability for use with different analysis tools such as MCRA and the tool for the calculation of summary statistics of the HBM data, which can be made available via the IPCHEM portal and/or integrated into the European HBM dashboard. More information about this data format and instructions on preparing data files compliant with this format can be found at the PARC HBM data harmonization web page. This page also contains a tool for validating data files prepared in this format and an example data file that can be uploaded to and used in MCRA for testing.

Excel data files provided to MCRA in this format are mapped during the file upload process to the internal data structure/format of MCRA. For this, MCRA uses a custom mapping/conversion procedure. For a large part, this mapping is fairly straightforward. However, for some fields and entities explicit choices are made that users need to be aware of.

• Survey/study

• Each data file corresponds to one HBM survey/study in MCRA.

- Survey *StartDate* refers to the first sampling date of all repeated samples.
- Survey *EndDate* refers to the last sampling date of all repeated samples.
- When all reported *country* values of the subjects are the same, then this is used as location of the survey/study.
- When a survey consists of two or more days, in case of repeated measurements or time points, the option to *Filter on repeated measurements* becomes available.

• Subjects/individuals

- Each subject maps to MCRA individuals of the study/survey.
- A selection of the subject/individual properties is mapped to the MCRA data format. Since MCRA does not support repeated properties, a choice has to be made when mapping individual properties that are repeated individual/subject properties in the harmonised HBM data format.
- Repeated recordings for each subject's *weight* are averaged to one value: *BodyWeight* (missing values for *weight* are supported in MCRA).
 - 1) missing values for *BodyWeight* are replaced by the average of the population.
 - 2) if no weight values are available, a nominal bodyweight of 70kg is used instead.
- Repeated recordings for each subject's height are averaged to one value: Height.
- From the repeated recordings for *smoking* only the last recording is taken: *Smoking_status*.
- Samples
 - Each sample maps to an MCRA sample.
 - The *matrix* code is translated to a *biological matrix* and *sampling type*, e.g. US translates to Urine and Spot, BP to Blood and Plasma.
 - *id_timepoint* refers to the sampling time and is translated to *DayOfSurvey*.
- Analytical methods
 - MCRA analytical methods are derived/reconstructed from the concentration data sheets of the harmonised HBM data format based on the reported detection limits (LOQs/LODs) and substances and the matrix of the samples.
- Sample concentrations/measurements
 - Concentrations without value (blanks) are recorded as missing value (MV).
 - Concentrations recorded as -10 are recorded as missing value (MV).
 - When both LOD and LOQ are known, concentrations recorded as -1 are interpreted as a value below limit of detection (x < LOD).
 - When both LOD and LOQ are known, concentrations recorded as -2 are interpreted as a value below limit of quantification and above limit of detection (LOD <= x < LOQ).
 - When LOD is NaN and LOQ is known, concentrations recorded as -3 are interpreted as values below the limit of quantification (x < LOQ).
 - When LOD is known and LOQ is NaN, concentrations recorded as -1 are interpreted as values below the limit of detection (x < LOD).

HBM concentrations settings

Selection settings

Name	Туре	Description
Exposure type	ExposureType	The type of exposure considered in the assessment; acute (short term) or chronic (long-term).
Sampling method	AlphaNumeric	The sampling method that should be included in the action.
Match HBM individuals selection to population definition options	IndividualSubsetType	Match HBM individuals selection to population definition. Use population definitions (default), ignore all population definitions use a selection of properties.
Select one or more individual(day) properties to filter the population	AlphaNumeric	Select one or more individual(day) properties to filter the individuals(days) in the population.
Use sampling weights	Boolean	If checked, individual sampling weights are used. If unchecked the individual sampling weights are not in the calculations.
Use complete analysed samples	Boolean	Select this option to remove samples from individuals for which one or more of the selected substances have not been analysed.
Exclude substances from sampling method	Boolean	Select this option to exclude one or more substances from the selected sampling methods.
Substances to exclude from a sampling method	HbmSamplingMethodSub- stance	The data of the selected substances will be excluded from analy
Filter on repeated measurements	Boolean	Select this option to make a selection of the time points to be included in the action. By default, all time points are included.
Selected measurements	AlphaNumeric	The measurements that should be included in the action.
Define populations based on specified individual properties	Boolean	Define a population by selecting specific ranges/values of individual properties. E.g., the female population between ages and 45 is composed of the properties gender (female) and age (between 18 and 45).
Individual day subset	IndividualDaySubsetDefinition	Individual day subset definition.
Individuals subset definitions	IndividualsSubsetDefinition	Contains a list of subset definitions to filter the population's individuals based on an individual's properties, by property nam and a custom filter query, for example a value range or a list of keywords.

Table 3.278: Selection settings for module HBM concentrations.

Output settings

Table 3.279:	Output	settings fo	r module	HBM	concentrations.
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Name	Туре	Description
Exclude privacy sensitive data from outputs	Boolean	Use this setting to not report the parts of the results (i.e., figure tables, or sections) that are marked as (potentially) privacy sensitive.
Lower percentage for variability (%)	Numeric	The default value of 25% may be overruled.
Upper percentage for variability (%)	Numeric	The default value of 75% may be overruled.

Uncertainty settings

Name	Туре	Description
Resample HBM individuals	Boolean	HBM individual data are resampled from the original database using the bootstrap methodology (Efron 1979, Efron & Tibshin 1993).

Table 3.280: Uncertainty settings for module HBM concentrations.

HBM concentrations as data

Data are provided in the form of surveys consisting of individuals from which the human monitoring samples taken. Substance concentration measurements are linked to analyses performed on the human monitoring samples. The data should also include information about the analytical methods that were used.

- HBM concentrations data formats
- HBM concentrations from data

3.4.16 Non-dietary exposures

Non-dietary exposures are the amounts of substances to which individuals in a population are exposed to via any of three non-dietary routes: dermal, inhalation or oral. Non-dietary exposures are the input for the estimation of *aggre-gate (internal) exposure distributions* from both dietary and non-dietary routes of exposure. Depending on the period individuals are exposed, non-dietary exposures are short-term or acute exposure and are expressed per individual-day. When the exposure takes place over an unspecified longer time period, the exposure is long-term or chronic and is expressed as an average exposure per individual. Examples are presented as case studies in Kennedy et al. (Kennedy et al. (2015a), Kennedy et al. (2015b), Kennedy and Butler Ellis (2017)). The R-code to generate these examples is available for general use.

Datasets are typically generated by external programs, e.g. *BROWSE*, *BREAM* or *PACEM*. The BROWSE and BREAM2 models both simulate distributions of potential exposure of residents and bystanders to pesticides sprayed on crops. Probability distributions are included to quantify variations in input parameters representing conditions during a spray event. PACEM is a probabilistic exposure model for substances present in personal care products *PCPs* or house cleaning procucts *HCPs*. BROWSE was an EU FP7 project that, in addition to bystanders and residents from boom-sprayers, includes various arable and orchard scenarios. It includes dermal, oral and inhalation routes of exposure and generates exposure files in the correct format for MCRA non-dietary exposure calculations. The underlying simulation of dermal spray deposits on bystanders and residents was taken from BREAM, although BROWSE includes post-processing to model indirect exposures, multiple routes and long-term exposure, see Kennedy and Butler Ellis (2017). Volatilisation is also included through the PEARL-OPS model (van den Berg et al. (2016)) to account for inhalation of vapours. BREAM2 is an updated version of the original BREAM model (Kennedy et al. (2012)) and software is available online (http://www.ssau.co.uk/BREAM2-calculator). Results from BREAM had been used as part of EFSA guidance on bystander and resident exposure. BREAM2 was recently shown to produce more realistic exposure distributions, when compared to measured dermal exposure (Butler Ellis et al. (2018)). Currently, the BROWSE software is outdated and replaced by BREAM2.

This module has as primary entities: Populations Substances

Output of this module is used by: Internal exposures

Non-dietary exposures from data

Non-dietary exposures data formats

Non-dietary exposures may be specified for multiple routes of exposure (dermal, oral and inhalation), for multiple substances, and for multiple exposure sources. Also, they can be provided as single deterministic exposure levels or as probabilistic exposure estimates and it is possible, but not mandatory, to specify uncertainty. The non-dietary exposures may be short term (acute) or longer term averages (chronic), and the user must ensure to supply appropriate non-dietary data for the type of exposure assessment of interest. For chronic assessments this means the non-dietary exposure is averaged over an appropriate time interval.

Non-dietary exposures are defined by non-dietary surveys to which dietary exposures are linked. For these surveys, individual properties can be specified to define non-dietary exposures for particular sub-groups of the population (e.g., specific age groups, or a specific gender). For each non-dietary survey a percentage of the target population that is not exposed from this source can be specified by means of a percentage. Uncertainty about non-dietary exposures can be specified by specifying multiple records for each individual in an additional table.

The use of multiple surveys can be used when multiple sources are relevant. For example, when modelling individuals taking part in various activities involving pesticide use or incidental exposures as a resident. Each non-dietary source is characterised in a particular user-selected or user-supplied non-dietary survey. By default, exposures from separate non-dietary surveys (sources) are considered to be independent events, but as explained below correlations between substances and/or activity types per individual can be represented if generated prior to uploading to MCRA. When including multiple non-dietary surveys it is possible to supply some with uncertainty/variability and others without variability/uncertainty according to the requirements and data availability.

When the user supplies probabilistic non-dietary exposure estimates (i.e., there is a distribution for the non-dietary exposure rather than a single nominal value), then this information will be propagated as part of the *exposure assessment*. Distributions may be included to represent variability, uncertainty or both, and in these cases the aggregate exposure estimates are reported with variability and/or uncertainty as appropriate. Multiple (uncertain) values from the non-dietary exposure distribution may be supplied per individual and per substance.

Exposures within a non-dietary survey may be expressed as correlated or independent for the different substances. For example, if the exposures are a mixture of substances in a known ratio (e.g. from a specific tank mix of pesticides), or if exposure to one substance strongly implies that exposure to another is likely, these relationships may be included in the non-dietary data supplied by the user. Inference for the matched-case scenario with uncertainty analysis can use exposure sets. These are specific sets of exposures defined for each individual and in any uncertainty iteration an individual will receive exactly one of the exposure sets for that individual. Alternatively, independence may be represented by generating sets drawn from independent distributions when generating these tables.

Non-dietary exposure data is provided per non-dietary surveys. Each non-survey has some general information about the exposed population and the origin of the non-dietary exposure data. Also, a number of properties, such as specific age groups, can be specified for a survey. To each non-dietary survey, non-dietary exposures can be linked. These exposures may originate from dermal, oral and/or inhalatory exposure routes.

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Non-dietary surveys

This table provides detail about non-dietary surveys (source of non-dietary exposure): description, location, date and unit of exposure).

Name	Туре	Description	Aliases	Required
idNonDietary- Survey	AlphaNumeric (50)	The survey identification number.	idNonDietary- Survey	Yes
Name	AlphaNumeric (100)	Name of the non-dietary survey.	Name	No
Description	AlphaNumeric (200)	Description of non-dietary survey.	Description	No
Location	AlphaNumeric (50)	The location of survey.	Location	No
Date	DateTime	The date of survey.	Date	No
NonDietary- IntakeUnit	ExternalExposure- Unit	The unit of the non-dietary exposure.	Unit, NonDietary- IntakeUnit, NonDietary- ExposureUnit	Yes
Percentage- Zeros	Numeric	The proportion zeros, specified as a percentage (%).	PercentageZeros	No
idPopulation	AlphaNumeric (50)	Unique identification code of the population.	IdPopulation, PopulationId	No

Table 3.281: Table definition for Non-dietary surveys.

Accepted table names: NonDietarySurveys, NonDietarySurvey.

Non-dietary survey properties

This table specifies demographic properties that apply to the individuals in the surveys. These properties could be used to link the individuals of a non-dietary survey with individuals from dietary surveys. That is, if demographic criteria are defined, only those individuals in the dietary survey that meet these criteria will be assigned non-dietary exposures. This table is not relevant when matching is switched on (i.e., when individuals are matched based on individual id).

Name	Туре	Description	Aliases	Required
Individual- PropertyName	AlphaNumeric (50)	Name of demographic criteria for non-dietary exposures in a particular survey e.g. age, gender, height (must correspond to a column name in Individuals table).	Individual- PropertyName	Yes
idNonDietary- Survey	AlphaNumeric (50)	The code of survey (must correspond to values in id column of the non-dietary surveys table).	idNonDietary- Survey	Yes
Individual- PropertyText- Value	AlphaNumeric (50)	Text value of the property e.g. male or female, smoker or non-smoker.	Individual- PropertyText- Value	No
Individual- Property- DoubleValue- Min	Numeric	Inclusive lower bound value of the property. E.g., a value of "18" for an individual property name called Age would mean that only individuals aged 18 and above receive the non-dietary exposures.	Individual- PropertyDouble- ValueMin	No
Individual- Property- DoubleValue- Max	Numeric	Inclusive upper bound value of property e.g. a value of "65" for an IndividualPropertyName called Age would mean that only individuals aged 65 and below receive the non-dietary exposures.	Individual- PropertyDouble- ValueMax	No

Table	3.282:	Table	definition for	Non-dietarv	survev	properties.
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Accepted table names: NonDietarySurveyProperties, NonDietarySurveyProperty.

Non-dietary exposures

This table defines nominal non-dietary exposure values (such as means) for individuals within the non-dietary surveys. It can also be used to specify non-dietary exposures for individuals within the food surveys. Each exposure comprises a non-dietary survey (source of exposure); a string identifying an individual, which may or may not correspond to the ID of an individual in a food survey; a substance; and dermal, oral and inhalation exposure values. Exposures are assumed to be external doses.

Name	Туре	Description	Aliases	Required
idIndividual	AlphaNumeric (50)	Non-dietary individual identification number. This id may 1) match with the individual ids of the dietary survey (dietary exposures matched to food survey individuals), 2) not match with the individual ids of the dietary survey (unmatched individuals), or contain a default exposure (indicated by idIndividual = 'General') linking the dietary exposures to individuals based on the demographic criteria defined in the non-dietary survey properties table.	idIndividual	Yes
idNonDietary- Survey	AlphaNumeric (50)	The code of the survey (must correspond to values in id column of non-dietary surveys table).	idNonDietary- Survey	Yes
idSubstance	AlphaNumeric (50)	The substance code.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
Dermal	Numeric	The dermal (non-dietary) exposure value.	Dermal	Yes
Oral	Numeric	The oral (non-dietary) exposure value.	Oral	Yes
Inhalation	Numeric	The inhalation (non-dietary) exposure value.	Inhalation	Yes

Table 3.28	33: Table	definition	for Non-	dietary ex	posures.
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Accepted table names: NonDietaryExposures, NonDietaryExposure.

Non-dietary exposure uncertainty records

This table may be used to supply uncertainty sets of multiple (uncertain) non-dietary exposure values for individuals within the non-dietary surveys. Multiple non-dietary values are generated by probabilistic exposure calculations i.e. when there is a distribution for the non-dietary exposure rather than a single nominal value. If this table is supplied, aggregate exposure estimates will be reported with uncertainty using the uncertainty set approach. Each exposure set comprises a non-dietary survey (source of exposure); an individual ID; a substance; and dermal, oral and inhalation exposure values. In addition, the id column is used to define the uncertainty set. Summarizing, an uncertainty set is identified by column id and contains all exposure sets defined for each individual. In each uncertainty run (outer loop) an uncertainty set is sampled and in each iteration (inner loop) nondietary individuals are sampled from this set.

Name	Туре	Description	Aliases	Required
idIndividual	AlphaNumeric (50)	Non-dietary individual identification number. The idIndividual value may correspond to an id in the Individuals table (dietary exposures matched to food survey individuals), may not correspond to an id in the Individuals table (unmatched individuals), or may contain a default exposure (indicated by idIndividual = 'General' - demographic criteria for the assignment of exposures are defined in the NonDietarySurveyProperties table). For matching to occur, the user will need to tick the option to 'match specific dietary survey individuals' in the user-interface. The software will then assign non-dietary exposures to the dietary individuals according to the values in this column. Any idIndividual values that do not correspond to individuals in the food survey will be ignored, unless a value 'General' is specified. Then the individual should meet the demographic criteria as defined in the NonDietarySurveyProperties table. If this box is left unticked, the non-dietary exposures will be randomly allocated to the dietary population provided they meet	idIndividual	Yes
idNonDietary- Survey	AlphaNumeric (50)	code of survey (must correspond to values in id column of NonDietarySurveys table)	idNonDietary- Survey	Yes
idCompound	AlphaNumeric (50)	Substance code (must correspond to values in id column of Substances table).	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
id	AlphaNumeric (50)	Uncertainty set identification number.	id	Yes
Dermal	Numeric	Dermal non-dietary exposure value.	Dermal	Yes
Oral	Numeric	Oral non-dietary exposure value.	Oral	Yes
Inhalation	Numeric	Inhalation (non-dietary)	Inhalation	Yes

Table 3.284: Table definition for Non-dietary exposure uncertainty records.

Accepted table names: NonDietaryExposuresUncertain, NonDietaryExposureUncertain.

Non-dietary exposures settings

Calculation settings

Table 3.285:	Calculation	settings for	or module	Non-dietary	exposures.
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Name	Туре	Description
NON-DIETARY: Population alignment method other non-dietary exposures	PopulationAlignmentMethod	Population aligment methods for nondietary exposure sources.
Match individuals of multiple non-dietary surveys	Boolean	If checked, then non-dietary exposures of individuals from different surveys with the same identifier (ID) are combined.

Uncertainty settings

Table 3.286: Uncertainty settings for module Non-dietary exp
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Name	Туре	Description
Resample non-dietary exposures	Boolean	Specifies whether non-dietary exposures are resampled. Note the non-dietary uncertainty is only ignored when individual uncertainty is set to false (uncheck box: do NOT resample individuals).

Non-dietary exposures uncertainty

In an aggregate exposure assessment, dietary and nondietary data are combined into an aggregate exposure distribution. The nondietary data are supplied in table *Non-dietary exposures*. In an uncertainty analysis, MCRA provides two ways to assess the uncertainty:

- 1. the uncertainty set approach
- 2. the bootstrap algorithm.

When table *Non-dietary exposure uncertainty records* is not supplied, the nondietary data in table *Non-dietary exposures* is resampled and the bootstrapped sets are used in the uncertainty run. More precisely, in each outer loop of the 2D Monte Carlo, within each nondietary survey (multiple surveys may be supplied), the nondietary individuals are resampled. Each individual represents a nondietary exposure set containing dermal and/or oral and/or inhalation exposure values for multiple substances. Bootstrapping is the default behaviour when the *Non-dietary exposure uncertainty records* table is missing. When uncertainty distributions supplied in this table represent sampling uncertainty (individual exposure sets are repeatedly sampled using the same nondietary exposure generator without changing the input parameters), then bootstrapping the data performs equally well and is more efficient.

Non-dietary exposures as data

Non-dietary exposures are collected in non-dietary surveys. Data may be specified on population level or individual level, and may or may not include variability and uncertainty.

- Non-dietary exposures data formats
- Non-dietary exposures from data

Inputs used: Active substances

Settings used

• Calculation Settings

See also Combining exposure sources.

3.4.17 Single value dietary exposures

Single value dietary exposures are based on the single value concentrations of substances, expressed per standard (kg) bodyweight and/or single value amounts of consumed modelled food. Depending on the exposure type, dietary exposures can be short-term/acute exposures.

This module has as primary entities: Populations Foods Substances

Output of this module is used by: Single value risks

Single value dietary exposures calculation

Either the short-(acute) or long-term (chronic) dietary exposure to a substances via food can be estimated as a single value calculated from single value inputs. This is often referred to as deterministic estimation. MCRA implements the IESTI, TMDI, IEDI and NEDI (Rees-Day) calculation methods that are also available in the EFSA PRIMo (Pesticide Residue Intake Model) tool revision 3 (EFSA (2018)).

The implementation in MCRA allows more choices than EFSA PRIMo by choosing other inputs or input combinations for the calculation formula. Moreover, the calculations can in all cases be adapted for processing factors or occurrence frequencies. For the chronic estimates, also the contributions per food or processed food are reported.

Acute single value dietary exposure assessment

The short term (acute) exposure assessment is usually the exposure related to a consumption of food over a single day. MCRA applies in principle the IESTI equations as shown in EFSA PRIMo revision 3 (EFSA (2018)), but the equations are extended with a factor OF to allow adaptation for an occurrence frequency lower than 1. So the inputs to the equations are not necessarily the same as used in PRIMo. For example, the large portion (LP) and body weight (BW) can be computed instead of just being standard values.

IESTI (International Estimated Short-Term Intake)

The IESTI (International Estimated Short-Term Intake) is calculated according to different equations depending on the unit weight of the raw agricultural commodity (RAC) and the unit weight of the edible portion (EP). The following cases are distinguished.

Case 1

refers to commodities with unit weight of the raw agricultural commodity $U_{RAC} \leq 25$ g (e.g. walnuts, strawberries and peas. It is also used for meat, liver, kidney, edible offal, eggs and for post-harvest uses in cereal grains, oilseeds and pulses).

Case 2a

for food product with a $\rm U_{RAC}>25~g,$ where the meal portion is $\rm > U_{ep}$ (unit weight edible portion).

Case 2b

for food products with a ${\rm U_{RAC}}>25$ g, where the meal portion is $\leq {\rm U_{ep}}.$

Case 3

for food products that are usually bulked or blended before they are consumed (e.g. cereals, pulses, oilseeds and milk).

The calculations are as follows.

Case 1

Case 2a

 $\frac{\mathbf{U}_{\text{ep}} \cdot \mathbf{HR} \cdot \mathbf{PF} \cdot \mathbf{CF} \cdot \mathbf{VF} \cdot \mathbf{OF} + (\mathbf{LP} - \mathbf{U}_{\text{ep}}) \cdot \mathbf{HR} \cdot \mathbf{PF} \cdot \mathbf{CF} \cdot \mathbf{OF}}{\mathbf{BW}}$

Case 2b

$$\frac{\text{LP} \cdot \text{HR} \cdot \text{PF} \cdot \text{CF} \cdot \text{VF} \cdot \text{OF}}{\text{BW}}$$

Case 3

$$\frac{\text{LP} \cdot \text{STMR} \cdot \text{PF} \cdot \text{CF} \cdot \text{OF}}{\text{BW}}$$

New Case 1 and 3:

New Case 2a and 2b

$$\frac{\text{LP} \cdot \text{MRL} \cdot \text{PF} \cdot \text{CF} \cdot \text{VF} \cdot \text{OF}}{\text{BW}}$$

Parameters used in the equations

MRL: Maximum residue level for the RAC concerned (default in mg/kg);

STMR: Supervised Trials Median Residue for raw agricultural commodity (RAC) concerned (default in mg/kg);

CF: Conversion factor residue definition enforcement to residue definition risk assessment (calculated as the ratio of residues according to the residue definition for risk assessment divided by the residue concentration according to the residue definition for enforcement);

OF: Use Frequency of the raw agricultural commodity (RAC),

BW: body weight of the population related to the LP (default in kg);

LP: Large portion reported (in kg/day) (97.5th percentile of eaters (or alternative percentile, depending on the number of reported eating occasions);

HR: Highest residue according to residue definition for enforcement in composite sample (default in mg/kg);

 U_{ep} : Unit weight of edible portion (in kg), provided by the country from which the LP was reported (or mean unit weight calculated from all available unit weight data, if no unit weight is available from the country matching the highest LP;

PF: Processing factor or peeling factor (calculated as the ratio of residues in processed/peeled product, divided by residue concentration in unprocessed/unpeeled product);

vF: variability factor, depending on the unit weight of the whole product (U_{RAC}) , different default vF are used in the calculations.

 $(U_{RAC}) < 25$ g, the calculations are performed according to case 1 (VF = 1).

 (U_{RAC}) between 25 and 250 g: VF = 7.

 (U_{RAC}) greater than 250: VF = 5.

In $IESTI_{new}$, a default VF of 3 is used.

In case the empirically derived variability factors are available, the default VF is to be replaced.
Alternative IESTI-styled assessments

If consumption survey data for a specific population are available, the LP values in the IESTI equations may be replaced by statistics calculated from these data (at the consumed modelled food level).

If concentration monitoring data (retrospective) or concentration field trial data (prospective) are available, the MRL, HR, STMR values in the IESTI equations may be replaced by statistics calculated from these data (at the consumed modelled food level).

In the current use of IESTI, the occurrence frequency (use frequency) OF is assumed to be 1. In alternative assessments, a more realistic estimate may be used. Such an estimate could be derived for example as the highest occurrence frequency observed in a retrospective assessment for either the same substance or the same food.

IESTI special cases

For some foods, chemicals are applied after harvest, i.c. post-harvest use. For those combinations of food and substance, Case 1 should be used in the calculation. However, commodities with post-harvest use like cereal grains, oilseeds and pulses are typically bulked or blended (Case 3). To overrule Case 3, specify in table *IESTI special cases* the food and substance combination with *PostHarvest* as application type. For those food and substance combinations with a unit weight of the raw agricultural commodity $U_{RAC} \leq 25$ g, Case 1 is applied. When substances are applied before harvest, i.c. pre-harvest use, Case 1 should be overruled by Case 3. Specify in table *IESTI special cases* the food and substance combination with *PreHarvest* as application type.

Chronic single value dietary exposure assessment

The long term (chronic) exposure assessment is usually the exposure related to a consumption over a longer period of time. MCRA applies in principle the TMDI, IEDI or NEDI (Rees-Day) equations as shown in EFSA PRIMo revision 3 (EFSA (2018)). However, the equations are extended with factors PF_i and OF_i to allow adaptation for processing factors and occurrence frequencies lower than 1. Also, the inputs to the equations are not necessarily the same as used in PRIMo. For example, the consumption statistics (MC, $p_{97.5}$) and body weight (BW) can be computed instead of just being standard values. Note that TMDI, IEDI and NEDI (Rees-Day) estimates are summations over foods (raw agricultural products). In addition to the summations, MCRA will also report the individual terms (single value dietary exposures per food).

TMDI (Theoretical Maximum Dietary Intake)

$$\sum_{X=i}^{n} \frac{MRL_i \cdot CF_i \cdot PF_i \cdot OF_i \cdot MC_i}{BW}$$

i, j, k, ...n: individual raw agricultural products

IEDI (International Estimated Dietary Intake)

$$\sum_{Y=i}^{n} \frac{STMR_i \cdot CF_i \cdot PF_i \cdot OF_i \cdot MC_i}{BW}$$

i, j, k, ...n: individual raw agricultural products

NEDI (National Estimated Dietary Intake): Rees-Day model (I)

$$\sum_{X=i}^{j} \frac{\textit{MRL}_{i} \cdot \textit{CF}_{i} \cdot \textit{PF}_{i} \cdot \textit{OF}_{i} \cdot \textit{p}_{97.5}\textit{consumption}_{i}}{BW} + \sum_{X=k}^{n} \frac{\textit{MRL}_{k} \cdot \textit{CF}_{k} \cdot \textit{PF}_{i} \cdot \textit{OF}_{i} \cdot \textit{MC}_{k}}{BW}$$

i, *j*: two raw agricultural products leading to the highest intake;

k, l, m, ...n: remaining raw agricultural commodities consumed

NEDI (National Estimated Dietary Intake): Rees-Day model (II)

$$\sum_{X=i}^{j} \frac{STMR_i \cdot CF_i \cdot PF_i \cdot OF_i \cdot p_{97.5} consumption_i}{BW} + \sum_{X=k}^{n} \frac{STMR_k \cdot CF_i \cdot PF_i \cdot OF_i \cdot MC_k}{BW}$$

i, *j*: two raw agricultural products leading to the highest intake;

k, l, m, ...n: remaining raw agricultural commodities consumed

Parameters used in the equations

MRL_i: Maximum residue level for the RAC concerned (default in mg/kg);

 $STMR_i$: Supervised Trials Median Residue for raw agricultural commodity (RAC) concerned (default in mg/kg);

 CF_i : Conversion factor residue definition enforcement to residue definition risk assessment (calculated as the ratio of residues according to the residue definition for risk assessment divided by the residue concentration according to the residue definition for enforcement);

 MC_i : mean consumption for a given raw agricultural product (RAC) calculated for the whole survey/subgroup of the survey, including processed products (recalculated to the unprocessed RAC) (default in kg/day);

 $p_{97.5}$ consumption_i for a given raw agricultural product (RAC), calculated from the individual consumption reported by the participants of the whole survey/subgroup of the survey, including processed products (recalculated to the unprocessed RAC) (default in kg/day);

BW: mean body weight of the population related to the LP or mean consumption (default in kg).

 OF_i : Occurrence Frequency of the substance on the food (typically, a raw agricultural commodity, RAC),

 PF_i : Processing factor or peeling factor (calculated as the ratio of residues in processed/peeled product, divided by residue concentration in unprocessed/unpeeled product);

Alternative TMDI-, IEDI- or NEDI-styled assessments

If consumption survey data for a specific population are available, the MC, $p_{97.5}$ consumption values in the IESTI equations may be replaced by statistics calculated from these data (at the consumed modelled food level).

If concentration monitoring data (retrospective) or concentration field trial data (prospective) are available, the MRL, STMR values in the IESTI equations may be replaced by statistics calculated from these data (at the consumed modelled food level).

In the current use of IESTI, the occurrence frequency (use frequency) OF is assumed to be 1. In alternative assessments, a more realistic estimate may be used. Such an estimate could be derived for example as the highest occurrence frequency observed in a retrospective assessment for either the same substance or the same food.

Single value dietary exposures data formats

Single value dietary exposures are IESTI etc.

Dietary exposure data is specified through dietary exposure models. To each dietary exposure model, exposure distributions are linked.

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Dietary exposure models

High level description of the dietary exposure models, specifying the id, name, description and the (reference) substance and exposure unit used for reporting the exposures. To this models, exposure percentiles and bootstrap values of the percentile may be linked.

Name	Туре	Description	Aliases	Required
idDietary- ExposureModel	AlphaNumeric (50)	Identifier of the dietary exposure model.	id, idDietary- Exposure, idExposure- Model	Yes
Name	AlphaNumeric (100)	The name of the dietary exposure model.	Name	No
Description	AlphaNumeric (200)	Description of dietary exposure model.	Description	No
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
ExposureUnit	AlphaNumeric (50)	The intake/exposure unit of the dietary exposures reported by this model. If not specified, then a default exposure unit of mg/kg BW/day is assumed.	Unit, ExposureUnit, IntakeUnit	Yes

Accepted table names: DietaryExposureModels, DietaryExposures.

Dietary exposure percentiles

Exposure percentiles linked to a dietary exposure model. The percentiles are reported in the unit specified by the exposure model to which they belong.

Name	Туре	Description	Aliases	Required
idDietary- ExposureModel	AlphaNumeric (50)	The code of the dietary exposure model to which this record belongs.	idDietary- ExposureModel	Yes
Percentage	Numeric	The percentage to which the percentile value belongs.	Individual- PropertyDouble- ValueMin	Yes
Exposure	Numeric	The percentile value. I.e., the exposure value belonging to the specified percentage.	Exposure	Yes

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Table 3.288:	Table	definition	IOL	Dietary	exposure	percentiles.

Accepted table names: DietaryExposurePercentiles.

Dietary exposure percentile bootstrap values

Uncertainty values, obtained from bootstrap runs, of the dietary exposure percentiles.

Name	Туре	Description	Aliases	Required
idDietary- ExposureModel	AlphaNumeric (50)	The code of the dietary exposure model to which this record belongs.	idDietary- ExposureModel	Yes
idUncertainty- Set	AlphaNumeric (50)	The uncertainty set identifier.	idUncertainty- Set, UncertaintyId	Yes
Percentage	Numeric	The percentage to which the percentile value belongs.	Individual- PropertyDouble- ValueMin	Yes
Exposure	Numeric	The percentile value. I.e., the exposure value belonging to the specified percentage.	Exposure	Yes

Table 3.289: Table definition for Dietary exposure percentile bootstrap values.

 $\label{eq:linear} Accepted \ table \ names: \ Dietary Exposure Percentiles Uncertain, \ Dietary Exposure Percentile Uncertains.$

Single value dietary exposures settings Selection settings

Name	Туре	Description
Exposure type	ExposureType	The type of exposure considered in the assessment; acute (shor term) or chronic (long-term).
Selected tier	SettingsTemplateType	Specifies all module settings should be set according to a pre-defined tier or using custom settings.

Table 3.290: Selection settings for module Single value dietary exposures.

Calculation settings

Name	Туре	Description
Single value dietary exposure calculation method	SingleValueDietaryExpo- suresCalculationMethod	Method for computing single value dietary exposures.
Apply processing factors	Boolean	Specified in table ProcessingFactor. If checked, processing fact are applied. Concentrations in the consumed food may be different from concentrations in the modelled food in monitorir programs (typically raw food) due to processing, such as peelin washing, cooking etc. If unchecked, no processing information used. This is in most (though not all) cases a worst-case assumption
Report consumptions and exposures per individual instead of per kg body weight	Boolean	Specifies whether body weights should be ignored and consumptions and exposures should be expressed per individual Otherwise, the consumptions and exposures are per kg body weight.
Use occurrence frequencies	Boolean	Account for occurrence frequencies for combinations of food a substance in the exposure calculations.
Use unit variability	Boolean	Controls whether to use unit variability.
Derived modelled foods for	ModelledFoodsCalculation-	The derived modelled foods for the specified concentration sou
concentration source	Source	that is used in the conversion algorithm to determine for which modelled food concentrations are available. This source is 1) a single value concentrations data source or 2) a sampled concentrations data source.

Table 3.291: Calculation settings for module Single value dietary exposures.

Output settings

Table 3.292: Output settings for module Single value dietary exposures.

Name	Туре	Description
Lower percentage for variability (%)	Numeric	The default value of 25% may be overruled.
Upper percentage for variability (%)	Numeric	The default value of 75% may be overruled.

Uncertainty settings

Table 3.293: Uncertainty settings for module Single value dietary expo-

sures.	

Name	Туре	Description
Lower uncertainty limit (%)	Numeric	Percentage lower bound, e.g. 2.5%.
Upper uncertainty limit (%)	Numeric	Percentage upper bound, e.g. 97.5%.

Calculation of single value dietary exposures

Single value dietary exposures are calculated from single value consumptions per modelled food and single value concentrations. Optionally, also processing factors, unit variability models and use frequencies are applied.

• Single value dietary exposures calculation

Inputs used: Single value consumptions Single value concentrations Processing factors Unit variability factors Occurrence frequencies

Settings used

• Calculation Settings

3.4.18 Single value non-dietary exposures

Single value non-dietary exposures are based on the single value concentrations or amounts of substances, as opposed to the distribution-based exposures of individuals in the non-dietary exposure module. Exposures are via any of the non-dietary routes: dermal, inhalation, or oral. Depending on the exposure type, single value non-dietary exposures can be short-term/acute or long term/chronic exposures. The exposures can be modelled as external exposures or internal exposures.

This module has as primary entities: Populations Substances

Single value non-dietary exposures from data

Single value non-dietary exposures calculation

Either the short-(acute) or long-term (chronic) non-dietary exposures to substances via inhalation, dermal or oral routes can be estimated.

Acute single value non-dietary exposure assessment

The short term (acute) exposure assessment is usually the exposure related to inhalation, dermal or oral contact of substances over a single day.

Chronic single value non-dietary exposure assessment

The long term (chronic) exposure assessment is the exposure related to dermal, oral or inhalation route(s) over a longer period of time (months or years).

Single value non-dietary exposures data formats

Single value non-dietary exposure estimates are supported from external exposures or internal exposures.

Single value non-dietary exposure data is provided per exposure scenario.

Download empty dataset template: Zipped CSV Excel

Exposure scenarios

This table provides details about exposure scenarios.

Name	Туре	Description	Aliases	Required
idExposure- Scenario	AlphaNumeric (50)	The identifier of the exposure scenario.	idExposure- Scenario, Exposure- ScenarioId	Yes
Name	AlphaNumeric (100)	Name of the exposure scenario.	Name	No
Description	AlphaNumeric (200)	Description of exposure scenario.	Description	No
idPopulation	AlphaNumeric (50)	Unique identification code of the population.	IdPopulation, PopulationId	No
ExposureType	ExposureType	The exposure type associated with the exposure scenario (i.e., chronic or acute).	ExposureType	Yes
ExposureLevel	TargetLevelType	The target level. I.e., internal or external. If omitted, external is assumed.	ExposureLevel, LevelExposure	No
ExposureRoutes	AlphaNumeric (100)	The exposure route(s) (only applicable if exposure level is external) for which exosure estimates are collected.	ExposureRoutes, Routes	No
ExposureUnit	ExternalExposure- Unit	The unit of the non-dietary exposure.	Unit, NonDietary- IntakeUnit, NonDietary- ExposureUnit	Yes

Table 3.294: Table definition for Exposure scenarios	5.
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Accepted table names: ExposureScenarios, ExposureScenarios.

Exposure determinants

This table is used to describe the custom properties used in the exposure estimates table.

Name	Туре	Description	Aliases	Required
idExposure- Determinant	AlphaNumeric (50)	The code of the exposure determinant.	IdExposure- Determinant, Exposure- DeterminantId	Yes
Name	AlphaNumeric (100)	The name of the exposure determinant.	Name	No
Description	AlphaNumeric (200)	Description of the exposure determinant.	Description	No
Туре	IndividualProperty- Type	This field specifies the type of the values of this exposure determinant. This type follows a controlled terminology, with possible values: Boolean, Categorical (default), Numeric, Nonnegative, Integer, NonnegativeInteger, Month, Datetime, Gender. If not specified, categorical is assumed.	Туре	No

Table 3.295: Table definition for Exposure determinants.

Accepted table names: ExposureDeterminants.

Exposure determinant combinations

This table is used to fill in the values of the exposure determinants.

Name	Туре	Description	Aliases	Required
idExposure- Determinant- Combination	AlphaNumeric (50)	The code of the exposure determinant combination.	IdExposure- Determinant- Combination, Exposure- Determinant- CombinationId, Exposure- Determinant- Combination	Yes
Name	AlphaNumeric (100)	The name of the exposure determinant combination.	Name	No
Description	AlphaNumeric (200)	Description of the exposure determinant combination.	Description	No

Table	3 296.	Table	definition	for	Exposure	determinant	combinations
1 abie	5.290.	rable	deminition	101	Exposure	ueterminant	combinations.

Accepted table names: ExposureDeterminantCombinations.

Exposure determinant values

This table describes exposure determinant values, for example determinants like face shields or application techniques like vehicle-mounted application of pesticides.

Name	Туре	Description	Aliases	Required
idExposure- Determinant- Combination	AlphaNumeric (50)	The code of the exposure determinant combination to which the determinant value is attached. The provided exposure estimate code should match with a code of the exposure estimate table.	idExposure- Determinant- Combination, Exposure- Determinant- CombinationId, Exposure- Determinant- Combination, Exposure- Determinant- Combination- Code	Yes
PropertyName	AlphaNumeric (50)	The identifier of the exposure determinant.	IdExposure- Determinant, idExposure- Determinant	Yes
TextValue	AlphaNumeric (100)	The value of the determinant as text value.		No
DoubleValue	Numeric	The value of the determinant as number.		No

Table 3.	297: Table	definition	for Exposu	re determinant	values.
14010 01		<i>ao</i> mininon	101 2		

Accepted table names: ExposureDeterminantValues, ExposureDeterminantVal.

Exposure estimates

This table defines non-dietary exposure values (such as means) for exposure scenarios. Exposures can be internal and external. Exposures are assumed to be external doses.

Name	Туре	Description	Aliases	Required
idExposure- Scenario	AlphaNumeric (50)	The identifier of an exposure scenario.	idExposure- Scenario, Exposure- ScenarioId	Yes
idExposure- Determinant- Combination	AlphaNumeric (50)	The identifier of an exposure determinant combination.	idExposure- Determinant- Combination, Exposure- Determinant- CombinationId, Exposure- Determinant- Combination	No
ExposureSource	AlphaNumeric (100)	The origin of substance or a mixture for the purposes of an exposure assessment, in immediate contact with outer surfaces of exposure, to which individuals in the target population may be exposed. Examples are food, drinking water, consumer products, indoor air, outdoor air, dust and soil.	Source	No
idSubstance	AlphaNumeric (50)	The substance code.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
ExposureRoute	ExposureRoute	The exposure route (only applicable if exposure level is external). If not specified and external level, then Dietary is assumed.	ExposureRoute	No
Value	Numeric	The exposure value.	Value, ExposureValue	Yes
EstimateType	AlphaNumeric (100)	An indication of the measure by which the exposure value has been obtained, e.g. a 95th percentile (P95) or as a mean (Mean).	Source	No

Table 3.298: Table definition for Exposure estimates.

Accepted table names: ExposureEstimates, ExposureEstimate.

Single value non-dietary exposures settings

Selection settings

Table 3.299: Selection settings for module Single value non-dietary expo-

Name	Туре	Description
Exposure type	ExposureType	The type of exposure considered in the assessment; acute (shor term) or chronic (long-term).

Calculation settings

Table 3.300: Calculation settings for module Single value non-dietary ex-

posures.

Name	Туре	Description
Configuration	AlphaNumeric	Select the configuration for computing the non-dietary exposur

Single value non-dietary exposures as data

Single-value non-dietary exposures estimates are collected in exposure scenarios.

- Single value non-dietary exposures data formats
- Single value non-dietary exposures from data

Settings used

• Calculation Settings

Calculation of single value non-dietary exposures

Single value non-dietary exposures are calculated from single value exposure values.

• Single value non-dietary exposures calculation

Settings used

• Calculation Settings

3.5 Hazard modules

Hazard data exist at two levels: at a lower level *dose response data* give *responses* measured in *test systems* from doses of *active substances*. Such data can be modelled with *dose response models*.

At a higher level *responses* can be linked to *effects*, optionally via *AOP networks*, using *effect representations*. If benchmark responses (BMRs) have been specified, *dose response models* can calculate Benchmark Doses (BMDs), which are the preferred Points of departure in hazard assessments. In addition, or alternatively, external *points of departure* can be specified for *active substances* and *effects*.

BMDs from *dose response models* and/or other *points of departure* can be converted to *hazard characterisations* at the intended level (external or internal dose, without or with safety factors), using *kinetic models, inter-species conversions* and/or *intra-species factors*. Finally, *hazard characterisations* can be translated to *relative potency factors*.

3.5.1 Active substances

Active substances are the substances that may lead with non-zero probability (P (AG)>0) to a specific *health effect* (adverse outcome). In the simplest case, all substances in the scope of the action will form one assessment group (AG) of active substances. In more advanced cases, the list of active substances is derived from possibly multiple assessment group memberships, which are scores for substances that determine whether a substance is included (score > 0) or excluded (score = 0) in the set of active substances. Substances with membership 0 are excluded from the list of active substances. Memberships scores between 0 and 1 are treated as probabilities of being in the set of active substances. Assessment group memberships can be either specified directly as data or derived from *QSAR membership models*, *molecular docking models*, or from availability of *points of departure*.

This module has as primary entities: Effects Substances

Output of this module is used by: Concentrations Single value concentrations Occurrence patterns Occurrence frequencies Substance conversions Non-dietary exposures Dust exposures Soil exposures Air exposures Consumer product exposures Absorption factors PBK model parametrisations Kinetic conversion factors Relative potency factors Hazard characterisations Inter-species conversions Intra species factors Concentration models Food conversions High exposure food-substance combinations Dietary exposures Internal exposures HBM analysis

Active substances from data

Active substances calculation

Depending on the *model settings*, the set of active substances for a specified effect is computed in several ways:

- 1. From the list of substances with available *points of departure (POD) data* for the specified effect. If there is a POD, then the substance is considered an active substance with membership 1. If not, the membership is 0, and the substance is excluded from the list of active substances.
- 2. From one or more in-silico (QSAR and/or molecular docking) models. The results of the in-silico models should be provided as *QSAR membership models data* and/or *molecular docking models data*. Binding energies from molecular docking models are first translated to crisp memberships using a threshold value. The results from multiple in-silico models can be combined in any of four membership calculation methods:
 - 1. (crisp, any) the substance is considered an active substance if any in-silico model indicates activity;
 - 2. (crisp, majority) the substance is considered an active substance if the majority of in-silico models indicates activity;
 - 3. (probabilistic, ratio) the membership probability is the fraction of in-silico models that indicate activity;
 - 4. (probabilistic, Bayesian) the membership probability is calculated using a Bayesian model according to Kennedy et al. (2020) and a specified prior probability (which is by default 0.5).

For substances within the scope of the assessment but without in-silico data, the default is to omit them from the assessment group. Set option *Include substances without membership information* to include them in the assessment group.

3. From a combination of 1 and 2, using either the union (OR) method or the intersection (AND method) of results.

Active substances data formats

Active substances as data have to be specified via assessment group (AG) memberships in an AG membership model. For each effect one or more AG membership models can be available, one of which should be chosen in assessments. The AG memberships can be crisp, i.e. a positive list of active substances (with default memberships 1, although it is also allowed to include the negative memberships with membership 0 explicitly) or probabilistic ($0 \le P \le 1$).

Assessment group membership models contain substance membership definitions for a given (health) effect. This data is described using two tables: the assessment group membership models table and the assessment group memberships table. The groups for a specified health effect are defined in the assessment group membership models table. The assessment group memberships table describes the substance memberships (or membership probabilities) in each group.

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Assessment group membership models

This table contains the definitions of the assessment group membership models. Each model contains a id, name, an optional description, and refers to its related health effect.

Name	Туре	Description	Aliases	Required
id	AlphaNumeric (50)	The unique identification code of the assessment group membership model.	id, idModel, Model, idAssessment- GroupModel, Assessment- GroupModel, idGroup- Membership- Model, Group- Membership- Model	Yes
Name	AlphaNumeric (100)	The name of the assessment group membership model.	Name	No
Description	AlphaNumeric (200)	Description of the assessment group membership model.	Description	No
idEffect	AlphaNumeric (50)	The effect code.	idEffect, EffectId, Effect	Yes
idIndex- Substance	AlphaNumeric (50)	The id/code of the index substance.	idIndex- Substance, idReference- Substance, IndexSubstance- Id, Reference- SubstanceId	No
Accuracy	Numeric	If applicable, the accuracy of the assessment group membership model memberships.	Accuracy	No
Sensitivity	Numeric	If applicable, the sensitivity of the assessment group membership model.	Sensitivity	No
Specificity	Numeric	If applicable, the specificity of the assessment group membership model.	Specificity	No
Reference	AlphaNumeric (200)	External reference(s) to sources containing more information about the assessment group model.	References	No

Table 3 301.	Table definition	for Asse	sement aroun	membershin	models
14016 5.501.	Table definition	IOI ASSC	ssment group	membership	mouels.

Accepted table names: AssessmentGroupMembershipModels, AssessmentGroupMembershipModel.

Assessment group memberships

Substances belong to an assessment group with certainty (probability 1), or the membership are uncertain. This table allows to specify membership probabilities for assessment group membership models. The probability should be a value between zero and one. For example, set to 1 or 0, or prior probabilities, or probabilities or 0/1 values estimated from QSAR, from Molecular Docking or from expert elicitation. The table can contain prior or posterior memberships. Default membership are specified with an empty idSubstance field.

Name	Туре	Description	Aliases	Required
idGroup- Membership- Model	AlphaNumeric (50)	The id of the assessment group memberships model or source.	Model, idModel, idAssessment- Group- Membership- Model, Assessment- Group- Membership- Model, idGroup- Membership- Model, Group- Membership- Model, idGroup-	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
Group- Membership	Numeric	Probability of the substance for belonging to the assessment group for the effect. If omitted, the default is 1, i.e. certain membership.	Group- Membership, Membership- Probability, Probability, Assessment- Group- Membership	Yes

Table 3.302:	Table definition	for Assessment	group mem	berships.
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 $\label{eq:constraint} Accepted \ table \ names: \ Assessment Group Memberships, \ Assessment Group Membership.$

Active substances settings

Selection settings

Table 3.303:	Selection	settings	for	module	Active	substances.

Name	Туре	Description
Multiple effects analysis	Boolean	Specifies whether the analysis should consider multiple effects. Otherwise, a single focal effect should be selected.
Include related effects of AOP network	Boolean	Include all related key events of the AOP network.

Calculation settings

Name	Туре	Description
Filter by certain assessment group membership	Boolean	Filter substances by certain assessment group membership.
Restrict active substances to substances with available PODs	Boolean	Restrict assessment group membership based on presence/abser of points of departure.
Restrict active substances to substances with available hazard characterisations	Boolean	Restrict assessment group membership based on presence/abser of hazard characterisations.
Derive memberships from QSAR membership data	Boolean	Specifies whether QSAR membership data is used for computing the assessment group memberships.
Derive memberships from molecular docking data	Boolean	Specifies whether molecular docking data is used for computing the assessment group memberhips.
Include substances without membership information	Boolean	For non-probabilistic methods: specifies whether substances for which no membership information is available in the specified inputs should be included in the assessment group.
Combination method memberships from available PODs and in-silico data	CombinationMethodMember- shipInfoAndPodPresence	Specifies whether to take the intersection or the union of the set substances with available PoDs and the set of substances with positive/probable (in-silico) membership score.
Membership calculation method	AssessmentGroupMembership- CalculationMethod	Calculation method for computing assessment group memberships: majority/any (crisp methods), ratio/Bayesian (probabilistic methods)
Default/prior membership probability	Numeric	Default substance membership probability for which no membership information is available in the specified inputs. Pri probability for Bayesian method.
Use probabilistic assessment group memberships	Boolean	Specifies whether substance memberships should be expressed terms of probabilities (probabilistic). Otherwise, substance memberships are expressed as in or out (crisp).

Table 3.304: Calculation settings for module Active substances.

Uncertainty settings

Table 3.305: U	Uncertainty	settings for	or module	Active	substances
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Name	Туре	Description
Resample assessment group memberships	Boolean	Specifies whether assessment group memberships of substances should be resampled using the assessment group membership probabilities.

Active substances as data

When provided as data, in the form of assessment group memberships, the active substances are derived from the specified memberships.

- Active substances data formats
- Active substances from data

Inputs used: AOP networks Points of departure Hazard characterisations

Settings used

• Calculation Settings

Calculation of active substances

Active substances and assessment group memberships may be computed from PoD presence of in-silico data.

• Active substances calculation

Inputs used: Molecular docking models QSAR membership models

Settings used

• Calculation Settings

3.5.2 AOP networks

Effects are related to each other using the toxicological concept of adverse outcome pathways (AOPs) and adverse outcome pathway networks (see https://aopwiki.org). Adverse Outcome Pathway (AOP) Networks specify how biological events (effects) can lead to an adverse outcome (AO) in a qualitative way through relations of upstream and downstream key events (KEs), starting from molecular initiating events (MIEs). Using AOPs, the adverse outcome (AO), e.g., liver steatosis, is linked to key events (KEs), e.g., triglyceride accumulation in the liver, and to molecular initiating events (MIEs), e.g., PPAR-alpha receptor antagonism. In general, multiple AOPs may lead to the same AO, and therefore AOP networks can be identified.

This module has as primary entities: Effects

Output of this module is used by: QSAR membership models Molecular docking models Active substances Relative potency factors Hazard characterisations Points of departure Effect representations

AOP networks from data

AOP Networks calculation

Find below a simplified AOP network based on a MCRA analysis for focal effect Steatosis-liver and related effects Fatty liver cells (tissue level), Cytoplasm displacement liver, Endoplasmatic reticulum stress liver, Mitochondrial disruption liver, Nucleus distortion liver and Triglyceride accumulation liver (cellular level).



Figure 3.71: AOP network liver steatosis

AOP networks data formats

AOP networks are described using two tables: the AOP networks table, and the effect relations table. The AOP networks table records the ids, names, descriptions, and other metadata of the AOP networks. The effect relations table describes the effects and effect relations (i.e., upstream and downstream key event relations) that are part of the AOP network.

Download empty dataset template: Zipped CSV Excel

AOP networks

Data format for specification of adverse outcome pathway (AOP) networks.

Name	Туре	Description	Aliases	Required
idAdverse- Outcome- Pathway- Network	AlphaNumeric (50)	Unique identification code of the AOP network.	idAOPN, idAOPNetwork, AOPN, AOPNetwork, Id	Yes
Name	AlphaNumeric (100)	Name of the AOP network.	Name	No
Description	AlphaNumeric (200)	Additional description or label of the AOP network.	Description	No
Reference	AlphaNumeric (200)	External reference(s) to sources containing more information about the AOP network. E.g., the AOP wiki, and the associated AOP wiki Ids.	Reference, References	No
idAdverse- Outcome	AlphaNumeric (50)	The identification code of the effect representing the adverse outcome of this AOP network.	idAdverse- Outcome, idAO, idEffect, Adverse- Outcome	Yes
RiskType	ExposureType	The exposure type of the adverse outcome. Acute or chronic.	RiskType	No

Accepted table names: AOPNetworks, AOPNetwork.

Effect relations

Dataformat for specification of the effect (key event) relationships of adverse outcome pathway (AOP) networks.

Name	Туре	Description	Aliases	Required
idAdverse- Outcome- Pathway- Network	AlphaNumeric (50)	Identification code of the AOP network for which this link is defined.	idAdverse- Outcome- Pathway- Network, idAOPN, idAOPNetwork, AOPN, AOPNetwork	Yes
idDownstream- KeyEvent	AlphaNumeric (50)	Identification code of the (triggered) effect of this relationship.	idDownstream- KeyEvent, idEffect, idKeyEvent, Effect, KeyEvent	Yes
idUpstream- KeyEvent	AlphaNumeric (50)	Identification code of the triggering effect of this relationship.	idTrigger, idUpstreamKey- Event, Trigger	Yes
Reference	AlphaNumeric (200)	External reference(s) to sources containing more information about the effect (key event) relationships.	Reference, References	No

Table 3.307: Table definition for Effect relations.

Accepted table names: EffectRelations, EffectRelation, EffectRelationships, EffectRelationship, KeyEventRelationship.

AOP networks settings

Selection settings

Name	Туре	Description
AOP Network Restrict AOP network by focal	AlphaNumeric Boolean	The AOP networks of interest. Restrict the AOP network to a specific sub-network, containing
upstream event		only the AOPs that include both the focal key event (KE) define here (which must be upstream of the AO) and the focal effect (adverse outcome, AO).
Focal upstream event	AlphaNumeric	The focal key event used for restricting the AOP network to a specific sub-network of interest.

Table 3.308: Selection settings for module AOP networks.

AOP networks as data

AOP networks can only be provided as data in the form of network definitions containing effect relations (key-event relationships) collections.

- AOP networks data formats
- AOP networks from data
- AOP networks calculation



Figure 3.72: AOP network

3.5.3 Burden of disease

Burden of disease data specify the burden of disease information related to specific effects and burden of disease indicators in specific populations.

This module has as primary entities: Populations Effects

Output of this module is used by: Environmental burden of disease

Burden of disease from data

Burden of disease data formats

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Burden of disease

Name	Туре	Description	Aliases	Required
Population	AlphaNumeric (50)	The population linked to this burden of disease.	Population	Yes
idEffect	AlphaNumeric (50)	Code of the effect linked to this burden of disease.	idEffect, EffectId, Effect, EffectCode	Yes
BodIndicator	BodIndicator	The burden of disease indicator linked to this burden of disease.	BodIndicator, Burdenof- DiseaseIndicator	Yes
Value	Numeric	The value of the burden of disease indicator.	Value	Yes

Table 3.307. Table definition for Durden of diseas	Table 3.309:	Table	definition	for	Burden	of	disease
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Accepted table names: BurdenOfDisease, BurdensOfDisease.

Burden of disease as data

Burden of disease definitions are provided as lists/catalogues of burden of disease information.

- Burden of disease data formats
- Burden of disease from data

3.5.4 Dose response data

Dose response data are data on response values of test systems at specified doses of substances (or mixtures of substances) from dose response experiments.

This module has as primary entities: Substances Test systems Responses

Output of this module is used by: Dose response models

Dose response data from data

Dose response data data formats

The meta-data of dose response experiments (such as name, description, etc.) are specified in the DoseResponseExperiments table.

For presenting the data of these experiments to the system, there are two formats: a single table format, see table *Dose* response data and a relational data format (three tables: *Dose response experiment doses, Experimental unit properties* and *Dose response experiment measurements*). Usually, the single table format will be the easier one. For internal use in MCRA, this single table data is converted to the relational data format.

Dose response data are used to extract assessment group membership or hazard doses. The meta-data of dose response experiments (such as name, description, etc.) are specified in the DoseResponseExperiments table. For presenting the data of these experiments to the system, there are two formats: a tabular format and a relational data format (three tables). Usually, the single table format will be the easier one. For internal use in MCRA, this single table data is converted to the relational data format.

Tabular dose response data format

In the tabular dose response data format, the substance doses and response measurements are provided in the same data table as columns and automatically parsed based on the specification of the substances and responses of the experiments in the dose response experiments table.

Download empty dataset template: Zipped CSV Excel

Dose response experiments

General information about the dose response experiments, such as the (unique) identifier, name, description, the used test-system, and the dose unit is stored in the table DoseResponseExperiments. If the data of an experiment is provided in a single table format, then the fields Time, Covariates, Substances, and Responses are used to map the column header names of the columns of the single data table to these their respective types.

Name	Туре	Description	Aliases	Required
idExperiment	AlphaNumeric (50)	Unique identification code of the dose effect experiment.	idExperiment, Id, Code	Yes
Name	AlphaNumeric (100)	Name of the dose effect experiment.	Name	No
Description	AlphaNumeric (200)	Description of the dose effect experiment.	Description	No
Date	DateTime	The starting date of the experiment.	Date	No
Reference	AlphaNumeric (200)	External reference, for instance, to the experiment protocol and/or supporting material.	Reference	No
Experimental- Unit	AlphaNumeric (100)	The name of the experimental unit of the experiment, e.g., rat, cage, litter, vial, cup, petridish.	Experimental- Unit	No
DoseRoute	AlphaNumeric (50)	For in-vivo test systems, the route in which the dose was administered	DoseRoute	No
Substances	AlphaNumeric	Code or comma separated list of the codes of the substances measured in the experiment. E.g., 'Cyproconazole, Thiram'. Required when presenting the dose-response data in a single table. Make sure that in table DoseResponseData the column headers exactly match these names.	idSubstance, SubstanceId, SubstanceCode, Substance, idSubstances, SubstanceIds, SubstanceCodes, Substances	Yes
DoseUnit	DoseUnit	Unit of the doses administered in this experiment.	DoseUnit	Yes
Responses	AlphaNumeric	Code or comma separated list of codes of the responses measured in the experiment. E.g., 'AngleM_PQ, Mortality'. Required when presenting the dose-response data in a single table. Make sure that in table DoseResponseData the column headers exactly match these names.	Responses, Response, idResponses, idResponse	Yes
Time	AlphaNumeric (100)	Identifier of the time field of the experiment. Required when presenting the dose-response data in a single table and responses are measured at multiple times. Make sure that in the table DoseResponseData the column header of the time-column exactly matches this name.	Time, Times	No
TimeUnit	TimeUnit	Unit of the time scale used in	TimeUnit	No
5. Hazard mod Covariates	ules AlphaNumeric (200)	the experiments. Comma separated list of the names/codes of the covariates of the experiment. E.g.	Covariates, Covariate	No

Table 3.310: Table definition for Dose response experiments.

Accepted table names: DoseResponseExperiments, DoseResponseExperiment.

Dose response data

Single (two-way) table data format for specifying data of dose response experiments (as alternative for the relational format). The column headers are dynamic and should be defined in the table DoseResponseExperiments through fields Substances and Responses (and, optionally, Covariates and Time). For responses given as aggregated statistics, also SD, CV, N and Uncertainty are specified as [Datatype:Response]. E.g., 'SD:Y', 'CV:Y', 'N:Y'. Uncertainty upper 95%limits are specified as 'UncertaintyUpper:Y'. For each quantal response an additional column 'N:[responsename]'is required with binomial totals (e.g. Mortality = 3, N:Mortality = 10).

Name	Туре	Description	Aliases	Required
idExperiment	AlphaNumeric (50)	Unique identification code of the dose effect experiment.	idExperiment, Experiment, Code	No
Experimental unit	AlphaNumeric (50)	Experimental unit numbers or identifiers. The column name of the experimental unit should be as specified in the dose response experiment record.	Experimental- Unit, Experimental- Units, Experimental unit	No
Substance(s)	AlphaNumeric (100)	One or more columns with doses for each substance, in the unit as specified in the dose response experiment table. The column name(s) should match the substance codes listed in the comma-separated list of the substances field of the dose response experiment record.		Yes
Response(s)	AlphaNumeric (100)	One or more columns with results for each response, in the unit(s) as specified in the dose response experiment table. The column name(s) should match the response codes listed in the comma-separated list of the responses field of the dose response experiment record.		Yes
Time	Numeric	The column containing the observed response times. The column name (header) should match that of the Time column in the dose response experiment record.		No
Covariate(s)	AlphaNumeric (100)	The column(s) containing additional properties of the experimental unit. The column name (header) should match the codes of the comma-separated covariates list in the dose response experiment record.		No

Table 3.311	Table defin	ition for D	Dose response	data.
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 $Accepted \ table \ names: \ Two Way Dose Response Data, \ Dose Response Data Two Way, \ Dose Response Data.$

Relational dose response data format

The relational dose response data format is the internal format of MCRA. In this format, the dose response experiments, doses, responses, and experimental units are provided in separate data tables.

Download empty dataset template: Zipped CSV Excel

Dose response experiments

General information about the dose response experiments, such as the (unique) identifier, name, description, the used test-system, and the dose unit is stored in the table DoseResponseExperiments. If the data of an experiment is provided in a single table format, then the fields Time, Covariates, Substances, and Responses are used to map the column header names of the columns of the single data table to these their respective types.

Name	Туре	Description	Aliases	Required
idExperiment	AlphaNumeric (50)	Unique identification code of the dose effect experiment.	idExperiment, Id, Code	Yes
Name	AlphaNumeric (100)	Name of the dose effect experiment.	Name	No
Description	AlphaNumeric (200)	Description of the dose effect experiment.	Description	No
Date	DateTime	The starting date of the experiment.	Date	No
Reference	AlphaNumeric (200)	External reference, for instance, to the experiment protocol and/or supporting material.	Reference	No
Experimental- Unit	AlphaNumeric (100)	The name of the experimental unit of the experiment, e.g., rat, cage, litter, vial, cup, petridish.	Experimental- Unit	No
DoseRoute	AlphaNumeric (50)	For in-vivo test systems, the route in which the dose was administered	DoseRoute	No
Substances	AlphaNumeric	Code or comma separated list of the codes of the substances measured in the experiment. E.g., 'Cyproconazole, Thiram'. Required when presenting the dose-response data in a single table. Make sure that in table DoseResponseData the column headers exactly match these names.	idSubstance, SubstanceId, SubstanceCode, Substance, idSubstances, SubstanceIds, SubstanceCodes, Substances	Yes
DoseUnit	DoseUnit	Unit of the doses administered in this experiment.	DoseUnit	Yes
Responses	AlphaNumeric	Code or comma separated list of codes of the responses measured in the experiment. E.g., 'AngleM_PQ, Mortality'. Required when presenting the dose-response data in a single table. Make sure that in table DoseResponseData the column headers exactly match these names.	Responses, Response, idResponses, idResponse	Yes
Time	AlphaNumeric (100)	Identifier of the time field of the experiment. Required when presenting the dose-response data in a single table and responses are measured at multiple times. Make sure that in the table DoseResponseData the column header of the time-column exactly matches this name.	Time, Times	No
TimeUnit	TimeUnit	Unit of the time scale used in the experiments	TimeUnit	No
5. Hazard mod Covariates	AlphaNumeric (200)	Comma separated list of the names/codes of the covariates of the experiment. E.g.	Covariates, Covariate	No 5

Table J.J12. Table definition for Dose response experiments.	Table 3.312:	Table definition	for Dose response	experiments.
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Accepted table names: DoseResponseExperiments, DoseResponseExperiment.

Dose response experiment doses

The table DoseResponseExperimentDoses describes the experiment design, being a complete specification of which doses of which substances were applied to which experimental unit and if relevant at what time.

Name	Туре	Description	Aliases	Required
idExperiment	AlphaNumeric (50)	Identification code of the experiment to which this design record belongs.	idExperiment, Experiment	Yes
idExperimental- Unit	AlphaNumeric (50)	Identification code of the experimental unit to which the dose is applied.	idExperimental- Unit, Experimental- Unit	Yes
Time	Numeric	The time of administration of the dose.	Time	No
idSubstance	AlphaNumeric (50)	Code of the substance that was administered.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
Dose	Numeric	The dose that was administered.	Dose	Yes

Table 3-3	13. Table	definition	for Dose	response e	xperiment	doses
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Accepted table names: DoseResponseExperimentDoses, DoseResponseExperimentDose.

Experimental unit properties

The table ExperimentalUnitProperties are used to specify additional properties of the experimental units of the experiment. For instance, the gender of the rat, in case rats are the experimental units.

Name	Туре	Description	Aliases	Required
idExperiment	AlphaNumeric (50)	Identification code of the experiment.	idExperiment, Experiment	Yes
idExperimental- Unit	AlphaNumeric (50)	Identification code of the experimental unit.	idExperimental- Unit, Experimental- Unit	Yes
PropertyName	AlphaNumeric (50)	Name of the experimental unit property.	Property, Name	Yes
Value	AlphaNumeric (100)	Value of the experimental unit property.	PropertyValue	No
OtherProperty		Other properties of experimental units are automatically parsed, using the column name (header) as property name.		No

Table 3.314: Table definition for Experimental unit properties.

Accepted table names: ExperimentalUnitProperties, ExperimentalUnitProperty.

Dose response experiment measurements

The table DoseResponseMeasurements describes the measurements that were done in the experiments. That is, for each response and experimental unit, at each observation time, one measurement should be recorded. If the response is an aggregated statistic, then this record may also include a standard deviation and number of units over which was aggregated.

Name	Туре	Description	Aliases	Required
idExperiment	AlphaNumeric (50)	Identification code of the experiment to which this measurement belongs.	idExperiment, Experiment	Yes
idExperimental- Unit	AlphaNumeric (50)	Identification code of the experimental unit from which the measurement is taken.	idExperimental- Unit, Experimental- Unit	Yes
idResponse	AlphaNumeric (50)	Identifier of the response that is measured.	idResponse, Response	Yes
Time	Numeric	Time of observation.	Time	No
ResponseValue	Numeric	The measured response.	ResponseValue, Value	Yes
SD:Response	Numeric	For aggregated responses, the standard deviation of the measurement.	SD:Response, ResponseSD	No
CV:Response	Numeric	For aggregated responses, the coefficient of variation (cv) of the measurement.	CV:Response, ResponseCV	No
N:Response	Numeric	For aggregated responses, the number of units over which was aggregated.	N:Response, ResponseN	No
Uncertainty- Upper:Response	Numeric	Optionally, measurement uncertainty quantification in terms of the upper value (i.e., an estimate of 95th percentile).	Uncertainty- Upper:Response, Response- Uncertainty- Upper, Uncertainty- Upper, Upper	No

Table 3.315: Table definition for Dose response experiment measurements.

 $\label{eq:constraint} Accepted \ table \ names: \ DoseResponseExperimentMeasurements, \ DoseResponseExperimentMeasurement, \ DoseResponseMeasurements, \ DoseResponseMeasurement.$

Dose response data settings

Selection settings

Table 3.316: Selection	settings for module	Dose response data.
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Name	Туре	Description
Merge dose response data of	Boolean	Specifies whether the dose response data of multiple experiment
multiple experiments		should be merged into one large dose response data set.

Dose response data as data

Dose response data are provided per experiment or study in which several responses (on in-vitro or in-vivo test systems) are measured from several administered substance doses.

- Dose response data data formats
- Dose response data from data

3.5.5 Dose response models

Dose response models are models fitted to dose response data and can be provided as data or calculated using a local or remote version of PROAST. The main results for hazard and risk assessment are benchmark doses (BMDs), related to a specified substance, response, optionally covariate value, and the benchmark response (BMR). Dose response models can be uploaded as data, retrieved from PROASTweb through *linked remote repositories*, or *calculated using an internal version of PROAST*.

This module has as primary entities: Test systems Responses Substances

Output of this module is used by: Hazard characterisations

Dose response models from data

Dose response models calculation

Dose response models are uploaded as data or retrieved from the PROASTweb through *linked remote repositories*. A second possibility is to compute dose response models using an integrated version of PROAST: for each response in a dose response experiment a dose response model is fitted. Depending on the type of data (e.g., response type, covariates y/n, single or multiple substances) a PROAST run is configured and executed. If *effect representations* are provided, then benchmark responses specified by the effect representations data are used, otherwise only the model fits will be computed without benchmark doses.

PROAST (copyright RIVM National Institute for Public Health and the Environment) is a software package for the statistical analysis of dose-response data. Its main purpose is dose-response modelling of toxicological data, and the derivation of a Benchmark dose (*BMD* or *BMDL*) in human risk assessment (or an ECx in ecotoxicological risk assessment). More generally, it can be used for (nonlinear) regression (with covariates) (see Slob (2002), Slob and Setzer (2013))

When PROAST is run without uncertainty, the *BMD* is estimated and the *BMDL* and *BMDU* are based on the profile likelihood method where the Fisher information matrix indicates the estimate's precision. When uncertainty is specified, the setting *Calculate a parametric benchmark dose confidence interval* becomes relevant.

- 1. checked: a benchmark dose interval (90% confidence) based on the profile likelihood method is estimated,
- 2. unchecked: the BMDL (p5) and BMDU (p95) are based on the 90% confidence interval of the bootstrapped BMD values.

In module *hazard characterisations* specify with setting *Use lower limit of BMD* whether a BMD or BMDL is used in the risk assessment.

Proast models

In Proast, a family of (nested) dose-response models are available that can be used for describing the change in any continuous endpoint as a function of dose. The likelihood ratio test is used to select one of the available models (model selection to prevent overparameterization).

- Model 1: y = a with a > 0
- Model 2: $y = a \cdot exp(x/b)$ with a > 0
- Model 3: $y = a \cdot exp(\pm (x/b)^d)$ with $a > 0, b > 0, d \ge 1$
- Model 4: y = a[c (c 1)exp(-x/b)] with a > 0, b > 0, c > 0
- Model 5: $y (c-1)exp(-(x/b)^d)$ with $a > 0, b > 0, c > 0, d \ge 1$



Figure 3.73: Dose response model: HepaRG Adipo72 72h.

where y is any continuous endpoint and x denotes the dose. In all models parameter a represents the level of the endpoint at dose 0, and b is considered as the parameter reflecting the efficacy of the substance or the sensitivity of the subject. At high doses model 4 and 5 level of to the value $a \cdot c$, so the parameter c can be interpreted as the maximum relative change. Model 3 and 5 have the flexibility to mimic threshold-like responses. All these model are nested to each other, except models 3 and 4, which both have three parameters.

In all models the parameter a is constrained to being positive for obvious reasons (it denotes the value of the endpoint at dose 0). The parameter d is constrained to values larger than (or equal to) 1, to prevent the slope of the function at dose 0 being infinite, which seems biologically implausible. The parameter b is constrained to be positive in all models. Parameter c in models 4 and 5 determines whether the function increases or decreases, by being larger or smaller than unity, respectively. To make model 3 a decreasing function a minus sign has to be inserted in the exponent (Slob (2002), Slob and Setzer (2013)).

Dose response models data formats

Dose response models are specified using three tables: the dose response models table holds the dose response model definitions (id, name, description) and other information about the dose response models. The dose response model benchmark doses table records the benchmark doses and (optionally) the model parameters for specific substances and covariates. The dose response model benchmark doses uncertainty table records results from bootstrap runs for the benchmark doses per substance/covariate combination.

Download empty dataset template: Zipped CSV Excel

Dose response models

Each dose response model has a unique id, a name (optional), and description (optional). Also, each dose response model is associated with a specific dose response experiment (idExperiment) from which the data used to create the model is obtained, a response (idResponse), one or more substances, and, optionally, specific covariates considered by the dose response model. The combination of the benchmark response type and the associated value define the benchmark response of the model. The dose unit specifies the unit used for the doses, and if applicable, the model equation can be specified.

Name	Туре	Description	Aliases	Required
idDose- ResponseModel	AlphaNumeric (50)	The unique identification code of the fitted dose response model.	idDose- ResponseModel, idModel	Yes
idExperiment	AlphaNumeric (50)	The identification code of the experiment from the dose response model.	experiment- Code, experimentId	Yes
Name	AlphaNumeric (100)	The name of the dose response model.	Name	No
Description	AlphaNumeric (200)	Description of the dose response model.	Description	No
Substances	AlphaNumeric	Code or comma separated list of the codes of the substances in the Dose Response Model. E.g., 'Cyproconazole, Thiram'.	Substances	Yes
idResponse	AlphaNumeric (50)	The response of the dose response model.	idResponse, Response	Yes
Covariates	AlphaNumeric	The covariates considered by the dose response model.	Covariates, Covariate	No
Benchmark- Response	Numeric	The value of the benchmark response or critical effect size.	Benchmark- Response, CriticalEffect- Size, CES	Yes
Benchmark- ResponseType	Benchmark- ResponseType	Specifies how the benchmark response is expressed. E.g., using a percent change in mean response or, for quantal response types, in terms of extra risk, additional risk, or ED50.	Benchmark- ResponseType, HazardEffect- SizeType, CriticalEffect- SizeType	No
LogLikelihood	Numeric	Loglikelihood of the model fit.	LogLikelihood	No
DoseUnit	AlphaNumeric (50)	The dose unit (if not specified, then mg/kg is assumed).	DoseUnit, UnitDose	No
ModelEquation	AlphaNumeric (500)	If available, the model equation of the dose response model (R model equation) or the identifier of the dose response model type.	ModelEquation, DoseResponse- ModelEquation, Equation	No

Table	3 317.	Table	definition	for D	ose resi	nonse	models
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Accepted table names: DoseResponseModels, DoseResponseModel.

Dose response model benchmark doses

The benchmark responses and benchmark doses belonging to the dose response models are recorded per substance/covariate in the dose response model benchmark doses table. Optionally, if the model equation of the dose response model has been specified in the dose response models table, the model parameter values for this specific substance/covariate can be specified here.

Name	Туре	Description	Aliases	Required
idDose- ResponseModel	AlphaNumeric (50)	The identification code of the dose response model to which this record belongs.	idDose- ResponseModel	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
Covariates	AlphaNumeric (500)	Comma separated list of the covariate values for which this benchmark dose applies.	Covariates, Covariate	No
Benchmark- Dose	Numeric	The (nominal) benchmark dose (BMD).	Benchmark- Dose, BMD, CED	Yes
Benchmark- DoseLower	Numeric	Benchmark dose lower uncertainty bound (BMDL).	Benchmark- DoseLower, BMDL, CEDL	No
Benchmark- DoseUpper	Numeric	Benchmark dose upper uncertainty bound (BMDU).	Benchmark- DoseUpper, BMDU, CEDU	No
Model- Parameter- Values	AlphaNumeric (500)	Parameter values for dose response models.	ParameterValues	No

Accepted table names: DoseResponseModelBenchmarkDoses.

Dose response model benchmark dose bootstraps

Empirical uncertainty values of the benchmark benchmark doses of dose response models can be recorded in the dose response model benchmark doses bootstraps table. The uncertainty set identifier (idUncertaintySet) can be specified to retain correlations between uncertainty records that originate from the same bootstrap run.

Name	Туре	Description	Aliases	Required
idDose- ResponseModel	AlphaNumeric (50)	The identification code of the dose response model to which this record belongs.	idDose- ResponseModel	Yes
idUncertainty- Set	AlphaNumeric (50)	The uncertainty set identifier.	idUncertainty- Set, UncertaintyId	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
Covariates	AlphaNumeric (500)	Comma separated list of the covariate values for which this benchmark dose applies.	Covariates	No
Benchmark- Dose	Numeric	Benchmark dose (BMD).	Benchmark- Dose, BMD, CED	Yes

Table 3.319: Table definition for Dose response model benchmark dose bootstraps.

Accepted table names: DoseResponseModelBenchmarkDosesBootstraps, DoseResponseModelBenchmarkDosesUncertain, DoseResponseModelBmdsUncertain.

Dose response models settings

Calculation settings

Table 3.320:	Calculation	settings for	module	Dose	response	models.
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Name	Туре	Description
Index substance	AlphaNumeric	The substance of interest or index substance.
Multiple substances analysis	Boolean	Specifies whether the assessment involves multiple substances.
Calculate a parametric benchmark dose confidence interval	Boolean	A parametric benchmark dose confidence interval (90%) is calculated by the profile likelihood method. If unchecked, the confidence interval is based on a bootstrap sample with uncertainty bounds p5 and p95.

Uncertainty settings

Table 3.321:	Uncertainty settings	for module Dose response models.
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Name	Туре	Description
Perform uncertainty analysis	Boolean	In probabilistic risk assessment of dietary exposure, distribution describe the variability in consumption within a given population of individuals and the variability of the occurrence and level of substances in the consumed foods. However, these calculations not consider the amount of uncertainty that is due to the limited size of the underlying datasets.
Resample hazard characterisations or RPFs	Boolean	Specifies whether to resample the hazard characterisations or relative potency factors. Requires hazard characterisation or RI uncertainty to be quantified in DoseResponseModelsUncertain RelativePotencyFactorsUncertain tables.
Iterations uncertainty analysis	Numeric	Specifies the number of uncertainty cycles (default 100).

Dose response models as data

Dose response models as data contain the details of fitted dose response models. The main elements for hazard and risk assessment are the benchmark doses (BMDs, BMDLs) related to specified substances, responses, and optionally covariate values for specified benchmark responses (BMR). These specifications can be provided in data files or can be retrieved/imported from PROAST output files on the PROAST website https://proastweb.rivm.nl/user/login using a PROASTweb user account and an application access key.

- Dose response models data formats
- Dose response models from data

Inputs used: Dose response data

Settings used

• Calculation Settings

Calculation of dose response models

Used as a calculator, dose response models are fitted to dose response data using an MCRA-internal version of PROAST. Currently, all available models appropriate for the response type will be fitted, and for the Hill and Exponential model families, the best fitting model based on maximum likelihood will be selected. The set of results for the calculation will include BMD(L)s etc. for all fitted models.

• Dose response models calculation

Inputs used: Effect representations

Settings used

• Calculation Settings

3.5.6 Effect representations

Effect representations specify the responses that can be used to measure specified effects and which response levels, the benchmark response (BMR), define the hazard limits for the effects.

This module has as primary entities: Effects Responses

Output of this module is used by: Hazard characterisations Dose response models

Effect representations from data

Effect representations data formats

Effect representations specify responses that may represent the effect.

Download empty dataset template: Zipped CSV Excel

Effect representations

One response can be set as the canonical response (golden standard). For a quantitative or stochastically qualitative canonical response a benchmark response should be defined.

Name	Туре	Description	Aliases	Required
idEffect idResponse	AlphaNumeric (50) AlphaNumeric (50)	Identifier of the effect Identifier of the response	idEffect idResponse	Yes Yes
Benchmark- Response	Numeric	The threshold response value that defines a hazard. For numeric responses (Continuous, Quantal, Count) the value that defines a hazard. For Binary responses 1 defines a hazard by default, unless redefined here.	BenchMark- Response, HazardEffect- Size, BMR, CriticalEffect- Size, CES	No
Benchmark- ResponseType	Benchmark- ResponseType	Specifies how the BenchMarkResponse is expressed, relative to the response at zero dose, or absolute. Required for numeric response types (Continuous, Quantal, Count). For qualitative responses (Ordinal, Categorical) Absolute is used.	Benchmark- ResponseType, HazardEffect- SizeType, CriticalEffect- SizeType	No

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Accepted table names: EffectRepresentations, EffectRepresentation.

Effect representations

Calculation settings

Name	Туре	Description
Multiple effects analysis	Boolean	Specifies whether the analysis should consider multiple effects. Otherwise, a single focal effect should be selected.
Include related effects of AOP network	Boolean	Include all related key events of the AOP network.

Table 3.323:	Calculation set	tings for mod	ule Effect represe	entations.
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Effect representations as data

Effect representations are provided as data in the form of specified combinations of effect and response, optionally with a benchmark response that defines a hazard limit for the effect.

- Effect representations data formats
- Effect representations from data

Inputs used: AOP networks

Settings used

• Calculation Settings

3.5.7 Exposure response functions

Exposure response functions define the relation between percentiles and percentile specific ratios for exposure effect related to environmental burden of disease.

This module has as primary entities: Substances Effects

Output of this module is used by: Environmental burden of disease

Exposure response functions from data Exposure response functions data formats

Download empty dataset template: Zipped CSV Excel

Exposure response functions

Name	Туре	Description	Aliases	Required
idModel	AlphaNumeric (50)	The unique identifier of the exposure response model.	idModel, Model, ModelId	Yes
Name	AlphaNumeric (100)	Name of the exposure response model.	Name	No
Description	AlphaNumeric (200)	Additional description or label of the exposure response model.	Description	No
idSubstance	AlphaNumeric (50)	The substance code.	idSubstance, Substance, SubstanceId	Yes
idEffect	AlphaNumeric (50)	Code of the effect linked to this exposure response function.	idEffect, EffectId, Effect, EffectCode	Yes
TargetLevel	TargetLevelType	The target level. I.e., internal or external. If omitted, external is assumed.	TargetLevel, LevelTarget	No
ExposureRoute	ExposureRoute	The exposure route(s) (only applicable if exposure level is external) for which this exposure response function is defined.	ExposureRoute, Route	No
Biological matrix	<i>BiologicalMatrix</i>	The biological matrix (e.g., blood, urine) for which this exposure response function is defined.	Biological- Matrix, MatrixSource, SourceMatrix	No
DoseUnit	DoseUnit	The unit of the result of the exposure response function.	DoseUnit, Unit	Yes
ExpressionType	ExpressionType	The expression type or adjustment method of the unit. This field specifies how the unit (target) is adjusted, e.g. for blood lipids for fat soluble biomarkers ('mg/g lipids') or the dilution level of the urine ('mg/g creatinine').	ExpressionType, Adjustment- Method	No
Population- Characteristic	Population- CharacteristicType	Population characteristic type.	Characteristic, idCharacteristic, Population- Characteristic, IdPopulation- Characteristic	No
Effect- ThresholdLower	Numeric (50)	Lower threshold for the effect for the specified population characteristic.	EffectThreshold- Lower	No
Effect- ThresholdUpper	Numeric (50)	Upper threshold for the effect for the specified population characteristic.	EffectThreshold- Upper	No
Effect metric	EffectMetric	The type of risk for which this exposure response function is defined.	EffectMetric, ResponseMetric, Metric	Yes
Exposure- ResponseType	ExposureResponse- Type	The type of specification used to define the relation between the exposure and the exposure response.	Exposure- ResponseType, ErType, ErfType	Yes
2 Exposure- Response- Specification	AlphaNumeric (200)	The specification for the exposure response. Either a mathematical expression when the exposure response type is function, or a single	Exposure- Response- Specification, ErfSpecification, ErSpecification	r 3. Modul e

Table 3.324: Table definition for Exposure response functions.
$\label{eq:constraint} Accepted \ table \ names: \ Exposure Response Functions, \ Response Functions, \ Exposure Effect Functions.$

Exposure response function subgroups

Exposure response functions defined per exposure category.

Name	Туре	Description	Aliases	Required
idModel	AlphaNumeric (50)	The unique identifier of the exposure response model.	idModel	Yes
idSubgroup	AlphaNumeric (50)	The unique identifier of the subgroup.	idSubgroup	Yes
ExposureUpper	Numeric (50)	Specifies the upper bound of the exposure percentile of the exposure response function subgroup. Exposure belong to a subgroup when the exposure is equal or lower than the specified upper bound and larger than the specified upper bound of the previous subgroup.	ExposureUpper, UpperExposure	No
Exposure- Response- Specification	AlphaNumeric (200)	The specification for the exposure response. Either a mathematical expression when the exposure response type is function, or a single value if the exposure response type is constant, perDoubling, or perUnit.	ErfSpecification, Exposure- Response- Specification, Exposure- Response- Function- Specification	Yes
Exposure- Response- Specification- Lower	AlphaNumeric (200)	Exposure response lower uncertainty bound. Either a mathematical expression when the exposure response type is function, or a single value if the exposure response type is constant, perDoubling, or perUnit.	Exposure- Response- Specification- Lower, ErfLower	No
Exposure- Response- Specification- Upper	AlphaNumeric (200)	Exposure response upper uncertainty bound. Either a mathematical expression when the exposure response type is function, or a single value if the exposure response type is constant, perDoubling, or perUnit.	Exposure- Response- Specification- Upper, ErfUpper	No

Accepted table names: ErfSubgroups, ErfSubgroup.

Exposure response functions

Uncertainty settings

Name	Туре	Description
Resample exposure response function values	Boolean	Specifies whether exposure response function values are resampled.

Table 3.326: Uncertainty settings for module Exposure response functions.

Exposure response functions as data

Exposure response function definitions are provided as lists/catalogues of exposure response function definitions.

- *Exposure response functions data formats*
- Exposure response functions from data

3.5.8 Hazard characterisations

Hazard characterisations are reference exposure values for active substances at the chosen biological target level (external or internal). Hazard characterisations may be specified for specific effects or for the critical effect as defined in hazard characterisation. Hazard characterisations are specified as external values (e.g. human based guidance values, such as ADI or ARfD) or are based on points of departure, such as BMD(L)s from dose-response models or externally specified points of departure (NOAEL, LOAEL, MDS). The computation may involve assessment factors, e.g. for inter-species conversion, intra-species variation or additional sources of uncertainty. The calculation may also use kinetic conversion factors, PBK models or absorption factors to convert external doses to internal concentration/doses or vice versa.

This module has as primary entities: Substances Effects Populations

Output of this module is used by: Active substances Relative potency factors Risks Single value risks

Hazard characterisations from data

In table *Hazard characterisations subgroups*, hazard characterisations are specified that are dependent on individual properties like age and/or *gender*.

For age, specify the lower bound of the age interval (in years) of the hazard characterisation subgroup. Individuals belong to a subgroup when the age of the individual is equal or greater than the specified lower bound and smaller than the specified lower age of the next subgroup.

Check option *Use hazard characterisations subgroup* (default) to use age and/or gender specific hazard characterisations (only visible when the datasource contains age/gender specific hazard characterisations information).

Hazard characterisations calculation

Hazard characterisations are defined as deterministic threshold values, e.g., *ADI*, *ARfD*, or as distributions (using probabilistic models). They are linked to an effect of interest or alternatively are defined for the critical effect. Hazard characterisations depend on the *exposure type* (acute or chronic) and the *target level* of the human body (external via some route of exposure or internal for a specific defined organ or biological matrix). Hazard characterisations are derived from *points of departure* provided as data and/or from *dose-response models*. The procedure for computing hazard characterisations has two main phases: 1) collection of all available hazard characterisations and imputation of missing hazard characterisations.

Collection of available hazard characterisation candidates involves collecting the appropriate points of departure data and/or dose-response models that are used for deriving the hazard characterisations. In MCRA, a distinction is made between three *methods for computing hazard characterisations*:

1. Calculation of hazard characterisations from *in-vivo* points of departure (*PoD*, e.g. *BMD*, *BMDL*, *NOAEL*, *LOAEL*).

- 2. Calculation of hazard characterisations from PoDs (in this case BMD) calculated from dose response data.
- 3. Calculation of hazard characterisations based on an *in-vivo PoD for the index substance and in-vitro RPFs from dose-response models for the other substances (IVIVE model)* (cumulative assessments only).

For all three methods, the collected points of departure and benchmark doses should be aligned with the target system. This alignment may involve various conversion steps for each point of departure and specific substance, and can be formally specified as:

$$\textit{HC} = f_{\texttt{expression-type}} \cdot f_{\texttt{kinetic}} \cdot \frac{1}{f_{\texttt{inter-species}}} \cdot \frac{1}{f_{\texttt{intra-species}}} \cdot \frac{1}{f_{\texttt{additional}}} \cdot \textit{PoD}$$

where:

- *HC* denotes the hazard characterisation.
- $f_{\text{expression-type}}$ denotes the *expression type correction factor*, e.g., for extrapolation from LOAEL or NOAEL, or from NOAEL to BMD.
- *f*_{kinetic} denotes the kinetic conversion factor for *conversion from internal to external or external to internal hazard characterisations*.
- $f_{\text{inter-species}}$ denotes the inter-species factor for *extrapolation from animal to human (inter-species)*.
- $f_{intra-species}$ denotes the intra-species factor for *extrapolation from the average to the sensitive human or* probabilistic calculation of the distribution of human individuals (intra-species).
- *f*_{additional} denotes the additional assessment factor for *extrapolation from the POD to the hazard characterisation in humans for sources where appropriate data or information is scarce or missing (additional).*
- *PoD* denotes the point of departure.

Note that inter- and intra-species extrapolation and the use of an *additional assessment factor* are optional. However, expression type correction and the kinetic conversion are always applied (when relevant) whatever option is chosen.

Occasionally, for some substances multiple hazard characterisations are available (e.g., obtained from multiple experiments) and for others substance hazard characterisations are still missing. Hence, two final steps remain to come to the final set of hazard characterisation:

- Aggregation over multiple available hazard characterisations. Set the selection method in case of multiple candidate hazard characterisations from MostToxic to Aggregate.
- Imputation of missing hazard characterisations. Check the option Imput missing hazard characterisations, to select the Imputation method.

Specify with setting *Use lower limit of BMD* whether a BMD or BMDL is used in the risk assessment. The BMDL is a parametric estimate based on the profile likelihood method or is the p5 estimate of the bootstrapped BMD values in an uncertainty run. See module *dose response models* for more information on the estimation of the BMDL.

In above figures the risk characterisation ratio (exposure/hazard) based on a BMDL is conservative compared to the ratio based on the BMD.

Hazard characterisation type extrapolation

Hazard doses, or points of departure can be of *various types*. E.g., *BMD* s, *NOAEL* s, or *LOAEL* s. When computing hazard characterisations, the type in which the hazard characterisations are expressed (i.e., the *hazard characterisation expression type*) should be specified explicitly. When points of departure from types different from the expression type are provided, these should be translated to the specified expression level. In the current implementation, the simple conversion factors shown in Table 3.327 are used, roughly based on the WHO guidance document on evaluating and expressing uncertainty in hazard characterisation, see WHO (2018).



Figure 3.74: Risk characterisation ratio (exposure/hazard) based on BMD, POCE = 0.18%.



Figure 3.75: Risk characterisation ratio (exposure/hazard) based on BMDL, *POCE* = 32%.

From	То	Conversion factor
BMD	NOAEL	1/3
BMD	LOAEL	1
NOAEL	BMD	3
NOAEL	LOAEL	1/3
LOAEL	BMD	1
LOAEL	NOAEL	1/3

Table 3.327: Conversion factors for hazard characterisation types.

Inter-species extrapolation

Hazard doses, or points of departure, are commonly only determined for animals, not for humans. In order to derive hazard characterisations for humans, the animal hazard doses need to be converted to toxicologically equivalent doses for humans. This extrapolation is usually expressed as a multiplication factor, and traditionally a factor of 10 is used (which is roughly obtained from the product of a factor of 3.2 for toxicokinetic variability and a factor 3.2 for toxicodynamic variability).

The following methods are available within MCRA:

- 1. No inter-species extrapolation: Assume that for all available points of departure, the animal hazard dose is equal to the human hazard dose. Effectively, this is equivalent to using a conversion factor of 1.
- 2. Default distribution: Use a conversion factor drawn from a default, substance and species independent log-normal uncertainty distribution specified (as *model settings*) by a geometric mean (GM) and geometric standard deviation (GSD). In the *nominal run*, the nominal value of this distribution (i.e., the geometric mean) is used as a conversion factor. In the *uncertainty analysis loop*, provided that inter-species extrapolation uncertainty is *included in the uncertainty analysis*, a single factor is drawn from the lognormal distribution.
- 3. Substance/species specific distributions: Use conversion factors drawn from substance/species specific lognormal uncertainty distributions specified (as *data*) by a geometric mean (GM) and geometric standard deviation (GSD). In the *nominal run*, a factor equal to the geometric mean is used for all combinations of substance and species. In the *uncertainty analysis loop*, provided that inter-species extrapolation uncertainty is *included in the uncertainty analysis*, a uncertainty factor is drawn from the lognormal distribution with $\mu = 0$ and $\sigma^2 = 1$, which is used to obtain correlated draws for all available inter-species conversion factor distributions. If the distribution parameters are missing for a specific substance/species, then the default distribution is used as a fallback.

Intra-species extrapolation of hazard characterisations

There is variation between individuals concerning their individual sensitivities to experience health effects. In some scenarios the aim is to perform assessments for the sensitive individuals instead of the average individuals for which the points of departure are derived. If this is the case, then extrapolation is required to translate hazard characterisations derived for the average individual to hazard characterisations for a sensitive individual. In traditional exposure assessments, a safety of 100 is commonly used as a margin of safety, that is assumed to be composed of a interspecies extrapolation factor (factor 10), and inter-individual extrapolation factor (factor 10). I.e., the hazard characterisation defined for the sensitive individual is defined as

$$HC_{\rm sens} = \frac{1}{f_{\rm intra-species}} \cdot HC_{\rm avg}$$

Here $f_{inter-species}$ denotes the intra-species factor. An alternative to using a fixed safety factor, is to define intraspecies variability explicitly using *a lognormal distribution*, characterised by a geometric mean (GM) equal to 1 and a geometric standard deviation (GSD). For *risks calculations*, this distribution could be used to sample individual hazard characterisations. This effectively converts the description of hazard characterisations to include variability, with an unbiased central value.

Additional assessment factors

In cases where where appropriate data or information is scarce or missing, an additional assessment factor (AF) is used for extrapolation from the *POD* to the hazard characterisation in humans. This factor can be used as a worst case value (preferably AF = 100) for inter- and intra-species extrapolation, but it may equally well serve as an additional extrapolation factor next to inter- and/or intra-species extrapolation. In the latter case the factor merely serves to account for differences in for example route to route, metabolic rates in interspecies, completeness and consistency of available data, reliability of alternative data (e.g. read-across) or quality of data in general.

In-vitro in-vivo extrapolation (IVIVE)

The in-vitro in-vivo extrapolation method implemented in MCRA is based on the following prerequisites:

- 1. For one substance, the index substance, a reliable point of departure is available for the adverse outcome of interest obtained from an *in-vivo* assay (i.e., external dose).
- 2. There are other substances for which there is a dose-response model available from an *in-vitro* assay on a response representing an early key event of the adverse outcome for these substances and the index substance.

In *IVIVE*, these RPFs, in combination with the known hazard characterisation of the index substance, can be used to derive hazard characterisations for the other substances as well. Figure 3.76 shows the conceptual model that forms the basis of the IVIVE methodology of MCRA.



KM = Kinetic Model (animal or human) MOE = Margin Of Exposure

Figure 3.76: Conceptual model IVIVE.

IVIVE for calculating internal hazard characterisations

- 1. Translate the (external) PoD of the index substance substance to an internal hazard characterisation for the human target system/*biological matrix*.
- 2. If the RPFs are obtained are obtained using mol-based specification of the doses, then convert the mol-based RPFs to mass-based RPFs. I.e.,

$$RPF_{mass-based,i} = RPF_{mol-based,i} \cdot \frac{MW_{ref}}{MW_i}$$

3. Derive internal hazard characterisations for the human target system for the other substances by multiplying the RPF obtained from dose-response modelling with the hazard characterisation of the index substance. I.e.,

$$HC_i = HC_{ref} \cdot RPF_{mass-based,i}$$

IVIVE for calculating external hazard characterisations

- 1. Translate the PoD of the index substance to an external human hazard characterisation (dietary/oral exposure route).
- 2. Derive an internal hazard characterisation for the index substance, with a target biological matrix representative for the response of the dose-response model.
- 3. If the RPFs are obtained are obtained using mol-based specification of the doses, then convert the mol-based RPFs to mass-based RPFs.
- 4. Derive internal hazard characterisations for the human target system for the other substances by multiplying the RPF obtained from dose-response modelling with the hazard characterisation of the index substance.
- Convert the internal hazard characterisations of the other substance to external hazard characterisations for the dietary/oral exposure route using.

Absorption or kinetic conversion of hazard characterisations

When the *hazard characterisation level* is internal and points of departure are available for external exposures (e.g., NOAELs from in-vivo animal studies) or when the hazard characterisation level is external and benchmark doses are available at the internal level, then *kinetic conversion factors* or *PBK models* are needed to translate the external doses to *equivalent internal doses at the target* or *biological matrix* of interest or *vice-versa*.

In MCRA, this alignment from internal to external or from external to internal is generally termed *conversion*, associated with a *kinetic conversion factor* or *PBK model*. The kinetic conversion factor is a multiplication factor needed to obtain a hazard characterisation on the target level from a hazard characterisation of the point of departure or benchmark dose. Depending on the chosen kinetic modelling tier, this kinetic conversion factor may be 1) assumed to be one, or derived from 2) absorption factors, 3) kinetic conversion factors or 4) using *PBK* models.

An important detail in the use of kinetic conversion factors for computing hazard characterisations is the order between kinetic conversion and inter-species extrapolation. Notice that when points of departure are determined for animals, a choice should be made regarding the order of inter-species extrapolation and kinetic modelling. That is, one may first choose to convert animal external point of departure to an internal hazard characterisation for that animal, using the available animal kinetic model. Alternatively, one may first extrapolate the animal external point of departure to a human external hazard characterisation, and thereafter apply the human kinetic model to obtain internal hazard characterisations. Currently, in MCRA only the latter approach is implemented.

Extrapolation from external to internal hazard characterisations

The calculation of internal hazard characterisations based on external hazard characterisations is similar to the procedure for *computing internal exposures*. In the simplest tier, equivalence can be assumed between internal and external hazard characterisations, and in higher tiers absorption factors, respectively PBK models can be used.

Calculation of internal doses using absorption factors

In the simplest form, internal doses are obtained from external exposure concentrations using absorption factors that are specified per combination of substance and route. That is, for a given substance, the internal hazard characterisation $HC_{int.dose}$ can be derived from an external hazard characterisation HC_{ext} as

$$HC_{\text{int,dose}} = f_{\text{abs},r} \cdot HC_{\text{ext},r}$$

Here, *r* denotes the route of the external exposure HC_{ext} , and $f_{abs,r}$ denotes the absorption factor of route *r*. Note that this model assumes that the external hazard characterisations are specified as concentrations (i.e., substance amount divided by the body weight).

Calculation of internal concentration using human PBK models

A more detailed alternative to absorption factors is to use kinetic conversion factors or one of the *advanced PBK models* available in MCRA. In the last approach, for each substance independently, an external exposure equivalent to the dose of the external hazard characterisation is presented to a representative simulated individual for a number of simulated days to the PBK model of the individual. This representative individual should represent the "average" individual of the population, with nominal physiological properties (e.g., an average bodyweight of 70kg). This yields a time course of the internal substance amount at the specified target or biological matrix from which a long term average substance amount (chronic) or peak substance amount (acute) can be obtained. By dividing this substance amount by the weight of the biological matrix, an internal concentration is obtained, which then represents the internal hazard characterisation.

More details on computing internal concentrations from external doses can be found in the description of the calculation of internal concentrations from external exposures in section *kinetic conversion factors* or *PBK models*. For both tasks, the procedure for computing internal exposures/doses is exactly and the same *kinetic model settings*, such as *dosing patterns* and *non-stationary period* period apply for calculation of internal hazard characterisations as well.

Calculation of internal doses using animal PBK models

In the above methods, the assumption is that the external points of departure (often obtained from experiments on animals) are first converted to external hazard characterisations for humans, and a human kinetic model is used for obtaining the internal hazard characterisations. As mentioned, an alternative approach is to use first the animal PBK models to derive an internal hazard characterisation specific for the tested animal species and thereafter extrapolate to humans. When there are more precise kinetic models available for the animal used in the experiments for obtaining the point of departure, this could be a preferred path.

1 Note

Notice that this procedure is not yet implemented.

Extrapolation from internal to external hazard characterisations

In some cases, hazard characterisations are available at the internal level whereas the specified *hazard characerisation level* is external. This situation may occur, for instance, in *in-vitro in-vivo extrapolation (IVIVE)*. Then, conversion is needed from the internal level to the external level, where the external level is implicitly defined as coming from the dietary/oral route of exposure.

For absorption factors, the external (dietary) hazard characterisation of a substance is simply computed by dividing the internal hazard characterisation by the dietary absorption factor, i.e.,

$$HC_{\rm ext,diet} = rac{HC_{\rm int,dose}}{f_{\rm abs,diet}}$$

For PBK models, reverse dosimetry is needed to find the corresponding external (dietary) dose that yields the internal concentration specified by the internal hazard characterisation. In MCRA, this is done using a bisection algorithm: external doses are systematically fed into the PBK model and convergence is reached when the resulting dose is within a specified precision of the value derived from the hazard characterisation.

Hazard characterisation imputation

Occasionally, for some substances causing the effect of interest, no data are available to determine the hazard characterisations, HCs, (e.g. no points of departure or dose response models). Instead of excluding these substances in quantitative analyses, missing HCs are imputed based on known HCs of similar substances. The imputed HCs are used for calculating relative potency factors or used in a risk assessment.

Munro P5 (TTC approach)

The Threshold of Toxicological Concern (TTC) is an example of a tier for extrapolation of hazard characterisations from other substances that is already in common use (see Munro et al. (1996)). The *Munro collection of NOELs/LOAELs* is a collection of *NOEL* s/*LOAEL* s for chemicals for the critical (i.e., first occurring) effect. In the TTC approach, the toxicity of an unknown substance is, depending on its Cramer class (see Cramer et al. (1976)), imputed by the 5th percentile NOAEL of the sub-collection of the corresponding Cramer class.

Two variations of this approach are to use the empirical NOAEL distribution itself (just sample from the NOAEL data), or to fit a distribution (e.g. lognormal) to the empirical data and sample from the parametric distribution. MCRA provides an implementation of the TTC approach that uses the empirical distribution. In the nominal run, this implementation imputes the hazard characterisations with a value equivalent to the TTC. In the uncertainty runs, NOAELs are sampled from the empirical distribution.

The TTC is a conservative estimate of the NOAEL for at least two reasons:

- 1. TTCs are calculated from a collection of NOELs for the critical (i.e., first occurring) effect within each study and often the effect of interest will not be the critical effect, and therefore higher NOAELs are expected.
- 2. The TTC is a low percentile and therefore a conservative estimate for a random class member with unknown NOAEL.

Munro central value

To avoid the conservatism of taking the 5th percentile in the Munro P5 approach, alternatively, a nominal (or central) value could be taken from the Munro collection for each Cramer class. For a nominal run without uncertainty, the expected contribution of a substance with missing hazard characterisation to the risk as quantified in the hazard index is obtained from

$$HI = SF \cdot \sum_{i}^{n} \frac{\exp_{i}}{HC_{i}}$$

Here SF are all combined safety factors. It follows from this equation that an unbiased estimate for the contribution from a substance with missing hazard characterisations is obtained by taking the harmonic mean from the available NOAELs:

$$\textit{NOAEL} = \left(\sum_{i=1}^n \frac{1}{\textit{NOAEL}_i}\right)^{-1}$$

This is the value to use in a nominal run without uncertainty for the Munro central value approach. For the uncertainty runs, this approach also uses random sampling from the empirical distribution of the corresponding Cramer class.

Available hazard characterisations distribution P5

Another conservative aspect of the TTC approach is that the Munro set lists NOELs/LOAELs for critical effects, not for the specific effect under study. Therefore an alternative is to use the effect-specific hazard characterisations of the substances for which these are available. This collection will have on average higher NOAELs than those of the Munro NOEL collection, because for many substances, the effect of interest will not be the critical effect.

Available hazard characterisations distribution central value

Similar to the Munro central value approach, a central value could also be obtained from the set of effect-specific hazard characterisations distribution for imputation of hazard characterisations. This approach may yield the most realistic, or unbiased imputation value for missing hazard characterisations.

Aggregation over multiple available hazard characterisations

In some scenarios, for a given substance and effect, multiple hazard characterisations are available e.g. different studies yield different *NOAEL* s. In such cases, a single hazard characterisation should be derived from the available candidates.

In a conservative approach, choose the lowest hazard characterisation (HC) of the available hazard characterisations. I.e.,

$$\mathrm{HC} = \min_{i=1,\dots,n} \mathrm{HC}_i$$

Alternatively, the candidates are aggregated into a new "average" hazard characterisation. For this, the harmonic mean, also used for obtaining central value estimates in the *imputation of missing hazard characterisations*, is a suitable approach.

$$\mathrm{HC} = \left(\sum_{i}^{n} \frac{1}{\mathrm{HC}_{i}}\right)^{-1}$$

Hazard characterisations data formats

Hazard characterisations provide reference threshold values associated with the hazard of interest. Examples are health-based guidance values such as ADI or ARfD, and points of departure such as BMD or NOAEL. Hazard characterisations may be dependent on e.g. age (or gender not implemented yet)

Download empty dataset template: Zipped CSV Excel

Hazard characterisations

Hazard characterisations are specified for combinations of hazard characterisation type, effect, substance, population type, target level, and exposure route (for external) or target organ (for internal). Effects can be specific, but can also be labelled as being the critical effect and used as such if this has been specified in the hazard characterisation settings.

Name	Туре	Description	Aliases	Required
idHazard- Characterisation	AlphaNumeric (50)	Id of the hazard characterisation.	id, idHazard- Characterisation	Yes
idEffect	AlphaNumeric (50)	Code of the (critical) effect linked to this hazard characterisation.	idEffect, EffectId, Effect, EffectCode	No
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceCode, SubstanceId, Substance	Yes
idPopulation- Type	AlphaNumeric (50)	The code of the population type for which this reference value is defined. If not specified, PS06A, Consumers is assumed.	idPopulation- Type, PopulationType, Population- Group, Population- Subgroup	No
TargetLevel	TargetLevelType	The target level. I.e., internal or external. If omitted, external is assumed	TargetLevel	No
ExposureRoute	ExposureRoute	The exposure route (only applicable if target level is external). If not specified, Dietary is assumed.	ExposureRoute	No
Matrix	BiologicalMatrix	The biological matrix or organ (should be specified when target level is internal).	Matrix, BiologicalMatrix	No
Expression type	ExpressionType	The expression type for (internal) hazard characterisations of which the concentrations are not directly specified at the level of the biological matrix. For instance, urinary expressions expressed in terms of the amount excreted per gram of creatinine or blood concentration expressed per gram of lipids.	ExpressionType	No
IsCriticalEffect	Boolean	Specifies whether this value is IsCriticalEffect the value associated with the critical effect. If omitted, No is assumed		No
ExposureType	ExposureType	The exposure type associated with the hazard characterisation (i.e., chronic or acute).	ExposureType	Yes
Hazard- Characterisation- Type	Hazard- Characterisation- Type	The type of the hazard characterisation (e.g., ARfD, ADI, NOAEL, BMD).	Hazard- Characterisation- Type	Yes
Qualifier	ValueQualifier	Qualifier of the hazard characterisation value, e.g. equal-to (=) or smaller-than (<). If omitted, = is assumed.	QualifierType	No
Value	Numeric	Reference value that characterises the hazard.	Value, Hazard- Characterisation- Value	Yes
5. Hazard modu	lles ^{seUnit}	Unit of the hazard characterisation value.	DoseUnit, Unit	Yes 5:
idPointOf- Departure	AlphaNumeric (50)	ric (50) The code of the point of idHazardDose, idPod		No

Table 3.328: Table definition for Hazard characterisations.

Accepted table names: HazardCharacterisations.

Hazard characterisations uncertainty

Often, the hazard characterisations found for a substance are uncertain. This table facilitates in specifying the hazard characterisation uncertainty in the form of a set of uncertainty values that may additionally be specified for a hazard characterisation model - substance combination.

Name	Туре	Description	Aliases	Required
idHazard- Characterisation	AlphaNumeric (50)	The hazard characterisation model code (must correspond to values in id column of Hazard characterisations table).	idHazard- Characterisation	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
Hazard- Characterisation- Value	Numeric	Reference value that characterises the hazard.	Hazard- Characterisation, Hazard- Characterisation- Value	Yes

Accepted table names: HazardCharacterisationsUncertain, HazardCharacterisationsUncertai, HazardDoseUncertain, HCUncertain.

Hazard characterisations subgroups

Hazard characterisations are dependent on e.g. age.

Name	Туре	Description	Aliases	Required
idSubgroup	AlphaNumeric (50)	The id of the subgroup.	idSubgroup	Yes
idHazard- Characterisation	AlphaNumeric (50)	Id of the hazard characterisation.	idHazard- Characterisation	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
AgeLower	Numeric (50)	Specifies the lower bound of the age interval (in years) of the hazard characterisation subgroup. Individuals belong to a subgroup when the age of the individual is equal or greater than the specified lower bound and smaller than the specified lower age of the next subgroup.	AgeLower, LowerAge	No
Gender	GenderType	The gender of the subgroup.	Gender	No
Value	AlphaNumeric (50)	The hazard characterisation of the subgroup.	Value	Yes

Table 3.330: Table definition for Hazard characterisations subgroups.

Accepted table names: HCSubgroups, HCSubgroup.

Hazard characterisations subgroups uncertainty sets

Hazard characterisations are dependent on e.g. age and may be uncertain.

Name	Туре	Description	Aliases	Required
idSubgroup	AlphaNumeric (50)	The id of the subgroup.	idSubgroup	Yes
idHazard- Characterisation	AlphaNumeric (50)	Id of the hazard characterisation.	idHazard- Characterisation	Yes
idUncertainty- Set	AlphaNumeric (50)	The id of the ucertainty set.	idUncertainty- Set, idUncertainty- Set, idUncertainty	Yes
Value	AlphaNumeric (50)	The uncertain hazard characterisation of the subgroup.	Value	Yes

Table 3.331:	Table	definition	for	Hazard	characterisations	subgroups	un-
certainty sets.							

Accepted table names: HCSubgroupsUncertain, HCSubgroupUncertain.

Hazard characterisations settings

Selection settings

Name	Туре	Description
Exposure type	ExposureType	The type of exposure considered in the assessment; acute (shor term) or chronic (long-term).
Multiple effects analysis	Boolean	Specifies whether the analysis should consider multiple effects. Otherwise, a single focal effect should be selected.
Target level	TargetLevelType	Select whether the assessment is at the level of external dose, internal (target) concentration or internal (systemic) dose. For a aggregate assessment, combining dietary and nondietary exposures, the target level is always internal concentration or internal (systemic) dose. When only dietary exposures are available, the target level is optional and may be an external (administered) dose, an internal (tissue) concentration or an internal (absorbed) dose.
Consider critical effect	Boolean	Specifies whether the analysis should look at critical effects such as specified in the Hazard characterisation data source.
Restrict active substances to substances with available hazard characterisations	Boolean	Restrict assessment group membership based on presence/abser of hazard characterisations.
Include related effects of AOP network	Boolean	Include all related key events of the AOP network.

Table 3.332: Selection settings for module Hazard characterisations.

Calculation settings

Name	Туре	Description
Selected tier	SettingsTemplateType	Specifies all module settings should be set according to a
		pre-defined tier or using custom settings.
Seed for pseudo-random	Numeric	A value of 0 will use a pseudo-random seed in each run, a value
number generator		0 will provide the same results in a repeated run.
Index substance	AlphaNumeric	The substance of interest or index substance.
Multiple substances analysis	Boolean	Specifies whether the assessment involves multiple substances.
Point of departure source	TargetDosesCalculationMethoa	characterisations: from points of departure (data), dose respons models or both.
Exposure routes	ExposureRoutes	Exposure routes included in assessment.
Apply kinetic conversions	Boolean	Convert substance concentrations from other biological matrice using kinetic conversion models. The substances for conversion are designated within the kinetic models module. Substance conversion proves valuable when a biomarker was not directly measured for a matrix of interest.
Restrict to single target	Boolean	Restrict to one biological matrix when computing substance
		hazard characterisations at the internal level.
Target biological matrix	BiologicalMatrix	The target biological matrix (internal compartment) for which
Specify the type of kinetic	InternalModelType	Specify the type of model to convert external exposure to the
model	internatioaet1 ype	internal level.
Expression type	PointOfDeparture	Specifies how hazard characterisations are expressed: as BMD, NOAEL, or the expression type is ignored.
Selection method in case of	TargetDoseSelectionMethod	Choose either the most toxic (default) or an aggregated hazard
multiple candidate hazard		characterisation when in nominal runs there are multiple available
characterisations		candidates. In uncertainty runs, multiple candidates are
		resampled.
Impute missing hazard	Boolean	If checked, missing hazard characterisations are imputed based
characterisations	Harman ID Investorian Made - I	Munro NOELs or on other available points of departure.
Imputation method	Type	unbiased central estimate from either the Munro set or the available POD collection.
Use lower limit of BMD	Boolean	Use the lower uncertainty limit of the benchmark dose. For an
		uncertainty run, the lower limit is the p5 of the bootstrap samp
		otherwise it is a parametric estimate based on the parameters o
		the dose response curve.
Use inter-species conversions	Boolean	Use inter-species conversion factors (default value, e.g. 10, or
I las inter ana size for stars	Declary	data).
Use intra-species factors	Boolean	Use intra-species conversion factors (default value, e.g. 10, or data)
Additional assessment factor	Numeric	Additional assessment factor for extrapolation of PODs to
	Trumerie	(human) hazard characterisations.
Use additional assessment	Boolean	Use additional assessment factor for extrapolation of PODs to
factor		(human) hazard characterisations.
Use hazard characterisations	Boolean	Hazard characterisations are dependent on subgroups (e.g. base
subgroup		on age or gender, see dataformats HCSubgroups).
Exposure events and dose	ExposureEventsGeneration-	Generates doses and events for PBK model
generation method	Method	
Number of simulated days	Numeric	The total duration (in days) of the PBK model simulations.
Number of finnal days skipped	Numeric	to be skipped when calculating the steady state or peak internal
		exposures from the simulated time series
Use PBK model parameter	Boolean	When specified, use parameter variability.
variability		1 , 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1,
Number of events per day for	Numeric	The daily dose is administered in equal portions (dose / number
the DERMAL dose		events) per event.
Number of events per day for	Numeric	The daily dose is administered in equal portions (dose / number
the INHALATION dose	N	events) per event.
Number of events per day for	Numeric	I ne daily dose is administered in equal portions (dose / number

Table 3.333: Calculation settings for module Hazard characterisations.

Uncertainty settings

Name	Туре	Description
Resample intra-species factor	Boolean	Specifies whether intra-species factors are resampled from a parametric uncertainty distribution.
Resample hazard characterisations or RPFs	Boolean	Specifies whether to resample the hazard characterisations or relative potency factors. Requires hazard characterisation or RI uncertainty to be quantified in DoseResponseModelsUncertain RelativePotencyFactorsUncertain tables.
Lower uncertainty limit (%)	Numeric	Percentage lower bound, e.g. 2.5%.
Upper uncertainty limit (%)	Numeric	Percentage upper bound, e.g. 97.5%.

Table 3.334: Uncertainty settings for module Hazard characterisations.

Hazard characterisations tiers

Overview

	Table 3.335:	Tier	overview for	r module	Hazard	characterisations.
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	N EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
۲ ا	Γ External g	External	External	External	External	External	External	External	External	Exte

Retrospective dietary CRA (EC 2018) - Acute / Tier I

Table 3.336: Tier definition for Retrospective dietary CRA (EC 2018) -

Acute /	Tier	I.
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Name	Setting
Target level	External

Retrospective dietary CRA (EC 2018) - Chronic / Tier I

Table 3.337: Tier definition for Retrospective dietary CRA (EC 2018) - Chronic / Tier I.

Name	Setting
Target level	External

Retrospective dietary CRA (EC 2018) - Acute / Tier II

Table 3.338: Tier definition for Retrospective dietary CRA (EC 2018) - Acute / Tier II.

Name	Setting
Target level	External

Retrospective dietary CRA (EC 2018) - Chronic / Tier II

Table 3.339: Tier definition for Retrospective dietary CRA (EC 2018) - Chronic / Tier II.

Name	Setting
Target level	External

Retrospective dietary CRA (EFSA 2012) - Optimistic

Use the optimistic model settings according to the EFSA Guidance 2012. Concentration values are sampled using a sample-based empirical distribution. Available processing factors are applied. No unit variability model should be applied.

Table 3.340: Tier definition for Retrospective dietary CRA (EFSA 2012)

- Optimistic.

Name	Setting
Target level	External

Retrospective dietary CRA (EFSA 2012) - Acute / Pessimistic

Acute probabilistic exposure assessment using the pessimistic model settings according to the EFSA Guidance 2012. Only processing factors > 1 are applied. For unit variability, the Beta distribution is applied.

Table 3.341: Tier definition for Retrospective dietary CRA (EFSA 2012) - Acute / Pessimistic.

Name	Setting
Target level	External

Retrospective dietary CRA (EFSA 2012) - Chronic / Pessimistic

Chronic probabilistic exposure assessment using the pessimistic model settings according to the EFSA Guidance 2012. Only processing factors > 1 are applied.

Table 3.342: Tier definition for Retrospective dietary CRA (EFSA 2012) - Chronic / Pessimistic.

Name	Setting
Target level	External

Retrospective dietary CRA (EFSA 2022) - Acute / Tier I

Table 3.343: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier I.

Name	Setting
Target level	External

Retrospective dietary CRA (EFSA 2022) - Chronic / Tier I

Table 3.344: Tier definition for Retrospective dietary CRA (EFSA 2022) - Chronic / Tier I.

Name	Setting
Target level	External

Retrospective dietary CRA (EFSA 2022) - Acute / Tier II

Table 3.345: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier II.

Name	Setting
Target level	External

Retrospective dietary CRA (EFSA 2022) - Chronic / Tier II

Table 3.346: Tier definition for Retrospective dietary CRA (EFSA 2022) - Chronic / Tier II.

Name	Setting
Target level	External

Prospective dietary CRA (EFSA 2023) - Acute / Tier II

Table 3.347: Tier definition for Prospective dietary CRA (EFSA 2023) - Acute / Tier II.

Name	Setting
Target level	External

Prospective dietary CRA (EFSA 2023) - Chronic / Tier II

Table 3.348: Tier definition for Prospective dietary CRA (EFSA 2023) - Chronic / Tier II.

Name	Setting
Target level	External

Prospective dietary CRA (EFSA 2024) - Acute / Tier II

Table 3.349: Tier definition for Prospective dietary CRA (EFSA 2024) - Acute / Tier II.

Name	Setting
Target level	External

Prospective dietary CRA (EFSA 2024) - Chronic / Tier II

Table 3.350: Tier definition for Prospective dietary CRA (EFSA 2024) - Chronic / Tier II.

Name	Setting
Target level	External

Hazard characterisations as data

Hazard characterisations (HCs) can be provided as data e.g., in the form of ADI or ARfD. For age dependent hazard characterisations, specify the HCs in table HCSubgroups and HCSubgroupsUncertain. Currently, only age dependent HCs are implemented.

- Hazard characterisations data formats
- Hazard characterisations from data

Inputs used: AOP networks Active substances Points of departure

Settings used

• Calculation Settings

Calculation of hazard characterisations

Hazard characterisations can be computed from points of departure. The computation may involve assessment factors, e.g. for inter-species conversion, intra-species variation or additional sources of uncertainty. The additional assessment factor can be used to bypass inter- and intra species conversion, or as an additional assessment factor to account for extrapolation for sources where appropriate data or information is scarce or missing (e.g. to implement a mixture assessment factor). The hazard characterisation calculation may also use kinetic conversion factors, PBK models or absorption factors to convert external doses to internal doses/concentrations or vice versa.

• Hazard characterisations calculation

Inputs used: Dose response models Effect representations Inter-species conversions Intra species factors Absorption factors Kinetic conversion factors PBK model parametrisations

Settings used

• Calculation Settings

3.5.9 Inter-species conversions

Inter-species conversions specify how to convert a hazard characterisation for a given species to a hazard characterisation for humans. In the simplest approach, this specifies a fixed inter-species factor. In a higher tier, this specifies a geometric mean (GM) and geometric standard deviation (GSD) for a lognormal uncertainty distribution of the interspecies factor. Inter-species conversion are specified per effect and can be general or substance-specific.

This module has as primary entities: Substances Effects

Output of this module is used by: Hazard characterisations

Inter-species conversions from data

Inter-species conversions data formats

Inter-species conversion models specify how to convert a hazard dose for a given species to a hazard dose for humans.

Download empty dataset template: Zipped CSV Excel

Inter-species model parameters

Inter-species extrapolation factors are described using a lognormal distribution specified by a geometric mean (GM) and geometric standard deviation (GSD). Inter-species factors are defined for an effect and a species and may optionally be specified specifically for a substance.

Name	Туре	Description	Aliases	Required
idEffect	AlphaNumeric (50)	The code of the effect for which this inter-species model is defined.	idEffect, EffectId, Effect	Yes
idSubstance	AlphaNumeric (50)	The code of the substance for which this inter-species model is defined.	idSubstance, SubstanceId, SubstanceCode, Substance	No
Species	AlphaNumeric (50)	Species	Species	Yes
InterSpecies- GeometricMean	Numeric	Interspecies geometric mean.	InterSpecies- GeometricMean, InterSpeciesGM	Yes
InterSpecies- Geometric- Standard- Deviation	Numeric	Interspecies geometric standard deviation.	InterSpecies- Geometric- Standard- Deviation, InterSpeciesGS- D	Yes
Standard- HumanBody- Weight	Numeric	The standard human body weight.	Standard- HumanBody- Weight	Yes
HumanBody- WeightUnit	Body Weight Unit	The unit of the human body weight specification (kg is assumed if not defined).	HumanBody- WeightUnit	No
Standard- AnimalBody- Weight	Numeric	The standard animal body weight.	Standard- AnimalBody- Weight	Yes
AnimalBody- WeightUnit	Body Weight Unit	The unit of the animal body weight specification (kg is assumed if not defined).	AnimalBody- WeightUnit	No

T 11 2 2 7 1	F 11 1 6 1.1	с. т	
Table 3.351:	Table definition	for Inter-species	model parameters.

Accepted table names: InterSpeciesModelParameters, InterSpeciesModelParameter, InterSpeciesFactors, InterSpeciesFactor.

Inter-species conversions settings

Selection settings

T 11	2 2 5 2	C 1		c ·	1 1 T .		•
Table	3 352	Selection	settings t	for mod	iule Infer	-species	conversions
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Name	Туре	Description
Interspecies factor geometric mean	Numeric	Interspecies factor geometric mean.
Interspecies factor geometric standard deviation	Numeric	Interspecies factor geometric standard deviation.
Restrict active substances to substances with available hazard characterisations	Boolean	Restrict assessment group membership based on presence/abse of hazard characterisations.

Calculation settings

Name	Туре	Description
Multiple substances analysis	Boolean	Specifies whether the assessment involves multiple substances.
Use inter-species conversions	Boolean	Use inter-species conversion factors (default value, e.g. 10, or data).

Table 3.353: Calculation settings for module Inter-species conversions.

Uncertainty settings

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Table 3 354.	Lincertainty	settings	for module	Inter_snecies	conversions
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Name	Туре	Description
Resample inter-species factor	Boolean	Specifies whether inter-species factors are resampled from a parametric uncertainty distribution.

Inter-species conversions as data

Data are provided in the form of a geometric mean (GM) and geometric standard deviation (GSD)

- Inter-species conversions data formats
- Inter-species conversions from data

Inputs used: Active substances

Settings used

• Calculation Settings

3.5.10 Intra species factors

Intra-species factors describe variation between individuals concerning their individual sensitivities to experience well-defined health effects. Traditionally the intraspecies factor is a fixed value, but the true distribution might be (very) uncertain. There is some support for assuming a lognormal distribution to describe the variability between individuals. In MCRA, intraspecies factors are sampled from a lognormal distribution, characterised by a geometric mean (GM) equal to 1 and a geometric standard deviation (GSD) thats needs to be given a value representing the intraspecies variability. GM is 1 by definition (50% of the population is assumed to be less sensitive than the average, 50% is mor sensitive) and has no uncertainty. On the other hand, there is uncertainty about the GSD. In MCRA it is assumed that this uncertainty is described by a Chi-square distribution with df degrees of freedom. By specifying a lower and upper bound for the p95 sensitive individual e.g. a lower value 2 and upper value 10 (meaning, the p95 individuals are between 2 and 10 times more sensible than the average human), a Chi-square distribution can be estimated where 1) the GSD specifies the variability and 2) the degrees of freedom specifies the uncertainty around the GSD.

This module has as primary entities: Substances Effects

Output of this module is used by: Hazard characterisations

Intra species factors from data

Intra species factors calculation

There is variation between individuals concerning their individual sensitivities to experience health effects. In some scenarios the aim is to perform assessments for the sensitive individuals instead of the average individuals for which the points of departure are derived. If this is the case, then extrapolation is required to translate hazard characterisations derived for the average individual to hazard characterisations for a sensitive individual.

Traditionally a fixed safety factor describes the variation between individuals. Little information is available about the true distribution of human sensitivities, so there is also a large uncertainty. In MCRA, the intra-species variability is modelled explicitly using *a lognormal distribution*, characterised by a geometric mean (GM) equal to 1 and a geometric standard deviation (GSD). This distribution is used to sample individual hazard characterisations. This effectively converts the description of hazard characterisations to include variability, with an unbiased central value. GM is 1 by definition (50% of the population is assumed to be less sensitive and 50% more sensitive than the average individual) and has no uncertainty. On the other hand, there is uncertainty about the GSD or *SD* = log(GSD). This uncertainty is described by a chi-square distribution with *df* degrees of freedom:

$$\frac{df\cdot SD^2}{\sigma^2}\sim \chi^2_{df}$$

where σ is the true standard deviation and χ^2_{df} denotes a chi-square distribution with df degrees of freedom. Thus df characterises the amount of uncertainty regarding the intra species variation quantified by (G)SD.

It is difficult to specify values for GSD and df. A practical way to do this is, define a p95-sensitive individual having an intra-species factor corresponding to the p95 percentile of the distribution describing the variability.

Assume that the p95-sensitive individuals are between e.g. 2 and 10 times more sensitive than the average human. The values 2 and 10 are to be interpreted as uncertainty bounds, the 2.5th and 97.5th percentiles of the uncertainty distribution for the $F_{intra, p95}$.

Then, p95-sensitive individuals are between $F_{intra, p95 p2.5}$ and $F_{intra, p95 p97.5}$ more sensitive than the average human, where the first index indicate the variability distribution and the second the uncertainty distribution.

For the lognormal distribution we have:

$$ln(F_{intra, p95}) = ln(GM) + 1.645 \cdot \sigma = 1.645 \cdot \sigma$$

And thus $\sigma = \frac{ln(F_{intra,p95})}{1.645}$. The uncertainty statement can be re-expressed as a 95% confidence interval for σ :

$$\frac{ln(F_{intra,\,p95\,p2.5})}{1.645} \le \sigma \le \frac{ln(F_{intra,\,p95\,p97.5})}{1.645}$$

A standard two-sided 95% confidence interval for σ derived from the χ^2 distribution is:

$$SD \cdot \sqrt{\frac{df}{\chi^2_{df,0.975}}} \leq \sigma \leq SD \cdot \sqrt{\frac{df}{\chi^2_{df,0.025}}}$$

Equating the two lower bounds as well as the two upper bounds we obtain two equations with two unknowns. Substituting for SD and rearranging we find that df can be found from:

$$\frac{\chi^2_{df,0.975}}{\chi^2_{df,0.025}} = \left(\frac{\ln(F_{intra,\,p95\,p97.5})}{\ln(F_{intra,\,p95\,p2.5})}\right)^2$$

And then GSD can be calculated as:

$$GSD = exp(SD) = (F_{intra, p95 \, p97.5}) \sqrt{\frac{\chi^2_{df, 0.025}}{df}} \ /1.645$$

A bisection algorithm can be used to find the value for df for which the chi-square ratio in the example is

$$\frac{\chi^2_{df,0.975}}{\chi^2_{df,0.025}} = \left(\frac{\ln(10)}{\ln(2)}\right)^2 = 11.035$$

This value is attained for df = 6.25. See also van der Voet et al. (2009)

Intra-species factors data formats

Intra-species factors.

Download empty dataset template: Zipped CSV Excel

Intra-species model parameters

Intra species factors.

Name	Туре	Description	Aliases	Required
idEffect	AlphaNumeric (50)	The effect code.	idEffect, EffectId, Effect	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	No
IntraSpecies- Lower- VariationFactor	Numeric	The lower uncertainty bound. The p95-sensitive-individual is 'lower bound' times more sensitive than the average individual.The lower and upper bounds are used to derive a geometric standard deviation (gsd) and degrees of freedom (df).	IntraSpecies- LowerVariation- Factor	No
IntraSpecies- UpperVariation- Factor	Numeric	The upper uncertainty bound. The p95-sensitive-individual is 'upper bound' times more sensitive than the average individual. The lower and upper bounds are used to derive a geometric standard deviation (gsd) and degrees of freedom (df).	IntraSpecies- UpperVariation- Factor	Yes
idPopulation	AlphaNumeric (50)	Unique identification code of the population.	IdPopulation, PopulationId	No

Table 3.355: Table definition for Intra-species model parameters.

Accepted table names: IntraSpeciesModelParameters, IntraSpeciesModelParameter, IntraSpeciesFactors, IntraSpeciesFactor.

Intra species factors settings

Selection settings

	e	•
Name	Туре	Description
Intra-species factor	Numeric	Intra-species factor.
Restrict active substances to	Boolean	Restrict assessment group membership based on presence/abser
substances with available		of hazard characterisations.
hazard characterisations		
	Name Intra-species factor Restrict active substances to substances with available hazard characterisations	NameTypeIntra-species factorNumericRestrict active substances to substances with available hazard characterisationsBoolean

Table 3.356: Selection settings for module Intra species factors.

Calculation settings

Name	Туре	Description
Multiple substances analysis	Boolean	Specifies whether the assessment involves multiple substances.

Uncertainty settings

Name	Туре	Description
Resample intra-species factor	Boolean	Specifies whether intra-species factors are resampled from a parametric uncertainty distribution.

Table 3.358: Uncertainty settings for module Intra species factors.

Intra species factors as data

In the simplest approach, intra-species factors are fixed factors. In a higher tier, lower and upper values for the intraspecies factor are used to derive a variability distribution (log-normal around 1) and an uncertainty distribution for the geometric standard deviation related to human variability in sensitivity.

- Intra species factors data formats
- Intra species factors from data
- Intra species factors calculation

Inputs used: Active substances

Settings used

• Calculation Settings

3.5.11 Points of departure

Externally specified points of departure can be used as an alternative to calculated BMDs from dose response models. Points of departure can be of various types, such as NOAEL, LOAEL or BMD. They can be used to construct the list of active substances, to derive relative potency factors, and to perform health impact assessments.

This module has as primary entities: Substances Effects

Output of this module is used by: Active substances Hazard characterisations

Points of departure from data

Points of departure data formats

Points of departure, such as NOAELS and BMDs, describe the critical/reference levels of substance dose in relation to the presence or absence of an effect. If available, the uncertainty of externally specified points of departure can be specified with uncertainty sets (empirical distributions representing possible values) in the points of departure uncertainty table.

Download empty dataset template: Zipped CSV Excel

Points of departure

Nominal points of departure should be presented in this table. Each point of departure should be linked to an effect using the effect code (idEffect) and to substances using the substance code (idSubstance).

Name	Туре	Description	Aliases	Required
idModel	AlphaNumeric (50)	The dose response model code.	idDose- ResponseModel, idModel, idPod, idPointOf- Departure	No
idEffect	AlphaNumeric (50)	The effect code.	idEffect, EffectId, Effect	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
Species	AlphaNumeric (50)	The species used to obtain this point of departure.	Species, System	No
Value	Numeric	Point of departure, can be of various types, e.g. NOAEL, LOAEL, BMD, CED	PointOf- Departure, LimitDose, HazardDose, Value, CED	Yes
Туре	PointOfDeparture- Type	The type of the point of departure, e.g. NOAEL, LOAEL, BMD.	Type, PODType, HazardDose- Type, LimitDoseType	No
DoseUnit	DoseUnit	The dose unit (if not specified, then mg/kg is assumed).	DoseUnit, Unit, UnitDose	No
Benchmark- Response	AlphaNumeric (100)	The benchmark response or effect size.	Benchmark- Response, CriticalEffect- Size, HazardEffect- Size	No
ExposureRoute	ExposureRoute	The route of dose administration used in the study to obtain this point of departure. If not specified exposure route = Dietary is assumed.	ExposureRoute, RouteExposure	No
IsCriticalEffect	Boolean	Specifies whether this value is the value associated with the critical effect. If omitted, No is assumed.	IsCriticalEffect	No
TargetLevel	TargetLevelType	The target level, i.e., internal or external. If omitted, external is assumed.	TargetLevel	No
Matrix	<i>BiologicalMatrix</i>	The biological matrix or organ (should be specified when target level is internal).	Matrix, BiologicalMatrix	No
Expression type	ExpressionType	The expression type for (internal) hazard characterisations of which the concentrations are not directly specified at the level of the biological matrix. For instance, urinary expressions expressed in terms of the amount excreted per gram of	ExpressionType	No
5. Hazard mod	ules	creatinine or blood concentration expressed per gram of lipids.		54
PublicationTitle	AlphaNumeric (250)	Title of the publication of the	PublicationTitle,	No

Table 3.359: Table definition for Points of departu	ire.
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Accepted table names: PointsOfDeparture, PointOfDeparture, HazardDoses, HazardDose.

Points of departure uncertainty

Often, the PODs found for a substance/effect combination are uncertain. This table facilitates in specifying the POD uncertainty in the form of a set of uncertainty values that may additionally be specified for a substance/effect combination.

Name	Туре	Description	Aliases	Required
idDose- ResponseModel	AlphaNumeric (50)	The dose response model code (must correspond to values in id column of DoseResponseModels table).	idDose- ResponseModel	Yes
idUncertainty- Set	AlphaNumeric (50)	The identification code of the uncertainty set. During an uncertainty iteration one set will be picked to be the POD value.	idUncertainty- Set, UncertaintyId	Yes
idEffect	AlphaNumeric (50)	The effect code.	idEffect, EffectId, Effect	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
Point of departure	Numeric	Point of departure, can be of various types, e.g. NOAEL, LOAEL, BMD, CED	PointOf- Departure, HazardDose, LimitDose, CED	Yes
DoseResponse- Model- Parameter- Values	AlphaNumeric (200)	A comma separated list of the values of the parameters of the model, format: a=1.2,b=3.4,c=5.6	DoseResponse- Model- Parameter- Values, ParameterValues	No

Table 3.360: Table definition for Points of departure uncertainty.

Accepted table names: PointsOfDepartureUncertain, PointOfDepartureUncertain, HazardDosesUncertain, HazardDoseUncertain.

Points of departure settings

Selection settings

Table 3.361: Selection settings	s for module Points of	departure.
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Name	Туре	Description
Multiple effects analysis	Boolean	Specifies whether the analysis should consider multiple effects. Otherwise, a single focal effect should be selected.
Include related effects of AOP network	Boolean	Include all related key events of the AOP network.

Uncertainty settings

J. J			
Name	Туре	Description	
Perform uncertainty analysis	Boolean	In probabilistic risk assessment of dietary exposure, distribution describe the variability in consumption within a given population of individuals and the variability of the occurrence and level of substances in the consumed foods. However, these calculations not consider the amount of uncertainty that is due to the limited size of the underlying datasets.	
Resample hazard characterisations or RPFs	Boolean	Specifies whether to resample the hazard characterisations or relative potency factors. Requires hazard characterisation or RI uncertainty to be quantified in DoseResponseModelsUncertain RelativePotencyFactorsUncertain tables.	

Table 3.362: Uncertainty settings for module Points of departure.

Points of departure as data

Points of departure are provided as data for combinations of substance and effect and each is minimally described by a reference value and a type (e.g., NOAEL or LOAEL). In addition, the exposure route, specifies, and references may be specified.

- Points of departure data formats
- Points of departure from data

Inputs used: AOP networks

3.5.12 Relative potency factors

Relative potency factors (RPFs) quantify potencies of substances with respect to a defined effect, relative to the potency of a chosen index substance. RPFs can be used to express combined exposures of multiple substances in terms of a the exposure value of the chosen index substance (i.e., in index substance equivalents). In MCRA, hazard characterisations, and therefore also RPFs are based on mass units (e.g., μ g), and not on mol units. RPFs can be different for different levels of the human organism (external, internal, specific compartment). RPFs can be given as data or computed from hazard characterisations. RPFs can be specified with uncertainty. Computation from uncertain hazard characterisations allows to include correlations between uncertain RPFs which originate from using the same index substance.

This module has as primary entities: Substances Effects

Output of this module is used by: Concentrations Concentration models High exposure food-substance combinations Dietary exposures Internal exposures Exposure mixtures HBM analysis Risks

Relative potency factors from data

Relative potency factors calculation

Relative potency factors (*RPF* s) describe the potency of substances with respect to a defined effect, relative to the potency of a chosen index or reference substance. RPFs are provided as data or computed from *hazard characterisa-tions*. The RPF for substance *i* is defined by the ratio of hazard characterisation value (*POD*) for the index substance (ref) and the hazard characterisation value for substance *i*. That is,

$$RPF_i = POD_{ref} / POD_i$$
.

When hazard characterisations are resampled in the uncertainty runs, RPFs are recomputed based on the bootstrapped hazard characterisations. In this way, RPF uncertainty is included in the uncertainty analysis.

Relative potency factors data formats

Relative potency factors quantify relative potencies of substances with respect to an effect and can be used to express combined exposures of multiple substances in terms of the exposure value of the chosen index substance (i.e., as index substance equivalents). Relative potency factors can be provided in case hazard characterisations are missing. If available, the uncertainty of externally specified RPFs can be specified with uncertainty sets (empirical distributions representing possible values) in an additional table.

Download empty dataset template: Zipped CSV Excel

Relative potency factors

Relative potency factors are linked to an effect using the effect code (idEffect) and to substances using the substance code (idSubstance).

Name	Туре	Description	Aliases	Required
idCompound	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
idEffect	AlphaNumeric (50)	The effect code.	idEffect, EffectId, Effect	Yes
RPF	Numeric	The relative potency factor.	RPF, Relative- PotencyFactor, Value	Yes
PublicationTitle	AlphaNumeric (250)	Title of the publication of the study in which this relative potency factor was established.	PublicationTitle, Title	No
Publication- Authors	AlphaNumeric	Author(s) of the publication of the study in which this relative potency factor was established.	Publication- Authors, Publication- Author, Author, Authors	No
PublicationYear	Integer	Year of the publication of the study in which this relative potency factor was established.	PublicationYear, Year	No
PublicationUri	AlphaNumeric	Uniform resource identifier of the reference publication.	URI, URL, PublicationURI, PublicationURL	No
Description	AlphaNumeric (200)	Additional description of the relative potency factor.	Description, Remark	No

Table 3.363: '	Table definition	for Relative	potency factors.
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Accepted table names: RelativePotencyFactors, RelativePotencyFactor.

Relative potency factor uncertainty

This table contains sets of values representing the uncertainty for relative potency factors.

Name	Туре	Description	Aliases	Required
idUncertainty- Set	AlphaNumeric (50)	The uncertainty set identification number. During each uncertainty iteration one set is used.	idUncertainty- Set, UncertaintyId	Yes
idEffect	AlphaNumeric (50)	The effect code (must correspond to values in id column of Effects table).	idEffect, EffectId, Effect	Yes
idSubstance	AlphaNumeric (50)	The substance code (must correspond to values in id column of Substances table).	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
RPF	Numeric	The relative potency factor.	RPF, Relative- PotencyFactor, Value	Yes

Accepted table names: RelativePotencyFactorsUncertain, RelativePotencyFactorUncertain.

Relative potency factors settings

Calculation settings

Table 3.365:	Calculation settings	for module	Relative	potency factors.
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Name	Туре	Description
Multiple effects analysis	Boolean	Specifies whether the analysis should consider multiple effects. Otherwise, a single focal effect should be selected.
Include related effects of AOP network	Boolean	Include all related key events of the AOP network.
Index substance	AlphaNumeric	The substance of interest or index substance.

Uncertainty settings

Table 3.366: Uncertainty settings for module Relative potency factors.

Name	Туре	Description
Resample hazard characterisations or RPFs	Boolean	Specifies whether to resample the hazard characterisations or relative potency factors. Requires hazard characterisation or RI uncertainty to be quantified in DoseResponseModelsUncertain RelativePotencyFactorsUncertain tables.
Lower uncertainty limit (%)	Numeric	Percentage lower bound, e.g. 2.5%.
Upper uncertainty limit (%)	Numeric	Percentage upper bound, e.g. 97.5%.

Relative potency factors as data

Data are provided in the form of a RPF for a specific substance and effect.

- Relative potency factors data formats
- Relative potency factors from data

Inputs used: Active substances AOP networks

Settings used

• Calculation Settings

Calculation of relative potency factors

RPFs are computed from hazard characterisations.

• Relative potency factors calculation

Inputs used: Hazard characterisations

Settings used

• Calculation Settings

3.6 In-silico modules

Two types of *in-silico* models are available: QSAR models specify assessment group memberships for active substances, as numbers in the interval [0,1]. This allows both crisp (0 or 1) and probabilistic memberships. Molecular docking models specify binding energies and thresholds which can be used to convert binding energies to assessment group memberships for active substances.

3.6.1 Molecular docking models

Molecular docking models specify binding energies for substances in specific molecular docking models related to a specific health effect (adverse outcome).

This module has as primary entities: Substances Effects

Output of this module is used by: Active substances

Molecular docking models from data

'In the field of molecular modeling, docking is a method which predicts the preferred orientation of one molecule to a second when a ligand and a target are bound to each other to form a stable complex.[1] Knowledge of the preferred orientation in turn may be used to predict the strength of association or binding affinity between two molecules using, for example, scoring functions....'

See In Wikipedia.

Molecular docking models data formats

Required data tables:

- Molecular docking models, to identify models for a specified effect (receptor)
- Molecular docking binding energies, to specify the binding energies per substance for the receptor

Contains definitions of molecular docking models for a given effect (molecular initiating event), for example parameters needed in the conversion of binding energies to group memberships or to relative potency factors. Substance specific binding energies are specified in the binding energies table.

Download empty dataset template: Zipped CSV Excel

Molecular docking models

Each docking model has a unique identifier, and optionally a name and a description. Each model is linked to an effect using the idEffect field and optionally a binding threshold and the number of receptors can be added. A reference to the source of the data can be stored in the reference field.

Name	Туре	Description	Aliases	Required
id	AlphaNumeric (50)	The unique identification code of the molecular docking model.	idMolecular- DockingModel, idBinding- EnergyModel	Yes
Name	AlphaNumeric (100)	The name of the molecular docking model.	Name	No
Description	AlphaNumeric (200)	Description of the molecular docking model.	Description	No
idEffect	AlphaNumeric (50)	The effect code, typically for the Molecular Initiating Event that is modelled	idEffect, EffectId, Effect	Yes
Threshold	Numeric	Threshold Molecular Docking binding energy (group membership = 1 when higher).		Yes
NumberOf- Receptors	Integer	Example parameter needed for translating Molecular Docking binding energies to RPFs.		No
Reference	AlphaNumeric (200)	External reference(s) to sources containing more information about the molecular docking model.	References	No

Table 3.367:	Table definition	for Molecular	docking models.

Accepted table names: MolecularDockingModels, MolecularDockingModel, BindingEnergyModels, BindingEnergyModel.

Molecular docking binding energies

Molecular docking model binding energies per substance

Table 3.368:	Table definition	for Molecular	docking bind	ling energies.
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Name	Туре	Description	Aliases	Required
idMolecular- DockingModel	AlphaNumeric (50)	The id of the molecular docking model or source.	idMolecular- Docking, Molecular- DockingModel	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
BindingEnergy	Numeric	Molecular Docking binding energy.	Molecular- Docking- BindingEnergy	Yes

Accepted table names: MolecularBindingEnergies, MolecularBindingEnergy, BindingEnergies, BindingEnergy, MolecularDockingBindingEnergy.

Molecular docking models

Calculation settings

Name	Туре	Description
Multiple effects analysis	Boolean	Specifies whether the analysis should consider multiple effects. Otherwise, a single focal effect should be selected.
Include related effects of AOP network	Boolean	Include all related key events of the AOP network.

Table 3.369: Calculation settings for module Molecular docking models.

Molecular docking models as data

Binding energies for substances in specific molecular docking models related to a specific health effect (adverse outcome) are provided as data.

- Molecular docking models data formats
- Molecular docking models from data

Inputs used: *AOP networks*

Settings used

• Calculation Settings

3.6.2 QSAR membership models

QSAR membership models specify assessment group memberships for active substances related to a specific health effect (adverse outcome). Memberships should be derived externally from Quantitative Structure-Activity Relationship (QSAR) models.

This module has as primary entities: Substances Effects

Output of this module is used by: Active substances

QSAR membership models from data

'Quantitative structure-activity relationship models (QSAR models) are regression or classification models used in the chemical and biological sciences and engineering. Like other regression models, QSAR regression models relate a set of "predictor" variables (X) to the potency of the response variable (Y), while classification QSAR models relate the predictor variables to a categorical value of the response variable....'

See In Wikipedia.

QSAR membership models data formats

Required data tables:

- QSAR membership models, to identify QSAR models for a specified health effect
- QSAR membership scores, to specify the memberships per substance per QSAR model

Note that only memberships 1 (include) and 0 (exclude) are allowed.

Substance membership models obtained from QSAR for a given (health) effect. The models are defined in the membership models table, and substance specific memberships are specified in the QSAR memberships table.

Download empty dataset template: Zipped CSV Excel

QSAR membership models

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This table contains the definitions of the QSAR membership models. Each model contains a id, name, an optional description, and refers to its related health effect.

Name	Туре	Description	Aliases	Required
id	AlphaNumeric (50)	The unique identification code of the QSAR membership model.	id, Model, ModelCode, idModel, QSARModel, idQSARModel, QSAR- Membership- Model, idQSAR- Membership- Model, Membership- Model, idMembership- Model,	Yes
Name	AlphaNumeric (100)	The name of the QSAR membership model.	Name	No
Description	AlphaNumeric (200)	Description of the QSAR membership model.	Description	No
idEffect	AlphaNumeric (50)	The effect code.	idEffect, EffectId, Effect	Yes
Accuracy	Numeric	Accuracy of the QSAR membership model.	Accuracy	No
Sensitivity	Numeric	Sensitivity of the QSAR membership model.	Sensitivity	No
Specificity	Numeric	Specificity of the QSAR membership model.	Specificity	No
Reference	AlphaNumeric (200)	External reference(s) to sources containing more information about the QSAR model.	References	No

Table 3.370	: Table	definition	for OSAR	membershir	models.
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Accepted table names: QSAR, QSARMembershipModels, QSARMembershipModel, QSARModels, QSARModel.

QSAR membership scores

Substance membership score according to the QSAR model.

Name	Туре	Description	Aliases	Required
idQSAR- Membership- Model	AlphaNumeric (50)	The id of the QSAR model.	Model, ModelCode, idModel, QSARModel, idQSARModel, QSAR- Membership- Model, idQSAR- Membership- Model, Membership- Model, idMembership- Model,	Yes
idSubstance	AlphaNumeric (50)	The code of the substance.	idSubstance, SubstanceId, SubstanceCode, Substance	Yes
Membership- Score	Numeric	QSAR membership score. Value should be 1 for positive membership, or 0 for negative membership.	Membership- Score, Membership, QSARScore, Score	Yes

Table 3.371: Table definition for QSAR me	embership scores.
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Accepted table names: QSARMembershipScores, QSARMembershipScore, QSARMemberships, QSARMembership.

QSAR membership models

Calculation settings

Table 3.372:	Calculation	settings for	module QSAR	membership n	nodels.
		0			

Name	Туре	Description
Multiple effects analysis	Boolean	Specifies whether the analysis should consider multiple effects. Otherwise, a single focal effect should be selected.
Include related effects of AOP network	Boolean	Include all related key events of the AOP network.

QSAR membership models as data

QSAR memberships models are provided as data, per QSAR model assessment group memberships for active substances related to a specific health effect are specified.

- QSAR membership models data formats
- QSAR membership models from data

Inputs used: AOP networks

Settings used

• Calculation Settings

3.7 Kinetic modules

Kinetic models convert exposures or hazard characterisations from one or more external routes or compartments to an internal compartment or biological matrix. Internal exposure can be systemic or related to a specific biological matrix in a kinetic model (target). The reverse conversion from internal to external is called reverse dosimetry.

In a simple tier, kinetic models are specified as absorption factors (systemic). In a higher tier, physiologically based (toxico)kinetic (PBK) models of a specified type (currently available is the EuroMix generic PBK model) are linked to MCRA. Kinetic model instances for specific substances and test systems (e.g. cypermethrin in the rat) are specified with parameter sets for the chosen kinetic model.

There are 3 tiers for relating external to internal exposures (doses):

- 1. Assume conservative absorption factors as suggested by EFSA (EFSA (2014), EFSA (2017a)): oral and inhalation 100%, dermal 50%. The internal exposure is systemic (*absorption factors data tables*).
- 2. Use externally provided kinetic conversion factors, converting the exposure to an internal target (biological matrix) or organ (*kinetic conversion factors data tables*).
- 3. Use one of the *implemented PBK models*, with instances for specific substances defined in data table *kinetic model instances* and model parameters specified in data table *kinetic model instance parameters*.

Given a chosen tier, the calculation will fall back to the next lower tier in case of missing data.

3.7.1 Absorption factors

Absorption factors are simple kinetic conversion factors used to convert external exposure from one or more routes to an internal dose (systemic).

This module has as primary entities: Substances

Output of this module is used by: Internal exposures Hazard characterisations

Absorption factors from data

Absorption factors data formats

Data tables:

• Absorption factors

Absorption factors are chemical- and route-specific values representing the fraction of external exposures passing the absorption barrier.

Kinetic absorption data format

Absorption factor data format.

Download empty dataset template: Zipped CSV Excel

Absorption factors

Absorption factors translating external (daily) exposures for specific routes to the resulting internal (systemic, daily) intakes.

Name	Туре	Description	Aliases	Required
idCompound	AlphaNumeric (50)	code of substance (must correspond to values in id column of Substances table)	idSubstance, SubstanceId, SubstanceCode, Substance	No
Route	<i>ExposurePathType</i>	Non-dietary route or pathway, use 'Oral', 'Dermal', or 'Inhalation' to specify the route.	Route, Pathway	Yes
Absorption- Factor	Numeric	absorption factor value	Absorption- Factor, Factor	Yes

Accepted table names: KineticAbsorptionFactors, KineticAbsorptionFactor, AbsorptionFactors, AbsorptionFactor.

Absorption factors settings

Calculation settings

Name	Туре	Description
Multiple substances analysis	Boolean	Specifies whether the assessment involves multiple substances.
Exposure routes	ExposureRoutes	Exposure routes included in assessment.
Restrict active substances to substances with available hazard characterisations	Boolean	Restrict assessment group membership based on presence/abser of hazard characterisations.
Default oral absorption factor for non-dietary exposure	Numeric	When there is no kinetic model and absorption factors are not specified in file, non-dietary oral exposures (external doses) are multiplied by this factor to determine the absorbed (internal) do
Default oral absorption factor for dietary exposure	Numeric	When there is no kinetic model and absorption factors are not specified in file, dietary exposures (external doses) are multiplie by this factor to determine the absorbed (internal) dose.
Default dermal absorption factor for non-dietary exposure	Numeric	When there is no kinetic model and absorption factors are not specified in file, dermal exposures (external doses) are multiplied by this factor to determine the absorbed (internal) dose.
Default inhalation absorption factor for non-dietary exposure	Numeric	When there is no kinetic model and absorption factors are not specified in file, inhalation exposures (external doses) are multiplied by this factor to determine the absorbed (internal) do

Table 3.374: Calculation settings for module Absorption factors.

Absorption factors as data

Absorption factors are chemical- and route-specific values representing the fraction of external exposures passing the absorption barrier. The corresponding exposure is referred to as internal dose.

- Absorption factors data formats
- Absorption factors from data

Inputs used: Active substances

Settings used

• Calculation Settings
3.7.2 Kinetic conversion factors

External exposure can be from one or more exposure routes: oral (dietary or non-dietary), dermal or inhalation and is converted to an internal concentration by a kinetic conversion factor. The internal concentration is related to a specific compartment (*biological matrix*) in a kinetic model.

This module has as primary entities: Substances

Output of this module is used by: Internal exposures HBM analysis Hazard characterisations

Kinetic conversion factors from data

In table *Kinetic conversion factors*, conversion factors are specified to convert biomarkers from one *biological matrix* to another biological matrix. This specification may include *expression type* and/or *exposure route*.

Kinetic conversion factors may be dependent on individual properties like age and *gender*. Specify in table *Kinetic conversion factor subgroups* (kinetic conversion factor subgroups) age and/or gender specific kinetic conversion factors.

For age, specify the lower bound of the age interval (in years) of the kinetic conversion factor subgroup. Individuals belong to a subgroup when the age of the individual is equal or greater than the specified lower bound and smaller than the specified lower age of the next subgroup.

Check option *Use kinetic conversion factors subgroup* (default) to use age and/or gender specific kinetic conversion factors in your assessment (only visible when the datasource contains age/gender specific subgroup information).

For more information, see *internal concentrations using kinetic conversion factors* in module Internal exposures for conversion to blood standardised concentrations by lipids and/or urine standardised or normalised concentrations by repectively creatinine or specific gravity.

For kinetic conversion factor uncertainty distributions see uniform, lognormal, beta and inverse uniform distribution.

Kinetic conversion factors data formats

Data tables:

• Kinetic conversion factors

Kinetic conversion factors convert external exposures from one or more external routes (dietary, non-dietary oral, dermal, inhalation) to an internal (target) compartment.

Kinetic conversion data format

The kinetic conversion factor data format.

Download empty dataset template: Zipped CSV Excel

Kinetic conversion factors

Kinetic conversion factors translating external (daily) exposures for specific routes to the resulting internal target intakes.

Name	Туре	Description	Aliases	Required
idKinetic- Conversion- Factor	AlphaNumeric (50)	Id of the kinetic conversion factor	idKinetic- Conversion- Factor, idConversion- Factor, idKCFactor	Yes
idSubstance- From	AlphaNumeric (50)	Code of substance (must correspond to values in id column of Substances table)	idSubstance- From, SubstanceId- From, SubstanceCode- From, SubstanceFrom	Yes
ExposureRoute- From	ExposureRoute	The exposure route (only applicable if target level is external). If not specified, Dietary is assumed.	ExposureRoute	No
Biological matrix from	BiologicalMatrix	If applicable, the source matrix of the human body (e.g., blood, urine) to convert from. When specified, the measurements are considered at the level of internal doses.	MatrixSource, SourceMatrix	No
DoseUnitFrom	DoseUnit	The unit of the substance dose to convert from.	UnitSource, SourceUnit	Yes
Expression- TypeFrom	ExpressionType	The expression type or adjustment method of the dose unit (from). This field specifies how the dose unit (source) is adjusted, e.g. for blood lipids for fat soluble biomarkers ('mg/g lipids') or the dilution level of the urine ('mg/g creatinine').	ExpressionType- From, Adjustment- MethodFrom	No
idSubstanceTo	AlphaNumeric (50)	Code of substance (must correspond to values in id column of Substances table)	idSubstanceTo, SubstanceIdTo, SubstanceCode- To, SubstanceTo	Yes
ExposureRoute- To	ExposureRoute	The exposure route (only applicable if target level is external) to convert to. If not specified, Dietary is assumed.	ExposureRoute	No
Biological matrix to	<i>BiologicalMatrix</i>	If applicable, the matrix of the human body (e.g., blood, urine) to convert to. When specified, the measurements are considered at the level of internal doses.	MatrixTarget, TargetMatrix	No
DoseUnitTo	DoseUnit	The unit of the substance dose to convert to.	UnitTarget, TargetUnit	Yes
Expression- TypeTo	ExpressionType	The expression type or adjustment method of the dose unit (to). This field specifies how the dose unit (target) is adjusted, e.g. for blood lipids for fat soluble	ExpressionType- To, Adjustment- MethodTo	No
62		biomarkers ('mg/g lipids') or the dilution level of the urine ('mg/g creatinine').	Chapte	r 3. Modules
Conversion-	Numeric	Conversion factor value	Conversion- Factor Factor	Yes

Table 3.375:	Table defi	nition for K	Linetic conve	ersion factors.

Accepted table names: KineticConversionFactors, KineticConversionFactor.

Kinetic conversion factor subgroups

Kinetic conversion factor subgroups.

Name	Туре	Description	Aliases	Required
idKinetic- Conversion- Factor	AlphaNumeric (50)	Id of the kinetic conversion factor	idKinetic- Conversion- Factor, idConversion- Factor, idKCFactor	Yes
Conversion- Factor	Numeric	Conversion factor value	Conversion- Factor, Factor	Yes
AgeLower	Numeric (50)	Specifies the lower bound of the age interval (in years) of the kinetic conversion factor subgroup. Individuals belong to a subgroup when the age of the individual is equal or greater than the specified lower bound and smaller than the specified lower age of the next subgroup.	AgeLower, LowerAge	No
Gender	GenderType	The gender of the kinetic conversion factor subgroup.	Gender, Sex	No
Uncertainty- Upper	Numeric	The upper value of the distribution. If the distribution is uniform, then it is the upper bound of the uniform distribution. If the distribution is log-normal, then the upper value is assumed to correspond with the p95 percentile of the distribution. When a distribution is assumed, this value should be greater than the conversion factor.	Uncertainty- Upper, Upper	No

T 11 2 27	TT 1 1 C '.'	C TZ' /'	•	C / 1
Table $3.3/6$	Lable definition	for Kinefic	conversion	factor subgroups
14010 010701	ruote definition	for remetic	conversion	nuctor subgroups.

Accepted table names: KineticConversionFactorSGs, KCFactorSubGroups.

Kinetic conversion factors settings

Calculation settings

Name	Туре	Description
Multiple substances analysis	Boolean	Specifies whether the assessment involves multiple substances.
Exposure routes	ExposureRoutes	Exposure routes included in assessment.
Restrict active substances to substances with available hazard characterisations	Boolean	Restrict assessment group membership based on presence/abse of hazard characterisations.
Use kinetic conversion factors subgroup	Boolean	Kinetic conversion factors are dependent on subgroups (e.g. ba on age or gender, see dataformats KineticConversionfactorsSG

Table 3.377: Calculation settings for module Kinetic conversion factors.

Uncertainty settings

Table 3.378: Uncertainty settings for module Kinetic conversion	n factors.
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Name	Туре	Description
Resample kinetic conversion factors	Boolean	Specifies whether kinetic conversion factors are resampled.

Kinetic conversion factors as data

Specify kinetic conversion factors as data.

- Kinetic conversion factors data formats
- Kinetic conversion factors from data

Inputs used: Active substances

Settings used

• Calculation Settings

3.7.3 PBK model parametrisations

External exposure can be from one or more exposure routes: oral (dietary or non-dietary), dermal or inhalation. Internal exposures are related to a specific *biological matrix*, in a kinetic model.

Use one of the *implemented kinetic models*, with instances for specific substances defined in data table *kinetic model instances* and model parameters specified in data table *kinetic model instance parameters*.

Given a PBK model the calculation will fall back to a lower tier, see *kinetic conversion factors module*, in case of missing data.

This module has as primary entities: Substances

Output of this module is used by: Internal exposures Hazard characterisations

PBK model parametrisations from data

To simulate the fate of a substance in a compartment, the *PBK* models in MCRA are run using different dosing patterns. In general, the length of the simulation period is specified in days.

The Exposure events and dose generation setting specifies how repeated dose patterns are applied.

• Use usual (average) exposures per route to specify a single dose per day, repeated over the entire simulation length. The dose applied is for an acute exposure assessment equal to a daily dose, for a chronic assessment equal to the average of all daily exposures of an individual (e.g. *OIM*).

• Use (randomly selected) daily exposures per route to specify more complex dosing patterns. The daily exposures (in a chronic context) are randomly allocated to the events, so for one event per day, the entire sequence of doses is a random sequence of the daily doses of one individual and, in case of a two 24hour recall, two different doses are sampled. When the specified number of events is > 1, the daily doses are divided by the specified number of events in a day and randomly allocated to the events. Note that for PBK models with a resolution in hours, the number of events should be equal to 1, 2, 3, 4, 6, 8 and 12.

For irregular dosing patterns use setting *Specify event timings*. Allowed numbers are 1, 2, 3,...,24 separated by spaces (e.g. 1 2 7 12).

Currently, option *Specify event timings* is only implemented for the oral route and for PBK models with resolution in hours.

PBK model parametrisations data formats

Data tables:

- Kinetic model instances
- Kinetic model instance parameters

PBK model parametrisations contain specifications for kinetic model definitions and their parameterisation for build in (C-code) and uploadable (SBML) kinetic models.

PBK model data format

The PBK model parametrisations data format specifies PBK model definitions and parameters.

Download empty dataset template: Zipped CSV Excel

PBK model instances

PBK model instances/parametrisations.

Name	Туре	Description	Aliases	Required
idModel- Instance	AlphaNumeric (50)	Unique identification code of the PBK model instance/parametrisation.	idModel- Instance, Id, Code	Yes
idModel- Definition	KineticModelType	Identifier of the PBK model definition of which this is an instance/parametrisation.	idModel- Definition, ModelDefinition	Yes
Species	AlphaNumeric (50)	The species to which this instance/parametrisation applies.	Species, System, TestSystem	Yes
Substances	AlphaNumeric (150)	Code or comma separated list of the codes of the substances. Only applies for substance-specific parameters.	idSubstance, idSubstances, SubstanceId, SubstanceCode, SubstanceCodes, Substance, Substance,	No
Reference	AlphaNumeric (100)	Reference or author.	References	No
Name	AlphaNumeric (100)	Name of the kinetic model instance/parametrisation.	Name	No
Description	AlphaNumeric (200)	Additional description of the model instance/parametrisation.	Description	No

Table 3.379: Table definition for PBK model instances.

Accepted table names: PbkModelInstances, PbkModelParametrisations, KineticModelInstances, KineticModelInstance.

PBK model instance parameters

PBK model instance/parametrisation parameter values.

Name	Туре	Description	Aliases	Required
idModel- Instance	AlphaNumeric (50)	Unique identification code of the kinetic model instance to which this parameter belongs	Id, Code	Yes
Parameter	AlphaNumeric (50)	Identifier of the parameter in the PBK model.		Yes
Description	AlphaNumeric (200)	Description of or reference for the parameter values in the model.		No
Value	Numeric	Nominal parameter value of the distribution.	Mean	Yes
Distribution- Type	PbkModelParameter- DistributionType	Parameter distribution type.	Distribution- Type, Distribution	No
CvVariability	Numeric	Variability of the parameter expressed as a coefficient of variation (CV).		No
CvUncertainty	Numeric	Uncertainty of the parameter expressed as a coefficient of variation (CV).		No

Table 3.380: Table definition for PBK model instance parameters .

Accepted table names: PbkModelInstanceParameters, PbkModelInstanceParameterValues, KineticModelInstanceParameters, KineticModelInstanceParameter.

PBK model parametrisations settings

Calculation settings

Name	Туре	Description
Multiple substances analysis	Boolean	Specifies whether the assessment involves multiple substances.
Exposure routes	ExposureRoutes	Exposure routes included in assessment.
Restrict active substances to substances with available hazard characterisations	Boolean	Restrict assessment group membership based on presence/abser of hazard characterisations.
Number of simulated days	Numeric	The total duration (in days) of the PBK model simulations.
Number of initial days skipped	Numeric	The initial period (or burn-in period, specified in days) that nee to be skipped when calculating the steady-state or peak internal exposures from the simulated time series.
Use PBK model parameter variability	Boolean	When specified, use parameter variability.
Number of events per day for the DERMAL dose	Numeric	The daily dose is administered in equal portions (dose / number events) per event.
Number of events per day for the INHALATION dose	Numeric	The daily dose is administered in equal portions (dose / number events) per event.
Number of events per day for the ORAL dose	Numeric	The daily dose is administered in equal portions (dose / number events) per event.
Specify the hours (events).	Numeric	Specify the hours on which exposure events are simulated. Allowed numbers are 1, 2, 3, 4,, 24 separated by spaces.
Specify event timings	Boolean	If checked, a specific sequence of hours on a day can be specifi on which exposure events are simulated. Otherwise the events a derived based on the number of exposure events per day.
Exposure events and dose generation method	ExposureEventsGeneration- Method	Generates doses and events for PBK model
PBK simulation method	PbkSimulationMethod	Specifies in which mode to run the PBK model
Expected life time in years	Numeric	The total expected lifetime (in years) of the PBK model.
Use bodyweight adjusted exposures	Boolean	Use bodyweight adjusted exposures in PBK model simulations.
Number of initial years skipped	Numeric	The initial period (or burn-in period, specified in years) that nee to be skipped when calculating the steady-state or peak internal exposures from the simulated time series.

Table 3.381: Calculation settings for module PBK model parametrisations.

Uncertainty settings

Table 3.382: Uncertainty settings for module PBK model parametrisations.

Name	Туре	Description
Resample PBK model parameter values	Boolean	Specifies whether PBK model parameter values are resampled.

PBK model parametrisations as data

Specify kinetic instances of PBK models and their parametrisations as data.

- *PBK model parametrisations data formats*
- PBK model parametrisations from data

Inputs used: Active substances PBK model definitions

Settings used

• Calculation Settings

Available kinetic models

Physiologically based kinetic (PBK) models, or kinetic models for short, are mathematical representations of the animal or human body aimed at describing and predicting the time course distribution of chemicals in tissues and organs. Those internal dose metrics can usefully replace external exposure dose in the derivation of the quantitative dose-response relationships and following risk assessments. PBK models can simulate both internal doses from exposure scenarios (forward dosimetry) and external dose from biomonitoring data (reverse dosimetry).

The following generic PBK models are currently implemented in MCRA:

- EuroMix generic PBK model (Tebby et al. (2020)).
- *bisphenol PBK model ETHZ* (Karrer et al. (2019)).

The MCRA interface allows to run PBK models for any number of days and supplies an option to skip an initial number of days (build-up phase PBK model) in the calculation of the internal exposure.

In the original bisphenol PBK model of Karrer et al. (2019), doses were applied on fixed time points and only for the first four days. For oral exposure, three dosings per day with t = 0, 6 and 12 h, for dermal exposure to PCPs and TP, two dosings per day with t = 0 and 12 h. The steady state was reached after dosing on four consecutive days. Therefore, the bisphenol PBK model described in *Karrer et al.* should be run for four days to reached steady state and the number of initial days to be skipped should be set to 0. Note, no dosing is applied from day 5th onwards.

These restrictions were relaxed in a new implementation of the bisphenol PBK model (General Model BPA Reimplementation). In this new version the number of days is unlimited. Dosings are specified through the user interface including the definition on the non-stationary period, see *settings PBK models*.

EuroMix Bisphenols PBPK model (v1)

EuroMix Bisphenols PBPK model by Karrer et al. (23 July 2018).

ld	Description	Unit
Oral	Oral exposure	nmoles
Dermal	Dermal exposure	nmoles
Inhalation	Inhalation exposure	nmoles

Table 3.383: Exposure routes (forcings)

ld	Description	ScalingFactor	Multiplication- Factor	Unit
CPlasmaOut	Concentration in plasma			nmol/L
CGonadOut	Concentration in gonads			nmol/L
AurinebpaOut	Cumulative excretion of BPA in urine			nmoles
AurinegOut	Cumulative excretion of BPA-g in urine			nmoles
AurineTotalOut	Cumulative excretion of BPA and metabolites in urine			nmoles

Table 3.384: Model outputs

ld	Description	Default	Unit	Туре
BW	Bodyweight		kg	Physiological
QCC	Cardiac output		L/min	Physiological
QgonadC	Fractional blood			Physiological
	flow to gonads			DI 1 1 1
QliverC	flow to liver			Physiological
QfatC	Fractional blood			Physiological
	flow to fat tissue			
QbrainC	Fractional blood			Physiological
	flow to brain			
QskinC	Fractional blood			Physiological
	flow to skin			
QmuscleC	Fractional blood			Physiological
	flow to gonads			
VplasmaC	Fractional volume of plasma			Physiological
VfatC	Fractional volume of			Physiological
	fat tissue			
VliverC	Fractional volume of			Physiological
	liver tissue			
VbrainC	Fractional volume of			Physiological
	brain tissue			
VskinC	Fractional volume of skin tissue			Physiological
VgonadC	Fractional volume of			Physiological
vgondue	gonads			i nysiologicui
VmuscleC	Fractional volume of			Physiological
	muscle tissue			
VrichC	Fractional volume of			Physiological
	richly perfused			
	tissue			
VbodygC	Distribution volume			Physiological
	of BPA-g			

ld	Description	Default	Unit	Туре
VbodysC	Distribution volume of BPA-s			Other
MW	Molecular weight		g/mol	MolecularWeight
pliver	Partition coefficient liver to blood			PartitionCoefficient
pfat	Partition coefficient fat to blood			PartitionCoefficient
pslow	Partition coefficient slowly perfused tissue to blood			PartitionCoefficient
prich	Partition coefficient richly perfused tissue to blood			PartitionCoefficient
pgonad	Partition coefficient gonads to blood			PartitionCoefficient
pbrain	Partition coefficient brain to blood			PartitionCoefficient
pskin	Partition coefficient skin to blood			PartitionCoefficient
geC	Gastric emptying		1/h/kg bw^-0.25	Physicochemical
k0C	Oral uptake from the stomach into the liver		1/h/kg bw^-0.25	Physicochemical
k1C	Oral uptake from the small intestine into the liver		1/h/kg bw^-0.25	Physicochemical
k4C	Fecal elimination from small intestine after oral administration		1/h/kg bw^-0.25	Physicochemical
kGlingC	Transport of glucuronide from enterocytes into serum		1/h/kg bw^-0.25	Physicochemical
kGIinsC	Transport of sulfate from enterocytes into serum		1/h/kg bw^-0.25	Physicochemical
kmgutg	Km of Glucuronidation in the gut		nM	Physicochemical
vmaxgutgC	Vmax of Glucuronidation in the gut		nmol/h/kg bw	Physicochemical
fgutg	Correction factor of glucuronidation in the gut			Physicochemical
kmguts	Km of Sulfation in the gut		nM	Physicochemical
vmaxgutsC	Vmax of Sulfation in the gut		nmol/h/kg bw	Physicochemical
fguts	Correction factor of sulfation in the gut			Physicochemical

Table 3.385 – continued from previous page

ld	Description	Default	Unit	Туре
met1g	Fraction of glucuronide in the liver taken up directly into serum (the rest undergoes EHR)			Physicochemical
met1s	Fraction of sulfate in the liver taken up directly into serum			Physicochemical
enterocytes	Sum of enterocytes weights in duodenum, jejunum and ileum		L	Physicochemical
kmliver	Km of Glucuronidation in the liver		nM	Physicochemical
vmaxliverC	Vmax of Glucuronidation in the liver		nmol/h/g liver	Physicochemical
fliverg	Correction factor of glucuronidation in the liver			Physicochemical
kmlivers	Km of Sulfation in the liver		nM	Physicochemical
vmaxliversC	Vmax of Sulfation in the liver		nmol/h/g liver	Physicochemical
flivers	Correction factor of sulfation in the liver			Physicochemical
EHRtime	Time until EHR		h	Physicochemical
EHRrateC	EHR of glucuronide		1/h/kg bw^-0.25	Physicochemical
k4C_IV	Fecal elimination of glucuronide from the EHR compartment		1/h/kg bw^-0.25	Physicochemical
kurinebpaC	Clearance, urine excretion of parent compound		L/h/kg bw^0.75	Physicochemical
kurinebpagC	Clearance, urine excretion of glucuronide		L/h/kg bw^0.75	Physicochemical
kurinebpasC	Clearance, urine excretion of sulfate		L/h/kg bw^0.75	Physicochemical
vreabsorptiong- C	Vmax for renal reabsorption of glucuronide		nmol/h/kg bw^0.75	Physicochemical
vreabsorptionsC	Vmax for renal reabsorption of sulfate		nmol/h/kg bw^0.75	Physicochemical
kreabsorptiong	Km for renal reabsorption of glucuronide		nM	Physicochemical
kreabsorptions	Km for renal reabsorption of sulfate		nM	Physicochemical

Table 3.385 – continued from previous page

ld	Description	Default	Unit	Туре
kenterobpagC	EHR of parent compound due to biliary excretion of glucuronide		1/h/kg bw^-0.25	Physicochemical
kenterobpasC	EHR of parent compound due to biliary excretion of sulfate		1/h/kg bw^-0.25	Physicochemical
D_o	oral dose		ng/kg bw/dosing	Other
dose_O	oral dose		nmol/kg bw/dosing	Other
EoA_O	Extent of oral absorption			Physiological
uptake_O	amount of oral uptake		nmol/dosing	Other
period_O	uptake period		h	Other
koa	uptake rate		nmol/h	Other
t0_O	time point at which dosing starts		h	Other
t1_0	time point at which dosing ends		h	Other
D_d	dermal dose from thermal paper (TP)		ng/kg bw/dosing	Other
EoA_D	Extent of dermal absorption from TP		C	Physiological
dose_D	dermal dose from TP		nmol/kg bw/dosing	Other
aHL_D	Half-life for dermal penetration		h	Other
uptake_D	amount of dermal uptake from TP		nmol/dosing	Other
period_D	Uptake period dermal exposure from TP		h	Other
kda	Uptake rate of dermal exposure from TP		nmol/h	Other
t0_D	Time points at which dermal dosing from TP starts		h	Other
t1_D	Time points at which dermal dosing from TP ends		h	Other
D_d2	Dermal dose from PCPs		ng/kg bw/dosing	Other
EoA_D2	Extent of dermal absorption from PCPs		c	Physiological
dose_D2	Dermal dose from PCPs		nmol/kg bw/dosing	Other
aHL_D2	Half-life for dermal penetration from PCPs		h	Other

Table 3.385 – continued from previous page

ld	Description	Default	Unit	Туре
uptake_D2	amount of dermal uptake from PCPs		nmol/dosing	Other
period_D2	Uptake period dermal exposure from PCPs		h	Other
kda2	uptake rate of dermal exposure from PCPs		nmol/h	Other
t0_D2	Time points at which dermal dosing from PCPs starts		h	Other
t1_D2	time points at which dermal dosing from TP ends		h	Other
QC	Cardiac output		L/h	Other
Qfat	Blood flow to the fat tissue		L/h	Other
Qliver	Blood flow to the liver tissue		L/h	Other
Qgonad	Blood flow to the gonads		L/h	Other
Qbrain	Blood flow to the brain		L/h	Other
Qskin	Blood flow to the skin tissue		L/h	Other
Qslow	Blood flow to the slowly perfused tissue		L/h	Other
Qrich	Blood flow to the richly perfused tissue		L/h	Other
Vliver	Volume of the liver		L	Other
Vfat	Volume of the fat tissue		L	Other
Vgonad	Volume of the gonads		L	Other
Vplasma	Volume of the plasma		L	Other
Vbrain	Volume of the brain		L	Other
Vskin	Volume of the skin		L	Other
Vslow	Volume of the slowly perfused tissue		L	Other
Vrich	richly perfused tissue		L	Other
Vbodyg	Volume of the distribution for BPAG		L	Other
Vbodys	Volume of the distribution for BPAS		L	Other
BW075	BW^0.75		kg^0.75	Other
BW025	BW^0.25		kg^0.25	Other
vmaxlivers- Cnew	scaled Vmax of Sulfation of BPA in the liver		nmol/h/kg bw^0.75	Other

Table 3.385 - continued from previous page	Э
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ld	Description	Default	Unit	Туре
vmaxliverCnew	scaled Vmax of Glucuronidation of BPA in the liver		nmol/h/kg bw^0.75	Other
vmaxgutgCnew	scaled Vmax of Glucuronidation of BPA in the gut		nmol/h/kg bw^0.75	Other
vreabsorptiong	scaled vmax of renal resorption of BPAG		nmol/h	Other
vreabsorptions	scaled vmax of renal resorption of BPAS		nmol/h	Other
EHRrate	scaled EHR of BPAG		1/h	Other
k0	scaled Uptake of BPA from the stomach into the liver		1/h	Other
ge	scaled Gastric emptying of BPA		1/h	Other
k1	scaled Uptake of BPA from small intestine into the liver		1 <i>/</i> h	Other
k4	scaled Fecal excretion of BPA after oral administration from small intestine		1/h	Other
k4_IV	scaled Fecal excretion of BPAG from the EHR compartment		1/h	Other
vmaxliver	rescaled and corrected vmax of BPA glucuronidation in the liver		nmol/h	Other
kGIing	scaled Uptake of BPAG from small intestine into serum		1/h	Other
met2g	Fraction of BPAG formed subject to EHR			Other
met2s	Fraction of BPAS formed subject to EHR			Other
kurinebpa	scaled Clearance of BPA via urine		L/h	Other
kurinebpag	scaled Clearance of BPAg via urine		L/h	Other
kurinebpas	scaled Clearance of BPAs via urine		L/h	Other
vmaxlivers	rescaled and corrected vmax of BPA sulfation in the liver		nmol/h	Other

Table 3.385 – continued from previous page

ld	Description	Default	Unit	Туре
kGIins	scaled Uptake of BPAS from small intestine into serum		1/h	Other
vmaxgutg	rescaled and corrected vmax of BPA glucuronidation in the gut		nmol/h	Other
vmaxguts	rescaled and corrected vmax of BPA sulfation in the gut		nmol/h	Other
kenterobpag	scaled EHR of BPA due to biliary excretion of BPAG		1/h	Other
kenterobpas	scaled EHR of BPA due to biliary excretion of BPAS		1/h	Other
t0_D1_day1	time points at which dermal dosing from Thermal paper starts		h	Other
t0_D2_day1	time points at which dermal dosing from Thermal paper starts		h	Other
t0_D1_day2	time points at which dermal dosing from Thermal paper starts		h	Other
t0_D2_day2	time points at which dermal dosing from Thermal paper starts		h	Other
t0_D1_day3	time points at which dermal dosing from Thermal paper starts		h	Other
t0_D2_day3	time points at which dermal dosing from Thermal paper starts		h	Other
t0_D1_day4	time points at which dermal dosing from Thermal paper starts		h	Other
t0_D2_day4	time points at which dermal dosing from Thermal paper starts		h	Other
t0_D21_day1	time points at which dermal dosing from PCPs starts		h	Other
t0_D22_day1	time points at which dermal dosing from PCPs starts		h	Other
t0_D21_day2	time points at which dermal dosing from PCPs starts		h	Other
t0_D22_day2	time points at which dermal dosing from PCPs starts		h	Other

Table 3.385 – continued from previous bade	Table	e 3.385 -	continued fr	om previous	page
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ld	Description	Default	Unit	Туре
t0_D21_day3	time points at which dermal dosing from PCPs starts		h	Other
t0_D22_day3	time points at which dermal dosing from PCPs starts		h	Other
t0_D21_day4	time points at which dermal dosing from PCPs starts		h	Other
t0_D22_day4	time points at which dermal dosing from PCPs starts		h	Other
t0_O1_day1	time points at which oral dosing starts		h	Other
t0_O2_day1	time points at which oral dosing starts		h	Other
t0_O3_day1	time points at which oral dosing starts		h	Other
t0_O1_day2	time points at which oral dosing starts		h	Other
t0_O2_day2	time points at which oral dosing starts		h	Other
t0_O3_day2	time points at which oral dosing starts		h	Other
t0_O1_day3	time points at which oral dosing starts		h	Other
t0_O2_day3	time points at which oral dosing starts		h	Other
t0_O3_day3	time points at which oral dosing starts		h	Other
t0_O1_day4	time points at which oral dosing starts		h	Other
t0_O2_day4	time points at which oral dosing starts		h	Other
t0_O3_day4	time points at which oral dosing starts		h	Other
ksiLiver	Ksi of glucuronidation in liver		nM	Other
ksiGut	Ksi of glucuronidation in gut		nM	Other
age	age	30		Other
gender	gender	0		Other
QCC_adult_f	QCC_adult_f			Other
Qgonad- C_adult_f	QgonadC_adult_f			Other
QliverC_adult_f	QliverC_adult_f			Other
QtatC_adult_f	QtatC_adult_f			Other
Qbrain- C_adult_f	QbrainC_adult_f			Other
QskinC_adult_f	QskinC_adult_f			Other
Qmuscle- C_adult_f	QmuscleC_adult_f			Other

Table 3.385 - continued from previous page

ld	Description	Default	Unit	Туре
Vplasma-	VplasmaC_adult f		[Other
C adult f	1			
VfatC adult f	VfatC adult f			Other
VliverC_adult_f	VliverC_adult_f			Other
Vbrain-	VbrainC_adult_f			Other
C_adult_f				
VskinC_adult_f	VskinC_adult_f			Other
Vgonad-	VgonadC_adult_f			Other
C_adult_f				
Vmuscle-	VmuscleC_adult_f			Other
C_adult_f				
VrichC_adult_f	VrichC_adult_f			Other
Vbodyg-	VbodygC_adult_f			Other
C_adult_f				
Vbodys-	VbodysC_adult_f			Other
C_adult_f				
QCC_adult_m	QCC_adult_m			Other
Qgonad-	QgonadC_adult_m			Other
C_adult_m				
Qliver-	QliverC_adult_m			Other
C_adult_m				
QfatC_adult_m	QfatC_adult_m			Other
Qbrain-	QbrainC_adult_m			Other
C_adult_m				
Qskin-	QskinC_adult_m			Other
C_adult_m				
Qmuscle-	QmuscleC_adult_m			Other
C_adult_m				
Vplasma-	VplasmaC_adult_m			Other
C_adult_m				
VfatC_adult_m	VfatC_adult_m			Other
Vliver-	VliverC_adult_m			Other
C_adult_m				
Vbrain-	VbrainC_adult_m			Other
C_adult_m				
Vskin-	VskinC_adult_m			Other
C_adult_m				
Vgonad-	VgonadC_adult_m			Other
C_adult_m				
Vmuscle-	VmuscleC_adult_m			Other
C_adult_m				
Vrich-	VrichC_adult_m			Other
C_adult_m				
Vbodyg-	VbodygC_adult_m			Other
C_adult_m				
Vbodys-	VbodysC_adult_m			Other
C_adult_m				
QC-	QCC_adolescent_f			Other
C_adolescent_f	0			
Qgonad-	Qgonad-			Other
C_adolescent_f	C_adolescent_f			
Qliver-	Qliver-			Other
C_adolescent_f	C_adolescent_f			0.1
Qfat-	QfatC_adolescent_f			Other
C_adolescent_f				

Table 3.385 – continued from previous page
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ld	Description	Default	Unit	Туре
Qbrain-	Qbrain-			Other
C adolescent f	C adolescent f			
Qskin-	Qskin-			Other
C_adolescent_f	C_adolescent_f			
Qmuscle-	Qmuscle-			Other
C_adolescent_f	C_adolescent_f			
Vplasma-	Vplasma-			Other
C adolescent f	C adolescent f			
Vfat-	VfatC adolescent f			Other
C_adolescent_f				
Vliver-	Vliver-			Other
C_adolescent_f	C_adolescent_f			
Vbrain-	Vbrain-			Other
C_adolescent_f	C_adolescent_f			
Vskin-	Vskin-			Other
C_adolescent_f	C_adolescent_f			
Vgonad-	Vgonad-			Other
C_adolescent_f	C_adolescent_f			
Vmuscle-	Vmuscle-			Other
C_adolescent_f	C_adolescent_f			
Vrich-	Vrich-			Other
C_adolescent_f	C_adolescent_f			
Vbodyg-	Vbodyg-			Other
C_adolescent_f	C_adolescent_f			
Vbodys-	Vbodys-			Other
C_adolescent_f	C_adolescent_f			
QC-	QCC_adolescent_m			Other
C_adolescent_m				
Qgonad-	Qgonad-			Other
C_adolescent_m	C_adolescent_m			
Qliver-	Qliver-			Other
C_adolescent_m	C_adolescent_m			
Qfat-	Qfat-			Other
C_adolescent_m	C_adolescent_m			
Qbrain-	Qbrain-			Other
C_adolescent_m	C_adolescent_m			
Qskin-	Qskin-			Other
C_adolescent_m	C_adolescent_m			
Qmuscle-	Qmuscle-			Other
C_adolescent_m	C_adolescent_m			
Vplasma-	Vplasma-			Other
C_adolescent_m	C_adolescent_m			
Vfat-	VfatC_adolescent_m			Other
C_adolescent_m				
Vliver-	Vliver-			Other
C_adolescent_m	C_adolescent_m			
Vbrain-	Vbrain-			Other
C_adolescent_m	C_adolescent_m			
Vskin-	Vskin-			Other
C_adolescent_m	C_adolescent_m			
Vgonad-	Vgonad-			Other
C_adolescent_m	C_adolescent_m			
Vmuscle-	Vmuscle-			Other
C_adolescent_m	C_adolescent_m			

Table 3.385 - continued from previous page

ld	Description	Default	Unit	Туре
Vrich-	Vrich-			Other
C adolescent m	C adolescent m			
Vbodyg-	Vbodyg-			Other
C_adolescent_m	C_adolescent_m			
Vbodys-	Vbodys-			Other
C_adolescent_m	C_adolescent_m			
QCC_child_f	QCC_child_f			Other
Qgonad-	QgonadC_child_f			Other
C_child_f				
QliverC_child_f	QliverC_child_f			Other
QfatC_child_f	QfatC_child_f			Other
Qbrain-	QbrainC_child_f			Other
C_child_f				
QskinC_child_f	QskinC_child_f			Other
Qmuscle-	QmuscleC_child_f			Other
C_child_f				
Vplasma-	VplasmaC_child_f			Other
C_child_f				
VfatC_child_f	VfatC_child_f			Other
VliverC_child_f	VliverC_child_f			Other
Vbrain-	VbrainC_child_f			Other
C_child_f				0.1
VskinC_child_f	VskinC_child_f			Other
Vgonad-	VgonadC_child_f			Other
C_child_f	Variate Calif.			Other
Vinuscie-	vinuscieC_cinita_i			Other
C_cillu_1 VrichC_child_f	VrichC child f			Other
Vhodva	VhodygC child f			Other
C child f	voouyge_ennu_r			Other
Vbodys-	VbodysC child f			Other
C child f	voodyse_ennd_r			other
OCC child m	OCC child m			Other
Ogonad-	OgonadC child m			Other
C child m	C 0*******_***			
Oliver-	OliverC child m			Other
C child m	、			
QfatC_child_m	QfatC_child_m			Other
Qbrain-	QbrainC_child_m			Other
C_child_m				
Qskin-	QskinC_child_m			Other
C_child_m				
Qmuscle-	QmuscleC_child_m			Other
C_child_m				
Vplasma-	VplasmaC_child_m			Other
C_child_m				
VfatC_child_m	VfatC_child_m			Other
Vliver-	VliverC_child_m			Other
C_child_m				
Vbrain-	VbrainC_child_m			Other
C_child_m				0.1
Vskin-	VskinC_child_m			Other
C_child_m	V 10 1 11			0.1
C child m	vgonauC_cnild_m			Oulei

Table 3.385 – continued from previous page

ld	Description	Default	Unit	Туре
Vmuscle- C_child_m	VmuscleC_child_m			Other
Vrich- C_child_m	VrichC_child_m			Other
Vbodyg- C_child_m	VbodygC_child_m			Other
Vbodys- C_child_m	VbodysC_child_m			Other

 Table 3.385 – continued from previous page

Model aliases: EuroMix_Bisphenols_PBPK_model_V1, PBPKModel_BPA.

EuroMix Bisphenols PBPK model (v2)

EuroMix Bisphenols PBPK model by Karrer et al. (2019).

ld	Description	Unit
Oral	Oral exposure	nmoles
Dermal	Dermal exposure	nmoles
Inhalation	Inhalation exposure	nmoles

Table 3.387: Model outputs

ld	Description	ScalingFactor	Multiplication- Factor	Unit
CPlasmaOut	Concentration in plasma			nmol/L
CGonadOut	Concentration in gonads			nmol/L
AurinebpaOut	Cumulative excretion of BPA in urine			nmoles
AurinegOut	Cumulative excretion of BPA-g in urine			nmoles
AurineTotalOut	Cumulative excretion of BPA and metabolites in urine			nmoles

Table 3.388: Model parameters

ld	Description	Default	Unit	Туре
BW	Bodyweight		kg	Physiological
QCC	Cardiac output		L/min	Physiological
QgonadC	Fractional blood flow to gonads			Physiological
QliverC	Fractional blood flow to liver			Physiological
QfatC	Fractional blood flow to fat tissue			Physiological

ld	Description	Default	Unit	Туре
QbrainC	Fractional blood flow to brain			Physiological
QskinC	Fractional blood flow to skin			Physiological
QmuscleC	Fractional blood flow to gonads			Physiological
VplasmaC	Fractional volume of plasma			Physiological
VfatC	Fractional volume of fat tissue			Physiological
VliverC	Fractional volume of liver tissue			Physiological
VbrainC	Fractional volume of brain tissue			Physiological
VskinC	Fractional volume of skin tissue			Physiological
VgonadC	Fractional volume of gonads			Physiological
VmuscleC	Fractional volume of muscle tissue			Physiological
VrichC	Fractional volume of richly perfused tissue			Physiological
VbodygC	Distribution volume of BPA-g			Physiological
VbodysC	Distribution volume of BPA-s			Other
pliver	Partition coefficient liver to blood			PartitionCoefficient
pfat	Partition coefficient fat to blood			PartitionCoefficient
pslow	Partition coefficient slowly perfused tissue to blood			PartitionCoefficient
prich	Partition coefficient richly perfused tissue to blood			PartitionCoefficient
pgonad	Partition coefficient gonads to blood			PartitionCoefficient
pbrain	Partition coefficient brain to blood			PartitionCoefficient
pskin	Partition coefficient skin to blood			PartitionCoefficient
geC	Gastric emptying		1/h/kg bw^-0.25	Physicochemical
k0C	Oral uptake from the stomach into the liver		1/h/kg bw^-0.25	Physicochemical
k1C	Oral uptake from the small intestine into the liver		1/h/kg bw^-0.25	Physicochemical
k4C	Fecal elimination from small intestine after oral administration		1/h/kg bw^-0.25	Physicochemical

Table	3.388 -	continued from	previous page

ld	Description	Default	Unit	Туре
kGIingC	Transport of glucuronide from enterocytes into serum		1/h/kg bw^-0.25	Physicochemical
kGIinsC	Transport of sulfate from enterocytes into serum		1/h/kg bw^-0.25	Physicochemical
kmgutg	Km of Glucuronidation in the gut		nM	Physicochemical
vmaxgutgC	Vmax of Glucuronidation in the gut		nmol/h/kg bw	Physicochemical
fgutg	Correction factor of glucuronidation in the gut			Physicochemical
kmguts	Km of Sulfation in the gut		nM	Physicochemical
vmaxgutsC	Vmax of Sulfation in the gut		nmol/h/kg bw	Physicochemical
fguts	Correction factor of sulfation in the gut			Physicochemical
metlg	Fraction of glucuronide in the liver taken up directly into serum (the rest undergoes EHR)			Physicochemical
met1s	Fraction of sulfate in the liver taken up directly into serum			Physicochemical
enterocytes	Sum of enterocytes weights in duodenum, jejunum and ileum		L	Physicochemical
kmliver	Km of Glucuronidation in the liver		nM	Physicochemical
vmaxliverC	Vmax of Glucuronidation in the liver		nmol/h/g liver	Physicochemical
fliverg	Correction factor of glucuronidation in the liver			Physicochemical
kmlivers	Km of Sulfation in the liver		nM	Physicochemical
vmaxliversC	Vmax of Sulfation in the liver		nmol/h/g liver	Physicochemical
flivers	Correction factor of sulfation in the liver			Physicochemical
EHRtime	Time until EHR occurs		h	Physicochemical
EHRrateC	EHR of glucuronide		1/h/kg bw^-0.25	Physicochemical

Table 3.388 – continued from previous page

ld	Description	Default	Unit	Туре
k4C_IV	Fecal elimination of glucuronide from the EHR compartment		1/h/kg bw^-0.25	Physicochemical
kurinebpaC	Clearance, urine excretion of parent compound		L/h/kg bw^0.75	Physicochemical
kurinebpagC	Clearance, urine excretion of glucuronide		L/h/kg bw^0.75	Physicochemical
kurinebpasC	Clearance, urine excretion of sulfate		L/h/kg bw^0.75	Physicochemical
vreabsorptiong- C	Vmax for renal reabsorption of glucuronide		nmol/h/kg bw^0.75	Physicochemical
vreabsorptionsC	Vmax for renal reabsorption of sulfate		nmol/h/kg bw^0.75	Physicochemical
kreabsorptiong	Km for renal reabsorption of glucuronide		nM	Physicochemical
kreabsorptions	Km for renal reabsorption of sulfate		nM	Physicochemical
kenterobpagC	EHR of parent compound due to biliary excretion of glucuronide		1/h/kg bw^-0.25	Physicochemical
kenterobpasC	EHR of parent compound due to biliary excretion of sulfate		1/h/kg bw^-0.25	Physicochemical
koa	uptake rate		nmol/h	Other
EoA_D	Extent of dermal absorption from TP			Physiological
aHL_D	Half-life for dermal penetration		h	Other
kda	Uptake rate of dermal exposure from TP		nmol/h	Other
EoA_D2	Extent of dermal absorption from PCPs			Physiological
aHL_D2	Half-life for dermal penetration from PCPs		h	Other
kda2	Uptake rate of dermal exposure from PCPs		nmol/h	Other
QC	Cardiac output		L/h	Other
Qfat	Blood flow to the fat tissue		L/h	Other
Qliver	Blood flow to the liver tissue		L/h	Other
Qgonad	Blood flow to the gonads		L/h	Other

Table 3.388 – continued from previous page

ld	Description	Default	Unit	Туре
Qbrain	Blood flow to the brain		L/h	Other
Qskin	Blood flow to the skin tissue		L/h	Other
Qslow	Blood flow to the slowly perfused tissue		L/h	Other
Qrich	Blood flow to the richly perfused tissue		L/h	Other
Vliver	Volume of the liver		L	Other
Vfat	Volume of the fat		L	Other
	tissue			
Vgonad	Volume of the gonads		L	Other
Vplasma	Volume of the plasma		L	Other
Vbrain	Volume of the brain		L	Other
Vskin	Volume of the skin		L	Other
Vslow	Volume of the slowly perfused tissue		L	Other
Vrich	richly perfused tissue		L	Other
Vbodyg	Volume of the distribution for BPAG		L	Other
Vbodys	Volume of the distribution for BPAS		L	Other
vreabsorptiong	scaled vmax of renal resorption of BPAG		nmol/h	Other
vreabsorptions	scaled vmax of renal resorption of BPAS		nmol/h	Other
EHRrate	scaled EHR of BPAG		1/h	Other
k0	scaled Uptake of BPA from the stomach into the liver		1/h	Other
ge	scaled Gastric emptying of BPA		1/h	Other
k1	scaled Uptake of BPA from small intestine into the liver		1/h	Other
k4	scaled Fecal excretion of BPA after oral administration from small intestine		1/h	Other
k4_IV	scaled Fecal excretion of BPAG from the EHR compartment		1/h	Other

Table 3.388 – continued from previous page	Table	3.388 -	continued	from	previous	page
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ld	Description	Default	Unit	Туре
vmaxliver	rescaled and corrected vmax of BPA glucuronidation in the liver		nmol/h	Other
kGIing	scaled Uptake of BPAG from small intestine into serum		1/h	Other
met2g	Fraction of BPAG formed subject to EHR			Other
met2s	Fraction of BPAS formed subject to EHR			Other
kurinebpa	scaled Clearance of BPA via urine		L/h	Other
kurinebpag	scaled Clearance of BPAg via urine		L/h	Other
kurinebpas	scaled Clearance of BPAs via urine		L/h	Other
vmaxlivers	rescaled and corrected vmax of BPA sulfation in the liver		nmol/h	Other
kGIins	scaled Uptake of BPAS from small intestine into serum		1/h	Other
vmaxgutg	rescaled and corrected vmax of BPA glucuronidation in the gut		nmol/h	Other
vmaxguts	rescaled and corrected vmax of BPA sulfation in the gut		nmol/h	Other
kenterobpag	scaled EHR of BPA due to biliary excretion of BPAG		1/h	Other
kenterobpas	scaled EHR of BPA due to biliary excretion of BPAS		1/h	Other
ksiLiver	Ksi of glucuronidation in liver		nM	Other
ksiGut	Ksi of glucuronidation in gut		nM	Other
age	age	30		Other
gender	gender	0		Other
QCC_adult_f	QCC_adult_f			Other
Qgonad- C_adult_f	QgonadC_adult_f			Other
QliverC_adult_f	QliverC_adult_f			Other
QtatC_adult_f	QtatC_adult_f			Other
C_adult_f	QbrainC_adult_f			Other

Table	3.388 -	continued	from	previous	page

ld	Description	Default	Unit	Туре
QskinC_adult_f	QskinC_adult_f			Other
Qmuscle-	QmuscleC_adult_f			Other
C_adult_f				
Vplasma-	VplasmaC_adult_f			Other
C_adult_f				
VfatC_adult_f	VfatC_adult_f			Other
VliverC_adult_f	VliverC_adult_f			Other
Vbrain-	VbrainC_adult_f			Other
C_adult_f				
VskinC_adult_f	VskinC_adult_f			Other
Vgonad-	VgonadC_adult_f			Other
C_adult_f				
Vmuscle-	VmuscleC_adult_f			Other
C_adult_f				
VrichC_adult_f	VrichC_adult_f			Other
Vbodyg-	VbodygC_adult_f			Other
C_adult_f				
Vbodys-	VbodysC_adult_f			Other
C_adult_f				
QCC_adult_m	QCC_adult_m			Other
Qgonad-	QgonadC_adult_m			Other
C_adult_m	~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~			
Qliver-	QliverC_adult_m			Other
C_adult_m				0.1
QfatC_adult_m	QfatC_adult_m			Other
Qbrain-	QbrainC_adult_m			Other
C_adult_m	01:011			0.1
Qskin-	QskinC_adult_m			Other
C_adult_m	On I C I k			Other
Qmuscle-	QmuscleC_adult_m			Other
C_adult_m	ValormoC odult m			Other
v piasilia-	vpiasinaC_aduit_in			Other
C_aduit_iii VfatC_adult_m	VfetC adult m			Other
Vliver	VliverC adult m			Other
C adult m	viiverC_aduit_iii			Oulei
Vbrain-	VbrainC adult m			Other
C adult m	voranic_aduit_iii			Oulei
Vskin-	VskinC adult m			Other
C adult m	v skine_uduit_m			other
Vgonad-	VoonadC adult m			Other
C adult m	·gonuce_udun_m			other
Vmuscle-	VmuscleC adult m			Other
C adult m	· musere c_waare_m			
Vrich-	VrichC adult m			Other
C adult m				
Vbodyg-	VbodygC adult m			Other
C_adult_m				
Vbodys-	VbodysC_adult_m			Other
C_adult_m	• • –			
QC-	QCC_adolescent_f			Other
C_adolescent_f				
Qgonad-	Qgonad-			Other
C_adolescent_f	C_adolescent_f			

 Table 3.388 - continued from previous page

ld	Description	Default	Unit	Туре
Qliver-	Qliver-			Other
C adolescent f	C adolescent f			
Qfat-	QfatC adolescent f			Other
C_adolescent_f				
Qbrain-	Qbrain-			Other
C_adolescent_f	C_adolescent_f			
Qskin-	Qskin-			Other
C_adolescent_f	C_adolescent_f			
Qmuscle-	Qmuscle-			Other
C_adolescent_f	C_adolescent_f			
Vplasma-	Vplasma-			Other
C_adolescent_f	C_adolescent_f			
Vfat-	VfatC_adolescent_f			Other
C_adolescent_f				
Vliver-	Vliver-			Other
C_adolescent_f	C_adolescent_f			
Vbrain-	Vbrain-			Other
C_adolescent_f	C_adolescent_f			
Vskin-	Vskin-			Other
C_adolescent_f	C_adolescent_f			
Vgonad-	Vgonad-			Other
C_adolescent_f	C_adolescent_f			
Vmuscle-	Vmuscle-			Other
C_adolescent_f	C_adolescent_f			
Vrich-	Vrich-			Other
C_adolescent_f	C_adolescent_f			
Vbodyg-	Vbodyg-			Other
C_adolescent_f	C_adolescent_f			
Vbodys-	Vbodys-			Other
C_adolescent_f	C_adolescent_f			
QC-	QCC_adolescent_m			Other
C_adolescent_m	0 1			0.1
Qgonad-	Qgonad-			Other
C_adolescent_m	C_adolescent_m			0.1
Qliver-	Qliver-			Other
C_adolescent_m	C_adolescent_m			Other
Qiat-	Qrat-			Other
C_adolescent_m	C_adolescent_m			Other
Quialli-	Quianii-			Other
C_adolescent_III	C_adolescent_III			Other
Qskill- C adalasaant m	QSKIII-			Other
C_autorescent_iii	C_autorescent_in			Other
C adolescent m	C adolescent m			Oulei
Vplasma	Vnlasma			Other
C adolescent m	C adolescent m			Oulei
Vfat-	VfatC adolescent m			Other
C adolescent m				Guidi
Vliver-	Vliver-			Other
C adolescent m	C adolescent m			Guior
Vbrain-	Vhrain-			Other
C adolescent m	C adolescent m			Guior
Vskin-	Vskin-			Other
C_adolescent_m	C_adolescent_m			

Table 3.388 – continued from previous page

ld	Description	Default	Unit T	уре
Vgonad-	Vgonad-		C	Other
C_adolescent_m	C_adolescent_m			
Vmuscle-	Vmuscle-		C	Other
C_adolescent_m	C_adolescent_m			
Vrich-	Vrich-		С	Other
C_adolescent_m	C_adolescent_m			
Vbodyg-	Vbodyg-		С	Other
C_adolescent_m	C_adolescent_m			
Vbodys-	Vbodys-		С	Other
C_adolescent_m	C_adolescent_m			
QCC_child_f	QCC_child_f		C	Other
Qgonad- C shild f	QgonadC_child_f		C	Other
OliverC child f	OliverC child f		0)ther
QfatC child f	OfetC child f			Other
Obrain-	ObrainC child f			Other
C_child_f	Quranic_cinid_r		C C	Juici
QskinC_child_f	QskinC_child_f		С	Other
Qmuscle-	QmuscleC_child_f		C	Other
C_child_f				
Vplasma- C_child_f	VplasmaC_child_f		C	Other
VfatC child f	VfatC child f		C)ther
VliverC child f	VliverC child f			Other
VhverC_ennu_r	VhrainC child f			Other
C_child_f	voranie_ennu_r			Juici
VskinC_child_f	VskinC_child_f		С	Other
Vgonad- C. child. f	VgonadC_child_f		С	Other
Vmuscle-	VmuscleC child f		C	Other
C child f			C C	
VrichC child f	VrichC child f		C	Other
Vbodyg-	VbodygC child f		C	Other
C_child_f			-	
Vbodys-	VbodysC_child_f		C	Other
C_child_f				
QCC_child_m	QCC_child_m		С	Other
Qgonad-	QgonadC_child_m		С	Other
C_child_m				
Qliver-	QliverC_child_m		С	Other
C_child_m				
QfatC_child_m	QfatC_child_m		С	Other
Qbrain-	QbrainC_child_m		С	Other
C_child_m				
Qskin-	QskinC_child_m		C	Other
C_child_m				
Qmuscle-	QmuscleC_child_m		С	Other
C_child_m				
Vplasma-	VplasmaC_child_m		С	Other
C_child_m				
VfatC_child_m	VfatC_child_m		C	Other
Vliver-	VliverC_child_m		C	Other
C_child_m			-	NI
v brain-	vbrainC_child_m		C	Jther
C_cniia_m				

Table 3.388 – continued from previous page

ld	Description	Default	Unit	Туре
Vskin-	VskinC_child_m	[Other
C_child_m				
Vgonad-	VgonadC_child_m			Other
C_child_m				
Vmuscle-	VmuscleC_child_m			Other
C_child_m				
Vrich-	VrichC_child_m			Other
C_child_m				
Vbodyg-	VbodygC_child_m			Other
C_child_m				
Vbodys-	VbodysC_child_m			Other
C_child_m				

Table 3.388 – continued from previous p	age
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Model aliases: EuroMix_Bisphenols_PBPK_model_V2, PBPKModel_BPA_Reimplementation.

EuroMix Generic PBTK model (v5)

Cosmos version 5 (adapted 9/11/2018)

	1	
ld	Description	Unit
Oral	Oral exposure	mmoles
Dermal	Dermal exposure	mmoles
Inhalation	Inhalatory exposure	mmoles

Table 3.389: Exposure routes (forcings)

Table	3.390:	Model	outputs
raute	5.570.	mouci	outputs

ld	Description	ScalingFactor	Multiplication- Factor	Unit
CVen	Venous blood	scVBlood	0.66667	mM
CArt	Arterial blood	scVBlood	0.33333	mМ
CFat	Fat tissues	scVFat		mМ
CPoor	Muscle tissues			mМ
CRich	Viscera	scVRich		mМ
CLiver	Liver	scVLiver		mМ
CSkin_u	Viable skin, unexposed			mM
CSkin_e	Viable skin, exposed	BSA, Height_vs, fsA_exposed		mM
CSkin_sc_u	Skin stratum corneum, unexposed			mM
CSkin_sc_e	Skin stratum corneum, exposed	BSA, Height_vs, fsA_exposed		mМ

Table 3.391: Model	parameters
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ld	Description	Default	Unit	Туре
BM BSA	Body mass Body skin surface area		kg dm2	Physiological Physiological

ld	Description	Default	Unit	Туре
scVFat	Fat as fraction of total body volume			Physiological
scVRich	Richly perfused tissues (viscera) as fraction of total body volume			Physiological
scVLiver	Liver as fraction of total body volume			Physiological
scVBlood	Blood as fraction of total body volume			Physiological
Height_sc	Skin thickness		decimeter	Physiological
Height_vs	Viable skin			Physiological
scFBlood	Total blood flow per unit mass		L/h/kg	Physiological
scFFat	Fat fraction of total blood flow going to compartments			Physiological
scFPoor	Poorly perfused tissues (muscles) fraction of total blood flow going to compartments			Physiological
scFLiver	Liver fraction of total blood flow going to compartments			Physiological
scFSkin	Skin fraction of total blood flow going to compartments			Physiological
Falv	Alveolar ventilation rate		L/h	Physiological
mic	Microsomal proteins content		mg/gr liver	Physiological
PCAir	Partition coefficient: blood over air			PartitionCoefficient
log_PCFat	Scaled parameter, partition coefficient: fat over blood			PartitionCoefficient
log_aPoor	Scaled parameter, partition coefficient: muscle over blood (poorly perfused tissue)			PartitionCoefficient
log_aRich	Scaled parameter, partition coefficient: viscera over blood (richly perfused tissue)			PartitionCoefficient
log_aLiver	Scaled parameter, partition coefficient: liver over blood			PartitionCoefficient
log_aSkin	Scaled parameter, partition coefficient: viable skin / blood			PartitionCoefficient

 Table 3.391 – continued from previous page

ld	Description	Default	Unit	Туре
log_aSkin_sc	Scaled parameter, partition coefficient: viable skin / stratum corneum			PartitionCoefficient
Kp_sc_vs	Diffusion rate from stratum corneum to viable skin		decimeter/h	Physicochemical
Ke	Renal excretion rate		L/h	Physicochemical
Michaelis	Flag for Michaelis-Menten vs linear metabolism (0 = linear)			Physicochemical
Vmax	Maximum rate of metabolism		mmoles/h/L liver	Physicochemical
Km	Michaelis-Menten constant		mM	Physicochemical
CLH	Hepatic clearance			Physicochemical
fup	Unbound fraction in blood			Physicochemical
Frac	Fraction absorbed by the gut			Physicochemical
kGut	Oral 1st order absorption rate constant		1/h	Physicochemical
Cinh	Inhalation			Other
Tinh	Inhalation duration			Other
OralDose			mmol	Other
DermalDose			mmol	Other
fSA_exposed	Fraction of skin surface area actually exposed			Physicochemical
FBlood	Blood flow			Other
FFat	Scaled parameters			Other
FPoor	Scaled parameters			Other
FRich	Scaled parameters			Other
FLiver	Scaled parameters			Other
FSkin	Scaled parameters			Other
VFat	Scaled parameters			Other
VRich	Scaled parameters			Other
VLiver	Scaled parameters			Other
VSkin_e	Scaled parameters			Other
VSkin_u	Scaled parameters			Other
VSkin_sc_e	Scaled parameters			Other
VSkin_sc_u	Scaled parameters			Other
VBlood	Scaled parameters			Other
VPoor	Scaled parameters			Other
VArt	Scaled parameters			Other
VVen	Scaled parameters			Other
FSkin_e	Scaled parameters			Other
FSkin_u	Scaled parameters			Other
PCFat	Partition coefficient: fat over blood			PartitionCoefficient

Table	3.391	- continued	from	previous page
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ld	Description	Default	Unit	Туре
PCPoor	Partition coefficient: muscle over blood (poorly perfused tissue)			PartitionCoefficient
PCRich	Partition coefficient: viscera over blood (richly perfused tissue)			PartitionCoefficient
PCLiver	Partition coefficient: liver over blood			PartitionCoefficient
PCSkin	Partition coefficient: viable skin / blood			PartitionCoefficient
PCSkin_sc	Partition coefficient: viable skin / stratum corneum			PartitionCoefficient
ResampledPC- Fat	Resampled value PCFat			PartitionCoefficient

 Table 3.391 – continued from previous page

Model aliases: EuroMix_Generic_PBTK_model_V5, CosmosV4, CosmosV5.

EuroMix Generic PBTK model (v6)

Cosmos version 6 (received 3/27/2019)

Table 3.392: Exposure routes (forcings)

ld	Description	Unit
Oral	Oral exposure	mmoles
Dermal	Dermal exposure	mmoles
Inhalation	Inhalatory exposure	mmoles

ld	Description	ScalingFactor	Multiplication- Factor	Unit
CTotal	Total concentration			mM
CVen	Venous blood concentration	scVBlood	0.66667	mM
CArt	Arterial blood concentration	scVBlood	0.33333	mM
CFat	Fat (adipose) tissue concentration	scVFat		mM
CPoor	Poorly perfused tissue (muscle) concentration			mM
CRich	Richly perfused tissue (viscera) concentration	scVRich		mM
CLiver	Liver concentration	scVLiver		mM
CSkin_u	Viable unexposed skin concentration			mM
CSkin_e	Viable exposed skin concentration	BSA, Height_vs, fsA_exposed		mM
CSkin_sc_u	Skin unexposed stratum corneum concentration			mM
CSkin_sc_e	Skin exposed stratum corneum concentration	BSA, Height_vs, fsA_exposed		mM

Table 3.393: Model outputs

Table 3.394: Model parameters

ld	Description	Default	Unit	Туре
BM	Body mass		kg	Physiological
BSA	Body surface area (internally scaled by an allometric scaling factor s = 70/BM^0.3)		dm2	Physiological
scVFat	Fat as fraction of total body volume			Physiological
scVRich	Richly perfused tissues (viscera) as fraction of total body volume			Physiological
scVLiver	Liver as fraction of total body volume			Physiological
scVBlood	Blood as fraction of total body volume			Physiological
Height_sc	Skin thickness		decimeter	Physiological
Height_vs	Viable skin			Physiological
scFBlood	Total blood flow per unit mass		L/h/kg	Physiological
scFFat	Fat fraction of total blood flow going to compartments			Physiological

ld	Description	Default	Unit	Туре
scFPoor	Poorly perfused tissues (muscles) fraction of total blood flow going to compartments			Physiological
scFLiver	Liver fraction of total blood flow going to compartments			Physiological
scFSkin	Skin fraction of total blood flow going to compartments			Physiological
Falv	Alveolar ventilation rate		L/h	Physiological
mic	Microsomal proteins content		mg/gr liver	Physiological
PCAir	Partition coefficient: blood over air			PartitionCoefficient
log_PCFat	Scaled parameter, partition coefficient: fat over blood			PartitionCoefficient
log_aPoor	Scaled parameter, partition coefficient: muscle over blood (poorly perfused tissue)			PartitionCoefficient
log_aRich	Scaled parameter, partition coefficient: viscera over blood (richly perfused tissue)			PartitionCoefficient
log_aLiver	Scaled parameter, partition coefficient: liver over blood			PartitionCoefficient
log_aSkin	Scaled parameter, partition coefficient: viable skin over blood			PartitionCoefficient
log_aSkin_sc	Scaled parameter, partition coefficient: viable skin stratum corneum over blood			PartitionCoefficient
Kp_sc_vs	Diffusion rate from stratum corneum to viable skin		decimeter/h	Physicochemical
Ke	Renal excretion rate		L/h	Physicochemical
Michaelis	Flag for Michaelis-Menten vs linear metabolism (0 = linear)			Physicochemical
Vmax	Maximum rate of metabolism		mmoles/h/L liver	Physicochemical
Km	Michaelis-Menten constant for metabolism		mM	Physicochemical

Table 3.394 – continued from previous page

ld	Description	Default	Unit	Туре
CLH	Hepatic Physicochemical clearance			Physicochemical
fub	Unbound fraction in blood			Physicochemical
Frac	Fraction absorbed by the gut			Physicochemical
kGut	Oral 1st order absorption rate constant		1/h	Physicochemical
Cinh	Inhalation			Other
Tinh	Inhalation duration			Other
OralDose			mmol	Other
DermalDose			mmol	Other
fSA_exposed	Fraction of skin surface area actually exposed			Physicochemical
FBlood	Blood flow			Other
FFat	Scaled parameters, blood flow to the fat			Other
FPoor	Scaled parameters, blood flow to poorly perfused tissues			Other
FRich	Scaled parameters, blood flow to richly perfused tissues			Other
FLiver	Scaled parameters, blood flow to the liver			Other
FSkin	Scaled parameters, blood flow to the skin			Other
VFat	Scaled parameters			Other
VRich	Scaled parameters, richly perfused tissue volume			Other
VLiver	Scaled parameters, liver volume			Other
VSkin_e	Scaled parameters, exposed skin volume			Other
VSkin_u	Scaled parameters, unexposed skin volume			Other
VSkin_sc_e	Scaled parameters, stratum corneum exposed skin volume			Other
VSkin_sc_u	Scaled parameters, stratum corneum unexposed skin volume			Other
VBlood	Scaled parameters, blood volume			Other
VPoor	Scaled parameters, poorly perfused tissue volume			Other

Table	3.394 -	continued	from	previous	page

ld	Description	Default	Unit	Туре
VArt	Scaled parameters, arterial blood volume			Other
VVen	Scaled parameters, venous blood volume			Other
FSkin_e	Scaled parameters, blood flow to exposed skin			Other
FSkin_u	Scaled parameters, blood flow to unexposed skin			Other
PCFat	Partition coefficient: fat over blood			PartitionCoefficient
PCPoor	Partition coefficient: muscle over blood (poorly perfused tissue)			PartitionCoefficient
PCRich	Partition coefficient: viscera over blood (richly perfused tissue)			PartitionCoefficient
PCLiver	Partition coefficient: liver over blood			PartitionCoefficient
PCSkin	Partition coefficient: viable skin over blood			PartitionCoefficient
PCSkin_sc	Partition coefficient: viable skin / stratum corneum			PartitionCoefficient
ResampledPC- Fat	Resampled value PCFat			PartitionCoefficient

 Table 3.394 – continued from previous page

Model aliases: EuroMix_Generic_PBTK_model_V6, CosmosV6.

PBK model chlorpyrifos (v1)

PBK model chlorpyrifos (v1)

Table 3.395:	Exposure	routes	(forcings)
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ld	Description	Unit		
Oral	Oral exposure	umoles		
ld	Description	ScalingFactor	Multiplication- Factor	Unit
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O_CV	Venous blood	VVc		uM
O_CP	Plasma from whole blood	VVc	0.6	uM
O_CU	Uterus tissue	VUc		uM
O_ACL	Cleared renally		0.03	umoles
O_CS	Slowly perfused tissue			umoles
O_CR	Richly perfused tissue			umoles
O_CF	Fat	VFc		umoles
O_CL	Liver	VLc		umoles
O_CK	Kidney	VKc		umoles
O_CM	Muscle	VMc		umoles
O_CH	Heart	VHc		umoles
O_CLu	Lung	VLuc		umoles
O_CBrb	Brain blood	VBrc	0.05	umoles
O_CBrt	Brain tissue	VBrc	0.95	umoles
O_CBr	Brain total	VBrc		umoles
O_CA	Arterial blood	VAc		umoles

Table 3.396: Model outputs

Table 3.397: Model parameters

ld	Description	Default	Unit	Туре
VLc	Fraction liver tissue of BW	0.0257		Physiological
VFc	Fraction fat tissue of BW	0.2142		Physiological
VLuc	Fraction lung tissue of BW	0.0076		Physiological
VAc	Fraction arterial blood of BW (0.074*1/4)	0.0198		Physiological
VVc	Fraction venous blood of BW (0.074*3/4)	0.0593		Physiological
VKc	Fraction kidney tissue of BW	0.004		Physiological
VMc	Fraction muscle tissue of BW	0.04		Physiological
VUc	Fraction uterus tissue of BW	0.0018		Physiological
VBrc	Fraction brain tissue of BW	0.02		Physiological
VHc	Fraction heart tissue of BW	0.0047		Physiological
QLc	Fraction of blood flow to liver	0.227		Physiological
QFc	Fraction of blood flow to fat	0.052		Physiological
QKc	Fraction of blood flow to kidneys	0.175		Physiological

ld	Description	Default	Unit	Туре
QMc	Fraction of blood flow to muscle	0.12		Physiological
QUc	Fraction of blood flow to uterus	0.2		Physiological
QBrc	Fraction of blood flow to brain	0.114		Physiological
QHc	Fraction of blood flow to heart	0.04		Physiological
MWP	Molecular weight	350.59	g/mol	Physicochemical
MWM1	Molecular weight	334 52	g/mol	Physicochemical
MWM2	Molecular weight	198.43	g/mol	Physicochemical
LogPP	Octanol/water	4.784	g, mor	PartitionCoefficient
LogPM1	Octanol/water	3.894		PartitionCoefficient
LogPM2	Octanol/water partition coefficient	1.856		PartitionCoefficient
Fa	Fraction absorbed. Obtained from Nolan 1984	0.7		Physicochemical
KaS	Absorption constant stomach. Obtained from Timchalk et al. 2002 (stomach; /h)	0.00000733	/h	Physicochemical
Kal	Absorption constant intestine. Obtained from Timchalk et al. 2002 (intestine; /h)	1.00033	/h	Physicochemical
KsI	Absorption constant (transfer stomach to intestine). Obtained from Timchalk et al. 2002 (transfer stomach to intestine; /h)	0.967749	/h	Physicochemical
fuP	Unbound fraction in plasma. Obtained from SimCyp	0.021		Physicochemical
fuM1	Unbound fraction in plasma. Obtained from SimCyp	0.15		Physicochemical
fuM2	Unbound fraction in plasma. Obtained from SimCyp	0.082		Physicochemical
BPP	B/P ratio obtained from Hsu, 2013. If no data available, set to 1	1.3		Physicochemical
BPM1	B/P ratio obtained from Hsu, 2013. If no data available, set to 1	2.7		Physicochemical

Table 3.397 - continued from pre	evious page
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ld	Description	Default	Unit	Туре
BPM2	B/P ratio obtained from Hsu, 2013. If no data available, set to 1	1		Physicochemical
KurineP	Urinary excretion rate constant (/h)	0	/h	Physicochemical
KurineM1	Urinary excretion rate constant (/h)	0	/h	Physicochemical
KurineM2	Urinary excretion rate constant (/h)	0.026	/h	Physicochemical
CYPabundance- CYP1A2	CYP1A2 abundance (pmolCYP/mg protein) ;(calculated based on database in Simcyp; sum of EM, PM and UM)	52		Physicochemical
CYPabundance- CYP2B6	CYP2B6 abundance (pmolCYP/mg protein) ;(calculated based on database in Simcyp; sum of EM, PM and UM)	15.8		Physicochemical
CYPabundance- CYP2C19	CYP2C19 abundance (pmolCYP/mg protein) ;(calculated based on database in Simcyp; sum of EM, PM and UM)	5.4		Physicochemical
CYPabundance- CYP3A4	CYP3A4 abundance (pmolCYP/mg protein) ;(calculated based on database in Simcyp; sum of EM, PM and UM)	137		Physicochemical
ISEFCYP1A2	Scaling factor CYP1A2 ISEF (Vmax) (non compound-specific) (calculated based on probe incubation, lab specific)	0.072		Physicochemical
ISEFCYP2B6	Scaling factor CYP2B6 ISEF (Vmax) (non compound-specific) (calculated based on probe incubation, lab specific)	0.476		Physicochemical

Table	3.397	- continued	from	previous page	

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ld	Description	Default	Unit	Туре
ISEFCYP2C19	Scaling factor CYP2C19 ISEF (Vmax) (non compound-specific) (calculated based on probe incubation, lab specific)	0.209		Physicochemical
ISEFCYP3A4	Scaling factor CYP3A4 ISEF (Vmax) (non compound-specific) (calculated based on probe incubation, lab specific)	0.107		Physicochemical
MPL	Scaling factor of human liver microsome in mg to gram liver (mg microsomal protein /g liver) (Al-Malahmeh, A. J et al., 2017) (Barter et al., 2007)	32	mg/g	Physicochemical
VMaxCYP1- A2mP1	Vmax of CYP1A2 at supersome level (pmol/min/pmol CYP) (Experimentally determined using supersomes, Chen et al., 2022)	3.963		Physicochemical
VMaxCYP2- B6mP1	Vmax of CYP2B6 at supersome level (pmol/min/pmol CYP) (Experimentally determined using supersomes, Chen et al., 2022)	7.755		Physicochemical
VMaxCYP2- C19mP1	Vmax of CYP2C19 at supersome level (pmol/min/pmol CYP) (Experimentally determined using supersomes, Chen et al., 2022)	2.744		Physicochemical
VMaxCYP3- A4mP1	Vmax of CYP3A4 at supersome level (pmol/min/pmol CYP) (Experimentally determined using supersomes, Chen et al., 2022)	17.78		Physicochemical

Table 3.397 - continued from previous page

ld	Description	Default	Unit	Туре
KmCYP1A2P1	Affinity constant of CYP1A2 (umoles/L) (Experimentally determined using supersomes, Chen et al., 2022)	0.61	umoles/L	Physicochemical
KmCYP2B6P1	Affinity constant of CYP2B6 (umoles/L) (Experimentally determined using supersomes, Chen et al., 2022)	0.14	umoles/L	Physicochemical
KmCYP2C19- P1	Affinity constant of CYP2C19 (umoles/L) (Experimentally determined using supersomes, Chen et al., 2022)	1.89	umoles/L	Physicochemical
KmCYP3A4P1	Affinity constant of CYP3A4 (umoles/L) (Experimentally determined using supersomes, Chen et al., 2022)	29.77	umoles/L	Physicochemical
VMaxCYP1- A2mP2	Vmax of CYP1A2 at supersome level (pmol/min/pmol CYP) (Experimentally determined using supersomes, Chen et al., 2022)	2.957		Physicochemical
VMaxCYP2- B6mP2	Vmax of CYP2B6 at supersome level (pmol/min/pmol CYP) (Experimentally determined using supersomes, Chen et al., 2022)	5.492		Physicochemical
VMaxCYP2- C19mP2	Vmax of CYP2C19 at supersome level (pmol/min/pmol CYP) (Experimentally determined using supersomes, Chen et al., 2022)	17.51		Physicochemical

Table 3.397 - continued from previous page

ld	Description	Default	Unit	Туре
VMaxCYP3- A4mP2	Vmax of CYP3A4 at supersome level (pmol/min/pmol CYP) (Experimentally determined using supersomes, Chen et al., 2022)	23.86		Physicochemical
KmCYP1A2P2	Affinity constant of CYP1A2 (umoles/L) (Experimentally determined using supersomes, Chen et al., 2022)	1.25	umoles/L	Physicochemical
KmCYP2B6P2	Affinity constant of CYP2B6 (umoles/L) (Experimentally determined using supersomes, Chen et al., 2022)	1.28	umoles/L	Physicochemical
KmCYP2C19- P2	Affinity constant of CYP2C19 (umoles/L) (Experimentally determined using supersomes, Chen et al., 2022)	1.37	umoles/L	Physicochemical
KmCYP3A4P2	Affinity constant of CYP3A4 (umoles/L) (Experimentally determined using supersomes, Chen et al., 2022)	18.13	umoles/L	Physicochemical
VMax3c	Maximum velocity constant (nmol/min/ml plasma) (Experimentally determined using microsomes, Chen et al., 2022)	37.98	nmoles/min/ml	Physicochemical
Km3	Affinity constant (umoles/L) (Experimentally determined)	627.9		Physicochemical
VMax4c	Maximum velocity constant (nmol/min/ml plasma) (Experimentally determined using microsomes, Chen et al., 2022)	1844	nmoles/min/ml	Physicochemical

Table 3.397 - continued from previous page
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ld	Description	Default	Unit	Туре
Km4	Affinity constant (umoles/L) (Experimentally determined)	289.8	umoles/L	Physicochemical
BW		70	kg	Physiological

	Table	3.397	- continued	from	previous	page
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Model aliases: PBK_Chlorpyrifos_V1.

1 Note

Additional kinetic models can be implemented, please contact the MCRA administrator.

EuroMix generic PBK model

Reference: Tebby et al. (2020)

In MCRA updated versions (version 4b, 6) of the PBK model developed at INERIS in the framework of the COSMOS project is used. The model describes the distribution of chemicals in venous blood, arterial blood, adipose tissues, poorly perfused tissues (muscles), gut lumen, liver, richly perfused tissues (other viscera), and skin. Each of those is described as a compartment (homogeneous virtual volume) in which distribution is instantaneous and limited only by the incoming blood flow or rate of entry in the compartment. Exposure can occur through the dermal route, ingestion or inhalation. The absorbed molecules can be excreted to urine, exhaled through the lung, or metabolized in liver.

The EuroMix generic PBK model is coded as a set of ordinary differential equations. There is one such equation per time-dependent chemical quantity of the model (so-called state variables). There are 13 state variables in the model: the quantity of chemical in venous blood (Q_{ven}) , in arterial blood (Q_{art}) , in adipose tissues (Q_{fat}) , in poorly perfused tissues (Q_p) , in well perfused tissues (Q_r) , in liver (Q_{liv}) , in unexposed skin $(Q_{s,u})$, in exposed skin $(Q_{s,e})$, in the stratum corneum of unexposed skin $(Q_{sc,u})$, in exposed stratum corneum $(Q_{sc,e})$, in gut lumen (Q_{gut}) , the quantity excreted to urine (Q_{ex}) , and the quantity metabolized (Q_{met}) . The model can predict, as a function of time, for given oral, dermal and/or inhalation exposures, all the above quantities and the corresponding concentrations as a function of time. Concentrations are obtained by dividing quantities by compartment volumes Tebby et al. (2020).

In Figure 3.78 a time course of the internal substance amount (μg) for Clothianidin in the liver is shown. For 50 consecutive days a bolus per day is submitted. The red line shows the substance amount varying over time. The green line displays the average of the peaks representing acute exposure, the blue line displays the steady state representing chronic exposure, all after skipping a nonstationary period of 10 days (the vertical black line).

From the substance amount, a concentration is computed by dividing it by the total compartment weight (i.e., the mass/volume of the compartment/organ).

In Figure 3.79, for a large number of individuals the internal exposure (acute, green dots) in the liver is plotted versus the external exposure ($\mu g/kgbw$). The diagonal represents the 1:1 ratio of internal vs external exposure.

Bisphenol model

Reference: Karrer et al. 2019: Karrer et al. (2019)

'Structural analogs such as the bisphenols S, F, and AF (BPS, BPF, BPAF) are used to replace the endocrine disrupting chemical bisphenol A (BPA), but they exert estrogenic effects in the same order of magnitude. In order to investigate the consequences of BPA restrictions, we assessed the cumulative risk from BPA, BPS, BPF, and BPAF in Europe before and after the first BPA restrictions in 2011. We modelled external exposures from food, personal care products (PCPs), thermal paper, and dust, using the models MCRA and PACEM for food and PCPs, respectively. We calculated internal concentrations of unconjugated BPs with substance-specific PBK models and cumulated concentrations by taking into account relative estrogenic potencies. Average cumulative exposure to unconjugated BPs was 3.8 and 2.1 ng/kg bw/day before and after restrictions, respectively. The decline was mostly caused by the replacement of BPA with BPS in thermal paper. Therefore, the margins of exposure (MOEs) for estrogenic effects were mostly higher after the restrictions. However, in high uncertainty percentiles the MOEs were partly lower than



Figure 3.77: Schematic representation of the EuroMix Generic PBK model.



Model CosmosV6





РВРК

Figure 3.79: Internal versus external exposure for Clothianidin in the liver (EuroMix Generic PBK model version 6).

before (e.g. the MOEs for the uncertainty P97.5 of the variability P99 were 2.6 and 1.9 before and after restrictions, respectively), which shows the higher uncertainty around exposures for substitutes compared to BPA.'

Abstract: Linking probabilistic exposure and pharmacokinetic modelling to assess the cumulative risk from the bisphenols BPA, BPS, BPF, and BPAF for Europeans. Authors: Cecile Karrer, Waldo de Boer, Christiaan Delmaar, Yaping Cai, Amélie Crépet, Konrad Hungerbühler, Natalie von Goetz



Figure 3.80: Graphical abstract 'Linking probabilistic exposure and pharmacokinetic modelling to assess the cumulative risk from the bisphenols BPA, BPS, BPF, and BPAF for Europeans.'

3.7.4 PBK model definitions

In this module, PBK models with code representation in SBML format (Systems Biology Markup Language) are uploaded (with extension *.sbml*). Currently only Level 3, Version 2 is supported.

SBML has three main purposes:

- enable the use of multiple software tools without having to rewrite models to conform to every tool's idiosyncratic file format;
- enable models to be shared and published in a form that other researchers can use even when working with different software environments;
- ensure the survival of models beyond the lifetime of the software used to create them.

See also wiki, SBML portal.

Output of this module is used by: PBK model parametrisations

PBK model definitions from data

The *PbkModelDefinitions* table is an internal MCRA table which is automatically generated after uploading an sbml file (with extension *.sbml*).

PBK model definitions data formats

Data tables:

· PBK model definitions

PBK model definitions specify uploadable PBK models.

PBK model data format

The PBK model definition data format specifies uploadable PBK models.

Download empty dataset template: Zipped CSV Excel

PBK model definitions

PBK model definitions.

Table 3.398: Table definition for PBK model definitions.

Name	Туре	Description	Aliases	Required
Id	AlphaNumeric (50)	Unique identifier/code of the model.	id, idModel, idPbkModel- Definitions	Yes
Name	AlphaNumeric (100)	The name of the model.	Name, PbkName	No
Description FilePath	AlphaNumeric (200) FileReference (200)	Description of the model. Path to SBML xml.	Description FilePath	No Yes

Accepted table names: PbkModelDefinitions, PbkModelDefinition.

PBK model definitions as data

PBK model definitions implement an uploadable version of PBK (SBML) models.

- PBK model definitions data formats
- PBK model definitions from data

3.8 Risk modules

Exposures and *hazard characterisations* are compared in risk metrics. If both exposure and hazard characterisation are characterised by a single value, the risk metric (e.g. a traditional risk characterisation ratio (hazard/exposure), hazard quotient or (exposure/hazard)) can be calculated using module Single value risks. Module Risks allows for probabilistic risk calculations. In both cases a threshold can be specified to assist in interpretation. The threshold value should be chosen in relation to the assessment factors used in the hazard characterisation, e.g. a threshold MOE=100 (of HI=0.01) is often used if no assessment factors have been used, but a threshold 1 would be appropriate if assessment factors have already been used to address relevant uncertainties.

3.8.1 Environmental burden of disease

Environmental Burden of Disease (EBD) assessment.

This module has as primary entities: Populations Substances

Environmental burden of disease calculation

See also: Environmental burden of disease methodology, Plass et al. (2024).

In Figure 3.81, the attributable burden (in *DALY* s) with uncertainty bounds are displayed together with the cumulative percentage (right axis). On the x-axis, the exposure is displayed according to the specified bin boundaries *see settings*. Specify custom bins or *ERF* defined bins (i.c. 5, 10, 25, 50, 75, 90, 95%).



Figure 3.81: Attributable burden & cumulative percentage

In Figure 3.82, the attributable burden is standardized by the population size (*see primary entity populations data format*) and expressed per 100.000 individuals.

In Figure 3.83, the attributable burden is standardized by the population size in each bin and expressed per 100.000 individuals. The population size in a bin is determined by the *bin boundaries* as specified in the interface (custom or *ERF* defined).

Environmental burden of disease settings



Figure 3.82: Standardized attributable burden per 100.000 individuals & cumulative percentage



Figure 3.83: Standardized exposed attributable burden per 100.000 individuals & cumulative percentage

Selection settings

Table 3.399: Selection settings for module Environmental burden of dis-

ease.

Name	Туре	Description
Exposure type	ExposureType	The type of exposure considered in the assessment; acute (shor term) or chronic (long-term).

Calculation settings

Table 3.400: Calculation settings for module Environmental burden of disease.

Name	Туре	Description
Target level	TargetLevelType	Select whether the assessment is at the level of external dose, internal (target) concentration or internal (systemic) dose. For a aggregate assessment, combining dietary and nondietary exposures, the target level is always internal concentration or internal (systemic) dose. When only dietary exposures are available, the target level is optional and may be an external (administered) dose, an internal (tissue) concentration or an internal (absorbed) dose.
Exposure calculation method	ExposureCalculationMethod	Method for obtaining exposure estimates. These can be modell exposures (e.g., external (dietary) exposures or internal exposure estimates obtained by aggregating dietary and non-dietary exposures) or exposure estimates derived from human (bio)monitoring data.
Burden of disease indicators	BodIndicator	Burden of disease indicators.
Specify the Bod approach	BodApproach	Specify the approach used for estimating the attributable burde of disease.
Specify the exposure grouping method	ExposureGroupingMethod	Specify the method by which the exposures are grouped in bins
Specify the bin boundaries.	Numeric	Specify the boundaries of the bins by which the exposures are grouped.

Uncertainty settings

Table 3.401: Uncertainty settings for module Environmental burden of dis-

ease.	

Name	Туре	Description
Lower uncertainty limit (%)	Numeric	Percentage lower bound, e.g. 2.5%.
Upper uncertainty limit (%)	Numeric	Percentage upper bound, e.g. 97.5%.

Calculation of environmental burden of disease

EBDs are calculated from human monitoring data.

• Environmental burden of disease calculation

Inputs used: Dietary exposures Internal exposures HBM analysis Exposure response functions Burden of disease Settings used

• Calculation Settings

3.8.2 Risks

Risks (health impacts) are defined as a function of exposure and hazard characterisation at a chosen biological level (external or internal). Risk metrics are either based on the ratio hazard/exposure (e.g., MOE(T)) or exposure/hazard (e.g., HI, HQ, and RPI).

This module has as primary entities: Substances Effects Populations

Output of this module is used by: Single value risks

Risks calculation

A (cumulative) risk assessment aims to characterise the health impact due to exposure to one or multiple substances causing common adverse health effects. The health impact is characterised by a distribution of individual risks, expressed by a *risk metric* i.e., a risk characterisation ratio (hazard/exposure) (H/E); or a risk characterisation ratio (exposure/hazard) (E/H), comparing exposures and hazard characterisations at the chosen level (external or internal). Hazard characterisations are included as single values or in a probabilistic way.



Figure 3.84: Risk characterisation ratio (exposure/hazard) total distribution (triazoles).

The aim is to specify the probability that a random individual from a defined (sub)population will have an exposure high enough to cause a particular health effect of a predefined magnitude, the critical effect size. The exposure level that results in exactly that critical effect in a particular person is that person's individual critical hazard dose (*ICED*). Individuals in a population typically show variation, both in their individual exposure and in their hazard characterisation. Both the variation in exposure and the variation in hazard characterisation are quantified in the form of probability distributions. Assuming independence between both distributions, they are combined by Monte Carlo methods. The proportion of the H/E ratio distribution below the (safety/uncertainty) threshold (or the proportion of the E/H ratio distribution above e.g. 0.01 (in general 1) is the probability of critical effect (*POCE*), here 13%, in the particular (sub)population. Uncertainties involved in the overall risk assessment (i.e., both regarding exposure

and hazard characterisation) are quantified using Monte Carlo and bootstrap methods. This results in an uncertainty distribution for any statistic of interest.



Figure 3.85: Uncertainty of percentiles. The boxplots for uncertainty show the p25 and p75 as edges of the box, and p2.5 and p97.5 as edges of the whiskers. The reference (nominal) value is indicated with the dashed black line, the median with the solid black line within the box. Outliers are displayed as dots outside the wiskers. Risk is specified as the characterisation ratio (exposure/hazard) (triazoles).

In Figure 3.86, the risk characterisation ratios (exposure/hazard) for a number of substances are shown. As shown, the distinction between variability (grey bars, 90% probability) and uncertainty (whiskers) is retained. This is discussed in van der Voet and Slob (2007) and van der Voet et al. (2009).

In Figure 3.87, hazards versus exposures are plotted for the same substances.

Risk metric calculation type

Currently, two types of risk metric calculation types are available. Both for the risk characterisation ratio (hazard/exposure) (ratio H/E) and (exposure/hazard) (ratio E/H):

- exposures are cumulated over substances using RPFs and the risk distribution is estimated based on the hazard characterisation of the reference substance and the cumulative exposure. All RPFs should be supplied. See also *cumulative RPF weighted risk distribution*.
- risk is calculated per substance as a ratio of each hazard characterisation and the exposure. Then, the risk is estimated as the cumulated *sum of ratios* over all substances. All hazard characterisations should be supplied. In formula:
- risk is calculated per substance as a ratio of each hazard characterisation and the exposure. Then, the risk is estimated as the cumulated *sum of ratios* over all substances. All hazard characterisations should be supplied. In formula:



Risk ratio (exposure/hazard)

Figure 3.86: Risk characterisation ratio (exposure/hazard) plot for multiple substances (triazoles). The threshold (0.01) is indicated with the left vertical line, the vertical black line on the right indicates threshold value = 1.

For the risk characterisation ratio E/H:

$$E/H = HI = \sum_{s=1}^{S} HQ_s$$

where summation is over the number of substances per individual(day).

For the risk characterisation ratio H/E:

$$H/E = MOET = \frac{1}{\sum_{s=1}^{S} \frac{1}{H/E_s}}$$

where summation is over the number of substances per individual(day).

The blue line indicates the median value of the cumulative risks in the population. Note that the cumulative sum of the medians of the risk characterisation ratios of the contributing substances, here metals (red areas) does not necessarily add up to the cumulative risk (blue line). Simulations with multivariate lognormal distributions without correlation suggest that for moderate percentiles like the median the cumulative risk is higher than the risk calculated as sum of ratios.

Inverse distribution

Risk can be calculated as a distribution of either a risk characterisation ratio hazard/exposure (H/E) or exposure/hazard (E/H), if at least one of the inputs exposure and hazard characterisation is a distribution. The risk distribution is characterised by percentiles. To accommodate for matching results of H/E and E/H in the case of percentiles, there is an option to calculate percentiles via the complementary percentile of the *inverse distribution* in order to handle numerical differences when calculating percentiles for a left or right tail. The option is especially useful for small data sets where percentile calculation is asymmetric in both tails. When set, the percentile is calculated as the inverse of the complementary percentage of the inverse distribution. E.g., the p_1 of the H/E distribution



Figure 3.87: Hazard vs. exposure plot for multiple substances. 95% bivariate confidence areas for target hazard dose distribution and exposure distribution. Inner ellipses express variability, outer ellipses uncertainty.



Figure 3.88: Cumulative risk characterisation ratios exposure/hazard (median) in the population (metals).

is calculated as $1/(p_{99}$ of 1/H/E distribution); the p_{99} of the E/H distribution is calculated as $1/(p_1$ of 1/E/H distribution).

Risk by food

The option *calculate risks by modelled foods* is available when the target dose level is external. Dietary exposures preserving all the information of exposures of modelled foods are used to calculate risks statistics for modelled foods and to calculate the percentages at risk of modelled foods in the background and foreground based on the specified *threshold* in the safety plot.

For co-exposure of substances, see maximum cumulative ratio (MCR) and the exposure mixtures module.

Risks settings

Calculation settings

Name	Туре	Description
Selected tier	SettingsTemplateType	Specifies all module settings should be set according to a
		pre-defined tier or using custom settings.
Seed for pseudo-random	Numeric	A value of 0 will use a pseudo-random seed in each run, a value
Exposure type	FraguesTups	0 will provide the same results in a repeated run.
Exposure type	Ехрозите Гуре	term) or chronic (long-term).
Multiple substances analysis	Boolean	Specifies whether the assessment involves multiple substances.
Compute cumulative risks	Boolean	Specifies whether to compute the combined/cumulative risk over all substances.
Health effect type	HealthEffectType	Specifies whether the health effect is a risk (negative) or benefit (positive).
Risk characterisation ratio	RiskMetricType	Report risks in terms of the ratio exposure/hazard (e.g., HI, HQ RPI) or as hazard/exposure (e.g., MOE(T)).
Show equivalent animal dose output	Boolean	Specifies whether equivalent animal doses should be reported in the output.
Risk threshold	Numeric	Threshold for interpretation of the risk metric. For instance, for the Margin of Exposure (MOE) a threshold of 100 is commonl used and for the Hazard Index (HI) the threshold is usually 1.
Use inverse distribution to calculate percentile	Boolean	Calculate percentile via the complementary percentage of the inverse distribution (default: no). Description: E.g., P0.1 of MOE(T) distribution is calculated via P99.9 of 1/MOE(T) distribution. Note: This option is provided because percentile calculation in small data sets is asymmetric in both tails.
Exposure routes	ExposureRoutes	Exposure routes included in assessment.
Target level	TargetLevelType	Select whether the assessment is at the level of external dose, internal (target) concentration or internal (systemic) dose. For a aggregate assessment, combining dietary and nondietary exposures, the target level is always internal concentration or internal (systemic) dose. When only dietary exposures are available, the target level is optional and may be an external (administered) dose, an internal (tissue) concentration or an internal (absorbed) dose.
Calculate risks by modelled foods, substances or a combination of the two	Boolean	When the dose target level is external, dietary exposures are directly used as input to risks. Dietary exposures preserve the information of exposure by modelled foods, substances or the combination. Summarizing this information may time consumi
Exposure calculation method	<i>ExposureCalculationMethod</i>	Method for obtaining exposure estimates. These can be modell exposures (e.g., external (dietary) exposures or internal exposure estimates obtained by aggregating dietary and non-dietary exposures) or exposure estimates derived from human (bio)monitoring data.
Cumulation setting	<i>RiskMetricCalculationType</i>	Specify method for computing cumulative risks of multiple substances (e.g., via RPF weighted exposures or as a sum of ratios).
Perform MCR analysis	Boolean	Perform a Maximum Cumulative Ratio (MCR) analysis to determine co-exposure between substances.
Substance weighting in mixtures	ExposureApproachType	Risk based: exposures in equivalents of the reference substance standardised: standardised exposures per substance have varian 1; or unweighted exposures: RPFs are equal to 1.
Cutoff MCR	Numeric	For selection of individual(day) exposures with maximum cumulative ratio (MCR = total exposure/maximum) above the cutoff.
Cutoff percentage in population ranked on total	Numeric	For selection of individual(day) exposures above the cutoff percentage in the set of individual(day)s ranked on total exposu
616 Display ratio total exposure/	Numeric	Chapter 3. Modules For MCR plot: specify ratio total exposure/ maximum for
Show tail perceptiles (MCP	Numeric	mutvidual(day) exposures . Give specific percentiles of exposure distribution $(\%)$ e.g. 07.5
plot) for:	1 (unierie	90 (space separated)

Table 3.402: Calculation settings for module Risks.

Output settings

Name	Туре	Description
Left margin safety plot	Numeric	Left margin of the risk value in risk characterisation plots / saf charts.
Right margin safety plot	Numeric	Right margin of the risk value in risk characterisation plots / safety charts.
Number of plot labels	Numeric	Maximum number of labels to plot in hazard versus exposure p
Number of substances in hazard vs. exposure plot	Numeric	Maximum number of substances to plot in hazard vs exposure plot.
Inclusion percentage variability interval	Numeric	The central percentage of the variability distribution to include intervals for exposure, hazard and MOE (e.g. 90 means p5-p9).
Exclude privacy sensitive data from outputs	Boolean	Use this setting to not report the parts of the results (i.e., figure tables, or sections) that are marked as (potentially) privacy sensitive.
Include drill-down on 9 individuals around specified percentile	Boolean	Specifies whether drilldown on 9 individuals is to be included i the output.
Show percentiles for	Numeric	Give specific percentiles of exposure distribution (%), e.g. 50 95 97.5 99 (space separated).
Percentage for drilldown	Numeric	Gives detailed output for nine individuals near this percentile of the exposure distribution.
Percentage for upper tail	Numeric	Gives detailed output for this upper percentage of the exposure distribution.
Number of levels of covariable to predict exposure	Numeric	Specify the number of levels, e.g. 20. The range of the covaria is divided by the number of levels: range = (max - min)/levels. For these covariable levels exposures are predicted.
Predict exposure at extra covariable levels	Numeric	Specify specific prediction levels in addition to the automatical generated prediction levels (space separated).
Lower percentage for variability (%)	Numeric	The default value of 25% may be overruled.
Upper percentage for variability (%)	Numeric	The default value of 75% may be overruled.
Report consumptions and exposures per individual instead of per kg body weight	Boolean	Specifies whether body weights should be ignored and consumptions and exposures should be expressed per individua Otherwise, the consumptions and exposures are per kg body weight.

Table 3.403: Output settings for module Risks.

Uncertainty settings

Table 3.404:	Uncertainty	settings for	module Risks.
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Name	Туре	Description
Lower uncertainty limit (%)	Numeric	Percentage lower bound, e.g. 2.5%.
Upper uncertainty limit (%)	Numeric	Percentage upper bound, e.g. 97.5%.

Risks tiers

Overview

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
C pr cu m la ti ri	true	true	true	true	true	true	true	true	true	true
H ef fe ty	Risk	Risk	Risk	Risk	Risk	Risk	Risk	Risk	Risk	Risk
R cl ac te i- sa ti ra	Hazard- Exposur- eRatio	Hazard- Exposur- eRatio	Hazard- Exposur- eRatio	Hazard- Exposur- eRatio	Hazard- Exposur- eRatio	Hazard- Exposur- eRatio	Hazard- Exposur- eRatio	Hazard- Exposur- eRatio	Hazard- Exposur- eRatio	Haza Expo eRat
R th	100	100	100	100	100	100	100	100	100	100
U in va di tr bu ti- ti- ti- tc c c c c la po	true	true	true	true	true	true	true	true	true	true
E po su ca cu la tio m	Modelled- Concen- tration	Modelled- Concen- tration	Modelled- Concen- tration	Modelled- Concen- tration	Modelled- Concen- tration	Modelled- Concen- tration	Modelled- Concen- tration	Modelled- Concen- tration	Modelled- Concen- tration	Mod Con tratic

Table 3.405: Tier overview for module Risks.

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
Ir cl si po ce ag va al it in te	99.8	99.8	99.8	99.8	99.8	99.8	99.8	99.8	99.8	99.8
C su tid of th sa da co fr th sa sa sa p	false	false	false	false	false	true	true	false	false	false
U uu va al it	true		true		false	true		true		true
M th ad	false	false	false	false	false		false	false	false	false
C va at m el	false	false	false	false	false	false	false	false	false	false
It en at su ve	false	false	false	false	false	false	false	false	false	false

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 2022 Acu Tier
R p c sı ti aı ez p sı p in di vi u in st oi p k; b w	false	false	false	false	false	false	false	false	false	false
Ig no sa pl w	true	true	true	true				true	true	true
E cl in di vi al w le th N di	false	true	false	true				false	true	false
T ta di st cc cc tr ti di	false	false	false	false	false		false	false	false	false

Table 3.405 - continued from previous page

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
F tee sa p c c c c ir t t t c c c t t t t t t t t t t t e : :	false	false	false	false		false	false	false	false	false
U su st co vo si ru	true	true	true	true				true	true	true
S st co vo si m	UseMost- Toxic	UseMost- Toxic	DrawRan- dom	DrawRan- dom				UseMost- Toxic	UseMost- Toxic	Drav dom
R ta al al cca su st at te ad ti su st al lc cca ti i	true	true	true	true				true	true	true

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
A cu fc su st au th ti ir su st cu vu si	false	false	true	true				false	false	true
F d p c s t s t a l c c t t i r c c s t t t t t t c c s t s t s t t s t t s t t s t t s t t s t t s t t s t t s t t s t t s t t s t t s t t s t t s t t s s t s s t s t s t s t s t s t s s t s t s t s t s t s t s t s s t s t s t s s t s t s t s t s t s t s t s s t s	false	false	false	false				false	false	false
U e: tr o la ti	true	true	true	true				false	false	false
T o f c e t r o la ti	10	10	10	10						

Table 3.405 - continued from previous page

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
R st tr o la ti to e M	true	true	true	true						
R st tr tr o la ti tc an tf ri u	true	true	true	true						
In p w te co te ti	true	true	true	true				true	true	true
V te co tr ti v ()	0.1	0.1	0.05	0.05				0.1	0.1	0.05

Table 3.405 – continued from previous page							ge			
N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
R st te ti ti ti ti ti ti ti ti ti ti ti ti ti	true	true	true	true				true	true	true
R si w ta in p ta ti ti to a ti ti ti	false	false	false	false				false	false	false
R Si W te in p ta ti to a si Si	false	false	false	false				true	true	true
A p p c in fa	true	true	true	true	true	true	true	true	true	true

										9-
	N EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
1	U false di tr bi ti	false	false	false	false	false	false	false	false	false
	Ig false nd pp cd in fa fa fa tc tc tt t	false	false	false	false	true	true	false	false	false
	D Empirical fa co co tr tr ti	Empirical	Empirical	Empirical	Empirical	NonDe- tect- SpikeLog- Normal	NonDe- tect- SpikeLog- Normal	Empirical	Empirical	Emp
	Ir false cl M fa bi m	false	false	false	false	true	true	false	false	false
	R false st L ir p ta ta ta tt tt tt	false	false	false		false	false	false	false	false
	C Replace- sc ByLOR va rc pj m	Replace- ByLOR	Replace- ByLOR	Replace- ByLOR	Replace- ByZero	Replace- ByLOR	Replace- ByLOR	Replace- ByLOR	Replace- ByLOR	Repl ByL

Table 3.405 - continued from previous page

1	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
H tt ff (() xx I cc ff X I C ff X I I I ff fx () I I I I I I I I I	I 0.5	0.5	0.5	0.5		1	1	0.5	0.5	0.5
S F b	true	true	true	true	true	true	true	true	true	true
I F F r iii V U U f f f a a a a v V U U U U U I iii V V U U I F F F F F F F F F F F F F F F F F		true	true	true	false	true	true	true	true	true

Table 3.405 - continued from previous page

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
C rec la ir p va u w sa sa p p te	true	true	false	false	false	true	true	true	true	false
U or rec fr q ci fc ir fc ir p ta ti	true	true	true	true	false			true	true	true
P m ri u co ta	false	false	false	false	false	true	false	false	false	false
A pi o c r e pi t e pi t e pi c c a	false	false	true	true				false	false	true
T go le	External	External	External	External	External	External	External	External	External	Exte
U va al it m	BetaDis- tribution		BetaDis- tribution			BetaDis- tribution		BetaDis- tribution		Beta tribu

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N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
E ti n n tı	Realistic		Realistic			Realistic		Realistic		Real
U a it p ra e te	Variabili- tyFactor		Variabili- tyFactor			Variabili- tyFactor		Variabili- tyFactor		Vari tyFa
N ty		OIM		OIM	OIM		OIM		OIM	
N (1 b o d in su v		2		2					2	
S u fi q to 1			true	true						true
R st u p c a a t t t t t t t u s o t c a t t t t t u s o t t c a t t t t t t t t t t t t t t t t			true	true						true
M F to (1 x N						1	1			

	N EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
	C m la ti se ti									
) 1 1 1 1 1 1 1 1 1	C co tr ti li fi fi te ez									
t 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	ce fa tc Ir cl fc ca									
	ca nr it ca ca tr ti									
	Fi ca m it st st ci ci ci ci ci ci									

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N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2012 Op- timistic	EFSA 2012 Acute Pes- simistic	EFSA 2012 Chronic Pes- simistic	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFS 202 Acu Tier
A ju m fa tc fc tc fc tc fc tc fc tc fc tc tr ti										
U de te m is tie st co ve si fc fc cc co m it										
Ir cl pr cd di ti fo cd ti fo cd fo ir n r fo r fo r r e pr m r										

Retrospective dietary CRA (EC 2018) - Acute / Tier I

Table 3.406:	Tier	definition	for	Retrospective	dietary	CRA	(EC	2018)	-
Acute / Tier	I.								

Name	Setting	From input tier	In module
Compute cumulative risks	true		(
Health effect type	Risk		
Risk characterisation ratio	Hazard-		
	Exposur-		
	eRatio		
Risk threshold	100		
Use inverse distribution to calculate percentile	true		
Exposure calculation method	Modelled- Concen- tration		
Inclusion percentage variability interval	99.8		
Consumptions on the same day come from the same sample	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Dietary exposures
Use unit variability	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Dietary exposures
Model-then-add	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Dietary exposures
Covariate modelling	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Dietary exposures
Iterate survey	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Dietary exposures

Name	Setting	From input tier	In module
Report consumptions and exposures per individual instead of per kg body weight	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Dietary exposures
Ignore sampling weights	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Consump- tions
Exclude individuals with less than N days	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations

Table 3.406 - continued from previous page
Name	Setting	From input tier	ln module
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Account for substance authorisations in substance conversions	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Use extrapolation rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Threshold for extrapolation	10	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Restrict extrapolations to equal MRLs	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Restrict extrapolations to authorised uses	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations

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Name	Setting	From input tier	In module
Impute water concentrations	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Water concentration value (µg/kg)	0.1	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Restrict water imputation to approved substances	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Processing factors

Table 3.406 - continued from previous page

Name	Setting	From	ln modulo
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EC 2018) -	Processing factors
Default concentration model	Empirical	Acute / Tier I Retrospec- tive dietary CRA (EC 2018) - Acute / Tian I	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Sample based	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models

Name	Setting	From input tier	In module
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Correlate imputed values with sample potency	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Parametric uncertainty	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Apply occurrence pattern percentages	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Occur- rence patterns
Target level	External	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Hazard character- isations
Unit variability model	BetaDis- tribution	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Unit variability factors

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Name	Setting	From input tier	In module
Estimates nature	Realistic	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Unit variability factors
Unit variability parameter	Variabili- tyFactor	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Unit variability factors

Table 3.406 - continued from previous page

Retrospective dietary CRA (EC 2018) - Chronic / Tier I

Table 3.407: Tier definition for Retrospective dietary CRA (EC 2018) - Chronic / Tier I.

Name	Setting	From input tier	In module
Compute cumulative risks	true		
Health effect type	Risk		
Risk characterisation ratio	Hazard- Exposur- eRatio		
Risk threshold	100		
Use inverse distribution to calculate percentile	true		
Exposure calculation method	Modelled- Concen- tration		
Inclusion percentage variability interval	99.8		
Consumptions on the same day come from the same sample	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Dietary exposures
Model type	OIM	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Dietary exposures
		continues or	n next page

Name	Setting	From input tier	ln module
Model-then-add	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Dietary exposures
Covariate modelling	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Dietary exposures
Iterate survey	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Dietary exposures
Ignore sampling weights	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Consump- tions
Exclude individuals with less than N days	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Consump- tions
N (number of days in survey)	2	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Consump- tions

Table 3.407 - continued from previous page

Name	Setting	From input tier	ln module
Total diet study concentration data	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Account for substance authorisations in substance conversions	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations

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Name	Setting	From input tier	In module
Use extrapolation rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Threshold for extrapolation	10	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Restrict extrapolations to equal MRLs	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Restrict extrapolations to authorised uses	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Impute water concentrations	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Water concentration value (µg/kg)	0.1	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations

Table 3.407 - continued from previous page

Name	Setting	From input tier	In module
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Restrict water imputation to approved substances	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Processing factors
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Processing factors
Default concentration model	Empirical	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models

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Name	Setting	From input tier	In module
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Sample based	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Correlate imputed values with sample potency	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models

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Name	Setting	From input tier	In module
Parametric uncertainty	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Apply occurrence pattern percentages	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Occur- rence patterns
Target level	External	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Hazard character- isations

Table 3.407 – continued from previous page

Retrospective dietary CRA (EC 2018) - Acute / Tier II

Table 3.408: Tier definition for Retrospective dietary CRA (EC 2018) - Acute / Tier II.

Name	Setting	From input tier	ln module
Compute cumulative risks	true		
Health effect type	Risk		
Risk characterisation ratio	Hazard-		
	Exposur-		
D 111111	eRatio		
Risk threshold	100		
Use inverse distribution to calculate percentile	true		
Exposure calculation method	Modelled-		
	Concen- tration		
Inclusion percentage variability interval	99.8		
Consumptions on the same day come	false	Retrospec-	Dietary
from the same sample		tive	exposures
		dietary	
		CRA (EC	
		2018) -	
		Acute /	
		Tier II	
		continues of	n next page

Name	Setting	From input tier	ln module
Use unit variability	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Dietary exposures
Model-then-add	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Dietary exposures
Covariate modelling	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Dietary exposures
Iterate survey	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Dietary exposures
Ignore sampling weights	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Consump- tions
Exclude individuals with less than N days	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Consump- tions

Table 3.408 - continued from previous page

Name	Setting	From input tier	ln module
Total diet study concentration data	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Use extrapolation rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Threshold for extrapolation	10	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Restrict extrapolations to equal MRLs	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Restrict extrapolations to authorised uses	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Impute water concentrations	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations

Table 3.408 - continued from previous page

Name	Setting	From input tier	ln module
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Restrict water imputation to approved substances	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Processing factors
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Processing factors
Default concentration model	Empirical	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models

Table	3.408 -	continued	from	previous	page
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Name	Setting	From input tier	In module
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Sample based	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Correlate imputed values with sample potency	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models

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Name	Setting	From	In
		input tier	module
Parametric uncertainty	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Apply occurrence pattern percentages	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Occur- rence patterns
Target level	External	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Hazard character- isations
Unit variability model	BetaDis- tribution	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Unit variability factors
Estimates nature	Realistic	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Unit variability factors

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Name	Setting	From input tier	ln module
Unit variability parameter	Variabili- tyFactor	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Unit variability factors

Table 3.408 - continued from previous page

Retrospective dietary CRA (EC 2018) - Chronic / Tier II

Table 3.409: Tier definition for Retrospective dietary CRA (EC 2018) - Chronic / Tier II.

Name	Setting	From input tier	ln module
Compute cumulative risks	true		1
Health effect type	Risk		
Risk characterisation ratio	Hazard- Exposur- eRatio		
Risk threshold	100		
Use inverse distribution to calculate percentile	true		
Exposure calculation method	Modelled- Concen- tration		
Inclusion percentage variability interval	99.8		
Consumptions on the same day come from the same sample	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Dietary exposures
Model type	OIM	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Dietary exposures
Model-then-add	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Dietary exposures

Name	Setting	From input tier	ln module
Covariate modelling	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Dietary exposures
Iterate survey	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Dietary exposures
Ignore sampling weights	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Consump- tions
Exclude individuals with less than N days	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Consump- tions
N (number of days in survey)	2	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Food con- versions

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Name	Setting	From input tier	In module
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Use extrapolation rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations

Table	3.409 -	continued	from	previous	page

Name	Setting	From input tier	In module
Threshold for extrapolation	10	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Restrict extrapolations to equal MRLs	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Restrict extrapolations to authorised uses	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Impute water concentrations	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Restrict water imputation to approved substances	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Processing factors
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Processing factors
Default concentration model	Empirical	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models

Table 3.409 - continued from previous page

Name	Setting	From input tier	ln module
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Sample based	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Correlate imputed values with sample potency	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Parametric uncertainty	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models

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Name	Setting	From input tier	In module
Apply occurrence pattern percentages	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Occur- rence patterns
Target level	External	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Hazard character- isations

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Retrospective dietary CRA (EFSA 2012) - Optimistic

Use the optimistic model settings according to the EFSA Guidance 2012. Concentration values are sampled using a sample-based empirical distribution. Available processing factors are applied. No unit variability model should be applied.

NameSettingFrom input tierIn moduleCompute cumulative riskstruemoduleHealth effect typeRiskinRisk characterisation ratioHazard- Exposur- eRatioinRisk threshold100inUse inverse distribution to calculate percentiletrueinExposure calculation methodModelled- Concen- trationinInclusion percentage variability interval99.8inConsumptions on the same day come from the same samplefalseRetrospec- tive dietary CRA (EFSA 2012) - OptimisticDietary exposuresUse unit variabilityfalseRetrospec- tive dietary CRA (EFSA 2012) - OptimisticDietary exposuresModel typeOIMRetrospec- tive dietary CRA (EFSA 2012) - OptimisticDietary exposuresModel-then-addfalseRetrospec- tive dietary CRA (EFSA 2012) - OptimisticDietary exposuresModel-then-addfalseRetrospec- tive dietary CRA (EFSA 2012) - OptimisticDietary exposuresModel-then-addfalseRetrospec- tive dietary CRA (EFSA 2012) - OptimisticDietary exposuresCovariate modellingfalseRetrospec- tive dietary (CRA (EFSADietary exposuresCovariate modellingfalseRetrospec- tive dietary (CRA (EFSADietary exposures
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Table 3.410: Tier definition for Retrospective dietary CRA (EFSA 2012) - Optimistic.

657

Retrospective dietary CRA (EFSA 2012) - Acute / Pessimistic

Acute probabilistic exposure assessment using the pessimistic model settings according to the EFSA Guidance 2012. Only processing factors > 1 are applied. For unit variability, the Beta distribution is applied.

Name	Setting	From input tier	In module
Compute cumulative risks Health effect type	true Risk		
Risk characterisation ratio	Hazard- Exposur- eRatio		
Risk threshold Use inverse distribution to calculate	100 true		
Exposure calculation method	Modelled- Concen- tration		
Inclusion percentage variability interval	99.8		
Consumptions on the same day come from the same sample	true	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Dietary exposures
Use unit variability	true	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Dietary exposures
Covariate modelling	false	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Dietary exposures
Iterate survey	false	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Dietary exposures

Table 3.411: Tier definition for Retrospective dietary CRA (EFSA 2012) - Acute / Pessimistic.

Name	Setting	From input tier	In module
Report consumptions and exposures per individual instead of per kg body weight	false	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Dietary exposures
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Processing factors
Ignore processing factors less than 1	true	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Processing factors
Default concentration model	NonDe- tect- SpikeLog- Normal	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Concen- tration models

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Name	Setting	From input tier	ln module
Include MRL fallback model	true	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Concen- tration models
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	1	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Concen- tration models
MRL Factor (f x MRL)	1	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Concen- tration models
Sample based	true	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Concen- tration models

Table 3.411 - continued from previous page

Name	Setting	From input tier	In module
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Concen- tration models
Correlate imputed values with sample potency	true	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Concen- tration models
Parametric uncertainty	true	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Concen- tration models
Target level	External	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Hazard character- isations
Unit variability model	BetaDis- tribution	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Unit variability factors
Estimates nature	Realistic	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Unit variability factors

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Name	Setting	From input tier	ln module
Unit variability parameter	Variabili- tyFactor	Retrospec- tive dietary CRA (EFSA 2012) - Acute / Pessimistic	Unit variability factors

Table 3.411 - continued from previous page

Retrospective dietary CRA (EFSA 2012) - Chronic / Pessimistic

Chronic probabilistic exposure assessment using the pessimistic model settings according to the EFSA Guidance 2012. Only processing factors > 1 are applied.

663

Name	Setting	From input tier	In module
Compute cumulative risks Health effect type Risk characterisation ratio	true Risk Hazard-		
	Exposur- eRatio		
Risk threshold Use inverse distribution to calc percentile	culate 100		
Exposure calculation method	Modelled- Concen- tration		
Inclusion percentage variability interval	y 99.8		
Consumptions on the same day from the same sample	y come true	Retrospec- tive dietary CRA (EFSA 2012) - Chronic / Pessimistic	Dietary exposures
Model type	OIM	Retrospec- tive dietary CRA (EFSA 2012) - Chronic / Pessimistic	Dietary exposures
Model-then-add	false	Retrospec- tive dietary CRA (EFSA 2012) - Chronic / Pessimistic	Dietary exposures
Covariate modelling	false	Retrospec- tive dietary CRA (EFSA 2012) - Chronic / Pessimistic	Dietary exposures
Iterate survey	false	Retrospec- tive dietary CRA (EFSA 2012) - Chronic / Pessimistic	Dietary exposures
Report consumptions and expo Bisk modulity	g body	Retrospec- tive	Dietary exposures
weight		dietary CRA (EFSA 2012)	

Table 3.412: Tier definition for Retrospective dietary CRA (EFSA 2012) - Chronic / Pessimistic.

Retrospective dietary CRA (EFSA 2022) - Acute / Tier I

Name	Setting	From input tier	In module
Compute cumulative risks Health effect type Risk characterisation ratio	true Risk Hazard- Exposur- eRatio		
Risk threshold Use inverse distribution to calculate percentile	100 true		
Exposure calculation method	Modelled- Concen- tration		
Inclusion percentage variability interval	99.8		
Consumptions on the same day come from the same sample	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Dietary exposures
Use unit variability	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Dietary exposures
Model-then-add	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Dietary exposures
Covariate modelling	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Dietary exposures

Table 3.413: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier I.

Name	Setting	From input tier	ln module
Iterate survey	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Dietary exposures
Ignore sampling weights	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Consump- tions
Exclude individuals with less than N days	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations

Table 3.413 – continued from previous page
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Name	Setting	From input tier	ln module
Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Account for substance authorisations in substance conversions	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Use extrapolation rules	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations

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Name	Setting	From input tier	ln module
Impute water concentrations	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Water concentration value (µg/kg)	0.1	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Restrict water imputation to approved substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Processing factors

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Name	Setting	From input tier	In module
Use distribution	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Processing factors
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Processing factors
Default concentration model	Empirical	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models

Table 3.413 - continued from previous page
Name	Setting	From input tier	ln module
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Sample based	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Correlate imputed values with sample potency	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Parametric uncertainty	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models

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Name	Setting	From input tier	In module
Apply occurrence pattern percentages	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Occur- rence patterns
Target level	External	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Hazard character- isations
Unit variability model	BetaDis- tribution	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Unit variability factors
Estimates nature	Realistic	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Unit variability factors
Unit variability parameter	Variabili- tyFactor	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Unit variability factors

Table 3.413 - continued from previous page

Retrospective dietary CRA (EFSA 2022) - Chronic / Tier I

Table 3.414: Tier definition for Retrospective dietary CRA (EFSA 2022) - Chronic / Tier I.

Name	Setting	From input tier	In module
Compute cumulative risks Health effect type	true Risk		

Name	Setting	From input tier	In module
Risk characterisation ratio	Hazard- Exposur- eRatio		
Risk threshold	100		
Use inverse distribution to calculate percentile	true		
Exposure calculation method	Modelled- Concen- tration		
Inclusion percentage variability interval	99.8		
Consumptions on the same day come from the same sample	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	<i>Dietary</i> <i>exposures</i>
Model type	OIM	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Dietary exposures
Model-then-add	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Dietary exposures
Covariate modelling	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Dietary exposures
Iterate survey	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	<i>Dietary</i> <i>exposures</i>

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Name	Setting	From input tier	In module
Report consumptions and exposures per individual instead of per kg body weight	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Dietary exposures
Ignore sampling weights	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Consump- tions
Exclude individuals with less than N days	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Consump- tions
N (number of days in survey)	2	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations

Table 3.414 - continued from previous page

Name	Setting	From input tier	ln module
Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Account for substance authorisations in substance conversions	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Use extrapolation rules	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations

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Name	Setting	From input tier	ln module
Impute water concentrations	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Water concentration value (µg/kg)	0.1	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Restrict water imputation to approved substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Processing factors

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Name	Setting	From input tier	ln module
Use distribution	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Processing factors
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Processing factors
Default concentration model	Empirical	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models

Name	Setting	From input tier	In module
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Sample based	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Correlate imputed values with sample potency	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Parametric uncertainty	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models

Table 3.414 - continued from previous page

Name	Setting	From input tier	ln module
Apply occurrence pattern percentages	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Occur- rence patterns
Target level	External	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Hazard character- isations

Table 3.414 – continued from previous page

Retrospective dietary CRA (EFSA 2022) - Acute / Tier II

Table 3.415:	Tier	definition	for	Retrospective	dietary	CRA	(EFSA	2022)
- Acute / Tier	II.							

Name	Setting	From input tier	In module
Compute cumulative risks	true		
Health effect type	Risk		
Risk characterisation ratio	Hazard-		
	Exposur-		
	eRatio		
Risk threshold	100		
Use inverse distribution to calculate percentile	true		
Exposure calculation method	Modelled-		
	Concen-		
	tration		
Inclusion percentage variability interval	99.8		
Consumptions on the same day come	false	Retrospec-	Dietary
from the same sample		tive	exposures
		dietary	
		CRA	
		(EFSA	
		2022) -	
		Acute /	
		Tier II	

Name	Setting	From input tier	In module
Use unit variability	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Dietary exposures
Model-then-add	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Dietary exposures
Covariate modelling	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Dietary exposures
Iterate survey	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Dietary exposures
Ignore sampling weights	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Consump- tions

Table 3.415 - continued from previous page

Name	Setting	From input tier	ln module
Exclude individuals with less than N days	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations

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Name	Setting	From input tier	ln module
Account for substance authorisations in substance conversions	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Use extrapolation rules	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Impute water concentrations	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations

Table 3.415 - continued from previous page

Name	Setting	From input tier	ln module
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Restrict water imputation to approved substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Processing factors
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Processing factors
Default concentration model	Empirical	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models

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Name	Setting	From input tier	In module
Include MRL fallback model	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Sample based	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models

Table 3.415 - continued from previous page

Name	Setting	From input tier	ln module
Correlate imputed values with sample potency	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Parametric uncertainty	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Apply occurrence pattern percentages	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Occur- rence patterns

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Name	Setting	From input tier	In module
Target level	External	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Hazard character- isations
Unit variability model	BetaDis- tribution	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Unit variability factors
Estimates nature	Realistic	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Unit variability factors
Unit variability parameter	Variabili- tyFactor	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Unit variability factors

Table 3.415 - continued from previous page

Retrospective dietary CRA (EFSA 2022) - Chronic / Tier II

Table 3.416: Tier definition for Retrospective dietary CRA (EFSA 2	2022)
- Chronic / Tier II.	

Name	Setting	From input tier	In module
Compute cumulative risks	true		
Health effect type	Risk		
Risk characterisation ratio	Hazard-		
	Exposur-		
	eRatio		
Risk threshold	100		
Use inverse distribution to calculate	true		
percentile			
Exposure calculation method	Modelled-		
	Concen-		
	tration		
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Name	Setting	From input tier	ln module
Inclusion percentage variability interval	99.8		
Consumptions on the same day come from the same sample	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Dietary exposures
Model type	OIM	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Dietary exposures
Model-then-add	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Dietary exposures
Covariate modelling	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Dietary exposures
Iterate survey	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Dietary exposures

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Name	Setting	From input tier	ln module
Ignore sampling weights	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Consump- tions
Exclude individuals with less than N days	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Consump- tions
N (number of days in survey)	2	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations

Table 3.416 - continued from previous page

Name	Setting	From input tier	ln module
Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Use extrapolation rules	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Impute water concentrations	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations

Table 3.416 – continued from previous page

Name	Setting	From input tier	In module
Water concentration value (µg/kg)	0.05	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Restrict water imputation to approved substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Processing factors

Table 3.416 - continued from previous page

Name	Setting	From input tier	ln module
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Processing factors
Default concentration model	Empirical	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models

Table 3.416 – continued from previous page

Name	Setting	From input tier	ln module
Sample based	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Correlate imputed values with sample potency	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Parametric uncertainty	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Apply occurrence pattern percentages	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Occur- rence patterns

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Name	Setting	From input tier	ln module
Scale up use frequency to 100%	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Occur- rence patterns
Target level	External	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Hazard character- isations

Table 3.416 - continued from previous page

Prospective dietary CRA (EFSA 2023) - Acute / Tier II

Table 3.417: Tier definition for Prospective dietary CRA (EFSA 2023) - Acute / Tier II.

Name	Setting	From input tier	In module
Compute cumulative risks	true		
Health effect type	Risk		
Risk characterisation ratio	Hazard-		
	Exposur-		
	eRatio		
Risk threshold	100		
Use inverse distribution to calculate percentile	true		
Exposure calculation method	Modelled-		
	Concen-		
	tration		
Cumulation setting	RPFWeighte		
Inclusion percentage variability	99.8		
interval			

Name	Setting	From input tier	ln module
Consumptions on the same day come from the same sample	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Dietary exposures
Use unit variability	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Dietary exposures
Model-then-add	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Dietary exposures
Covariate modelling	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Dietary exposures
Iterate survey	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Dietary exposures

Table 3.417 - continued from previous page

Name	Setting	From input tier	ln module
Ignore sampling weights	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Consump- tions
Exclude individuals with less than N days	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Consump- tions
Total diet study concentration data	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Food con- versions
Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Use extrapolation rules	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Impute water concentrations	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations

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Name	Setting	From input tier	ln module
Water concentration value (µg/kg)	0.05	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Restrict water imputation to approved substances	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Include focal commodity concentrations	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Focal commodity substance occurrence percentage	20	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Adjustment factor for the focal food/substance concentration	1	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Use deterministic substance conversions for focal commodity	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Apply processing factors	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Processing factors
Use distribution	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Processing factors
Ignore processing factors less than 1	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Processing factors
Default concentration model	Empirical	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models

Table 3.417 - continued from previous page

Name	Setting	From input tier	ln module
Include MRL fallback model	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Sample based	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models

Table 3.417 – continued from previous pace
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Name	Setting	From input tier	ln module
Correlate imputed values with sample potency	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Parametric uncertainty	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Apply occurrence pattern percentages	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Occur- rence patterns

Table 3.417 - continued from previous page

Name	Setting	From input tier	ln module
Target level	External	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Hazard character- isations
Unit variability model	BetaDis- tribution	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Unit variability factors
Estimates nature	Realistic	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Unit variability factors
Unit variability parameter	Variabili- tyFactor	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Unit variability factors

Table 3.417 - continued from	previous page
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Prospective dietary CRA (EFSA 2023) - Chronic / Tier II

Table 3.418: Tier definition for Prospective dietary CRA (EFSA 2023) - Chronic / Tier II.

Name	Setting	From input tier	In module
Compute cumulative risks	true		
Health effect type	Risk		
Risk characterisation ratio	Hazard-		
	Exposur-		
	eRatio		
Risk threshold	100		
Use inverse distribution to calculate percentile	true		
Exposure calculation method	Modelled-		
	Concen-		
	tration		
Cumulation setting	RPFWeighte		
		a susting of a second	

Name	Setting	From input tier	ln module
Inclusion percentage variability interval	99.8		
Consumptions on the same day come from the same sample	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Dietary exposures
Model type	OIM	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	<i>Dietary</i> <i>exposures</i>
Model-then-add	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Dietary exposures
Covariate modelling	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Dietary exposures
Iterate survey	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Dietary exposures

Table 3.418 - continued from previous page

Name	Setting	From input tier	ln module
Ignore sampling weights	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Consump- tions
Exclude individuals with less than N days	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Consump- tions
N (number of days in survey)	2	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Consump- tions
Total diet study concentration data	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Food con- versions
Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Use extrapolation rules	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Impute water concentrations	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Restrict water imputation to approved substances	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Include focal commodity concentrations	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Focal commodity substance occurrence percentage	20	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Adjustment factor for the focal food/substance concentration	1	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Use deterministic substance conversions for focal commodity	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Apply processing factors	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Processing factors
Use distribution	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Processing factors
Ignore processing factors less than 1	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Processing factors

Table 3.418 - continued from previous page
Name	Setting	From input tier	ln module
Default concentration model	Empirical	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Include MRL fallback model	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Sample based	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models

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Name	Setting	From input tier	In module
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Correlate imputed values with sample potency	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Parametric uncertainty	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Apply occurrence pattern percentages	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Occur- rence patterns

Table 3.418 - continued from previous page

Name	Setting	From input tier	ln module
Restrict use percentage up-scaling to authorised uses	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Occur- rence patterns
Target level	External	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Hazard character- isations

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Prospective dietary CRA (EFSA 2024) - Acute / Tier II

Table 3.419: Tier definition for Prospective dietary CRA (EFSA 2024) - Acute / Tier II.

Name	Setting	From input tier	In module
Compute cumulative risks	true		
Health effect type	Risk		
Risk characterisation ratio	Hazard- Exposur- eRatio		
Risk threshold	100		
Use inverse distribution to calculate percentile	true		
Exposure calculation method	Modelled- Concen- tration		
Cumulation setting	RPFWeighte		
Inclusion percentage variability interval	99.8		
Consumptions on the same day come from the same sample	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Dietary exposures

Name	Setting	From input tier	ln module
Use unit variability	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Dietary exposures
Model-then-add	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Dietary exposures
Covariate modelling	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Dietary exposures
Iterate survey	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Dietary exposures
Ignore sampling weights	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Consump- tions

Table 3.419 - continued from previous page

Name	Setting	From input tier	ln module
Exclude individuals with less than N days	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Consump- tions
Total diet study concentration data	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Food con- versions
Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Use extrapolation rules	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Impute water concentrations	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations

Table 3.419 - continued from previous page

Name	Setting	From input tier	ln module
Restrict water imputation to the five most toxic substances	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Restrict water imputation to approved substances	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Include focal commodity concentrations	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Focal commodity substance occurrence percentage	20	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Adjustment factor for the focal food/substance concentration	1	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Use deterministic substance conversions for focal commodity	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Include processed derivatives of the focal food in measurement replacement	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Apply processing factors	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Processing factors
Use distribution	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Processing factors
Ignore processing factors less than 1	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Processing factors
Default concentration model	Empirical	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models

Table 3.419 - continued from previous page

Name	Setting	From input tier	ln module
Include MRL fallback model	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Sample based	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models

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Name	Setting	From input tier	In module
Correlate imputed values with sample potency	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Parametric uncertainty	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Apply occurrence pattern percentages	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Occur- rence patterns

Table 3.419 - continued from previous page

Name	Setting	From input tier	ln module
Target level	External	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Hazard character- isations
Unit variability model	BetaDis- tribution	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Unit variability factors
Estimates nature	Realistic	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Unit variability factors
Unit variability parameter	Variabili- tyFactor	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Unit variability factors

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Prospective dietary CRA (EFSA 2024) - Chronic / Tier II

Table 3.420: Tier definition for Prospective dietary CRA (EFSA 2024) - Chronic / Tier II.

Name	Setting	From input tier	In module
Compute cumulative risks	true		
Health effect type	Risk		
Risk characterisation ratio	Hazard-		
	Exposur-		
	eRatio		
Risk threshold	100		
Use inverse distribution to calculate percentile	true		
Exposure calculation method	Modelled-		
	Concen-		
	tration		
Cumulation setting	RPFWeighte		
		a a salina sa a sa	

Name	Setting	From input tier	In module
Inclusion percentage variability interval	99.8		
Consumptions on the same day come from the same sample	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Dietary exposures
Model type	OIM	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Dietary exposures
Model-then-add	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Dietary exposures
Covariate modelling	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Dietary exposures
Iterate survey	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Dietary exposures

Table 3.420 - continued from previous page

Name	Setting	From input tier	In module
Ignore sampling weights	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Consump- tions
Exclude individuals with less than N days	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Consump- tions
N (number of days in survey)	2	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Consump- tions
Total diet study concentration data	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Food con- versions
Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Use extrapolation rules	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations

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Name	Setting	From input tier	ln module
Impute water concentrations	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Restrict water imputation to approved substances	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Include focal commodity concentrations	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations

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Name	Setting	From input tier	ln module
Focal commodity substance occurrence percentage	20	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Adjustment factor for the focal food/substance concentration	1	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Use deterministic substance conversions for focal commodity	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Include processed derivatives of the focal food in measurement replacement	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Apply processing factors	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Processing factors
Use distribution	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Processing factors

Table 3.420 - continued from previous page

Name	Setting	From input tier	ln module
Ignore processing factors less than 1	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Processing factors
Default concentration model	Empirical	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Include MRL fallback model	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models

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Name	Setting	From input tier	ln module
Sample based	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Correlate imputed values with sample potency	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Parametric uncertainty	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Apply occurrence pattern percentages	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Occur- rence patterns

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Name	Setting	From input tier	ln module
Scale up use frequency to 100%	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Occur- rence patterns
Target level	External	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Hazard character- isations

Table 3.420 – continued from previous page

Calculation of risks

Risk (health impact) is quantified as exposure relative to hazard characterisation. Exposures or hazards can be single values or distributions, the risk metric is a distribution if at least one of the inputs is a distribution (if both are single values, see the module single value risks). Risk metrics are valid for a specific biological level (external or internal at a specific biological matrix).

• Risks calculation

Inputs used: Dietary exposures Internal exposures Hazard characterisations HBM analysis Relative potency factors

Settings used

• Calculation Settings

Risks are expressed as distribution of risk characterisation ratios (hazard/exposure) or)exposure/hazard). The distribution is summarised by percentiles, and by the probability to exceed the specified threshold value (e.g. 1 or 100). The hazard vs. exposure plot compares the exposures and the hazard characterisation for individuals or individualdays in a population. Exposures, hazard characterisations and risks can be acute or chronic. The default unit for exposures and hazard characterisations is $\mu g/kgBW/day$, but this can be changed by choosing non-default units for consumptions, concentrations and/or body weight.

By using probabilistic tiers for both exposure and hazard characterisation, the calculated risk distribution is equal to the Integrated hazard/exposure (IMOE) distribution, as described for the Integrated Probabilistic Risk Assessment (IPRA) approach in van der Voet and Slob (2007) and van der Voet et al. (2009).

3.8.3 Single value risks

Single value risks are risk estimates obtained from combining single value exposures with single value hazard characterisations or as a percentile from a risk distribution.

This module has as primary entities: Substances Effects Populations

Single value risks calculation

Single value risks can be calculated in two ways.

- 1. From single value risks: single value exposures are combined with (single value) hazard characterisations.
- 2. As percentile from risks distribution: a percentile can be selected from a risks distribution.

See below for a more detailed explanation.

Combining single value exposures and hazard characterisations

Single value risks are computed by combining *single value exposures* by route/source and substance with (single value) *hazard characterisations* by substance. They are computed as risk characterisation ratios hazard/exposure, hazard quotient or risk characterisation ratios exposure/hazard, or as a percentage of the reference dose (100 * exposure/hazard characterisation).

Single value risks from individual risks

In this option, a percentage point can be specified for the chosen risk metric, e.g. the risk characterisation ratio hazard/exposure (H/E) or exposure/hazard (E/H). The corresponding percentile is calculated from the distribution of individual *risks*. The default percentiles are a risk characterisation ratio hazard/exposure at 0.1% or a ratio exposure/hazard at 99.9%. Specify whether the risk metric is calculated using the inverse distribution or not. This option is provided because percentile calculation in small data sets is asymmetric in both tails. When this option is set, the percentile is calculated as the inverse of the complementary percentage of the inverse distribution. E.g., the $p_{0.1}$ of the H/E distribution is calculated as $1/(p_{99.9}$ of 1/H/E distribution); the $p_{99.9}$ of the E/H distribution is calculated as $1/(p_{0.1}$ of 1/E/H distribution).

Adjustment factors and uncertainty specification

Many sources of uncertainty that may affect input data, model assumptions and assessment methodology do not enter the assessment. In EFSA (2020a) and EFSA (2020b), thirty-four sources of uncertainty were identified and the impact of each source on the ratio H/E was quantified. Some uncertainties tend to overestimate the ratio H/E, others tend to underestimate it. Following the guidance of the EFSA Scientific Committee, specific ratio E/H and/or ratio H/E percentiles are adjusted using adjustment factors for exposure and hazard, e.g. from expert elicitation. They may be available as fixed values or as parametric uncertainty distributions. In the nominal run, the percentile is adjusted with the median of the uncertainty distribution. In each uncertainty run, adjustment factors are sampled from the uncertainty distribution. In the MCRA interface, for both exposure and hazard distribution separately, a fixed value or a parametric uncertainty distribution is specified. The available parametric uncertainty distributions are the same as available in the SHELF package that was used by EFSA. The SHeffield ELicitation Framework (SHELF) is a package of documents, templates and software to carry out elicitation of probability distributions for uncertaint quantities from a group of experts (http://www.tonyohagan.co.uk/shelf/).

To summarize, in the nominal run, the median of the uncertainty distribution is taken as adjustment factor. In each uncertainty run, the adjustment factor value is sampled from the uncertainty distribution. The adjustment factors for exposure and hazard are multiplied yielding the overall adjustment factor.

Options for specifying uncertainty distributions are:

- Gamma(a, b) with offset c, with shape and rate parameters a and b > 0.
- Beta(a, b) scaled to the interval [c, d], with shape parameters a and b > 0.
- Lognormal(μ , s) with offset c. Parameters μ and s specify the mean and standard deviation of the underlying normal.

 Log Student t(μ, s, ν) with offset d. Parameters μ and s specify the mean and standard deviation of the underlying normal, ν the degrees of freedom, ν > 0.

In the figures below, the consensus distributions of the experts for the combined impact of the quantified uncertainties affecting exposure are shown. When the larger part of the distribution is above 1, it is more likely that resolving the uncertainties will increase the median estimate. The vertical red line marks f = 1, where resolving the uncertainties would not change the calculated estimate.



Figure 3.89: Scaled gamma (a=3.26, b=3.56, offset=0.9), table 6, EFSA (2020a).

Background-only adjustment factor

When exposures are calculated by *combining focal food/substance concentrations with background concentrations*, it may be appropriate to have a separate adjustment for the foreground and background. A pragmatic solution agreed with EFSA is to estimate the contribution of the foreground in the tail above the selected percentile. Suppose this contribution is c. Note that c will vary in uncertainty runs. Then, the adjustment factor should be multiplied by (1-c), i.e. no adjustment for the focal part.

The calculation proceeds as follows:

$$\begin{array}{ll} p_{H/E, \mathrm{adjusted}} &= p_{H/E} \cdot (c + (1 - c) \cdot \mathrm{AdjustmentFactor}_{\mathrm{exposure}} \cdot \mathrm{AdjustmentFactor}_{\mathrm{hazard}}) \\ p_{E/H, \mathrm{adjusted}} &= \frac{p_{E/H}}{c + (1 - c) \cdot \mathrm{AdjustmentFactor}_{\mathrm{exposure}} \cdot \mathrm{AdjustmentFactor}_{\mathrm{hazard}}} \end{array}$$

Note that when the focal substance measurements are converted to active substances using *substance conversions* or *deterministic substance conversions*, then *c* is the sum of the contributions of the focal food in and all active substances to which the substance translates.

In Figure 3.93, an example is shown where the risk characterisation ratio (hazard/exposure) is adjusted for the exposure and hazard distribution based on expert elicitation. The median adjustment factors for exposure and hazard are respectively, 1.77 and 3.01. The overall adjustment factor is 5.33.



Figure 3.90: Scaled beta (a=2.37, b=4.26, lowerbound=0.5, upperbound=6), table 7, EFSA (2020a).



Figure 3.91: Scaled lognormal ($\mu = 0.705$, s = 0.566, offset=1), table 8, EFSA (2020b).



Figure 3.92: Scaled logstudents t ($\mu = -0.593$, s = 0.367, $\nu = 3$, offset=0.5), table 9, EFSA (2020b).



Figure 3.93: Risk characterisation ratio H/E (model) and adjusted ratio H/E (model + expert) with uncertainty bounds.

Single value risks settings

Calculation settings

Name	Туре	Description
Selected tier	SettingsTemplateType	Specifies all module settings should be set according to a pre-defined tier or using custom settings.
Single value risk calculation method	SingleValueRiskCalculation- Method	Calculate single value from exposures and hazard or from an individual risks distribution.
Exposure type	ExposureType	The type of exposure considered in the assessment; acute (short term) or chronic (long-term)
Multiple substances analysis	Boolean	Specifies whether the assessment involves multiple substances.
Compute cumulative exposures	Boolean	Specifies whether the assessment involves multiple substances a results should be cumulated over all substances.
Health effect type	HealthEffectType	Specifies whether the health effect is a risk (negative) or benefit (positive).
Risk characterisation ratio	RiskMetricType	Report risks in terms of the ratio exposure/hazard (e.g., HI, HQ RPI) or as hazard/exposure (e.g., MOE(T)).
Percentage for percentile	Numeric	Percentage for percentile (default 0.1 for MOE(T) or 99.9 for H HQ, RPI).
Use inverse distribution to calculate percentile	Boolean	Calculate percentile via the complementary percentage of the inverse distribution (default: no). Description: E.g., P0.1 of MOE(T) distribution is calculated via P99.9 of 1/MOE(T) distribution. Note: This option is provided because percentile calculation in small data sets is asymmetric in both tails.
Apply adjustment factors to the specified risk percentile	Boolean	Specify adjustment factors, e.g. based on expert knowledge elicitation, to a specified MOE(T) percentile (default 0.1%). If selected risk metric is HI, HQ, RPI, the adjustment factors shot still be specified for the complementary percentile of MOE(T) (e.g. P0.1 of MOE(T) if P99.9 of HI, HQ, RPI is selected).
Adjustment type related to exposure	AdjustmentFactorDistribution- Method	Specify the factor and/or distribution of the adjustment factor f the MOE(T) percentile. Default is no adjustment. Alternatives a fixed factor or an uncertainty distribution. If distributions are selected, default values are set based on EFSA cumulative risk reports 2020.
Parameter A (Fixed factor, mean Lognormal or LogStudent-t, or shape parameter Beta or Gamma)	Numeric	This parameter can be: 1) the fixed adjustment factor; 2) for Lognormal or LogStudent-t, the mean of the underlying normal distribution; 3) For Beta or Gamma. the shape parameter.
Parameter B (standard deviation Lognormal or LogStudent-t or second shape parameter Beta or rate parameter Gamma)	Numeric	This parameter can be: 1) for Lognormal or LogStudent-t, the standard deviation of the underlying normal distribution; 2) For Beta, the second shape parameter; 3) for Gamma, the rate parameter.
Parameter C (Lower bound Beta, offset Gamma or Lognormal or degrees of freedom Logstudent-t)	Numeric	This parameter can be: 1) for Beta, the lower bound value; 2) for Gamma or Lognormal, the offset; 3) for LogStudent-t, the degree of freedom.
Parameter D (Upper bound Beta or offset LogStudent-t)	Numeric	This parameter can be: 1) for Beta, the upper bound value; 2) f LogStudent-t, the offset.
Adjustment type related to hazard	AdjustmentFactorDistribution- Method	Specify the factor and/or distribution of the adjustment factor f the MOE(T) percentile. Default is no adjustment. Alternatives a fixed factor or an uncertainty distribution. If distributions are selected, default values are set based on EFSA cumulative risk reports 2020.
Parameter A (Fixed factor, mean Lognormal or LogStudent-t, or shape parameter Beta or Gamma)	Numeric	This parameter can be: 1) the fixed adjustment factor; 2) for Lognormal or LogStudent-t, the mean of the underlying normal distribution; 3) For Beta or Gamma. the shape parameter.
Parameter B (standard 3.8. Hisk modules deviation Lognormal or LogStudent-t or second shape parameter Beta or rate parameter Gamma)	Numeric	This parameter can be: 1) for Lognormal or LogStudent-t, the standard deviation of the underlying normal distribution; 2) For Beta, the second shape parameter; 3) for Gamma, the rate parameter.

Table 3.421: Calculation settings for module Single value risks.

Uncertainty settings

Table 3.422: Uncertainty settings for module Single value risks.

Name	Туре	Description
Lower uncertainty limit (%)	Numeric	Percentage lower bound, e.g. 2.5%.
Upper uncertainty limit (%)	Numeric	Percentage upper bound, e.g. 97.5%.

Single value risks tiers

Overview

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFSA 2022 Acute Tier II	EFSA 2022 Chronic Tier II	EFSA 2023 Acute Prospec- tive Tier II	EFS 202 Chi Pro tive II
S gl va ri ca cu la ti m	FromIndi- vidual- Risks	FromIndi- vidual- Risks	FromIndi- vidual- Risks	FromIndi- vidual- Risks	FromIndi- vidual- Risks	FromIndi- vidual- Risks	FromIndi- vidual- Risks	FromIndi- vidual- Risks	FromIndi- vidual- Risks	Fro vidu Risl
P ce aş fc p ce	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1
Ig no sa pl w	true	true	true	true	true	true	true	true	true	true
E cl ir di vi u al w le th N di	false	true	false	true	false	true	false	true	false	true

Table 3.423: Tier overview for module Single value risks.

							· ·			
N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFSA 2022 Acute Tier II	EFSA 2022 Chronic Tier II	EFSA 2023 Acute Prospec- tive Tier II	EFS 202 Chro Pros tive II
T ta di st co co tr ti di	false	false	false	false	false	false	false	false	false	false
F tee sa p c c c c irr th c c c c t tr ti t ir ti ti ti ti ti ti ti ti ti ti ti ti ti	false	false	false	false	false	false	false	false	true	true
U st co vo si rt	true	true	true	true	true	true	true	true	true	true
S st co vo si n	UseMost- Toxic	UseMost- Toxic	DrawRan- dom	DrawRan- dom	UseMost- Toxic	UseMost- Toxic	DrawRan- dom	DrawRan- dom	DrawRan- dom	Drav dom

Table 3.423 - continued from previous page

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFSA 2022 Acute Tier II	EFSA 2022 Chronic Tier II	EFSA 2023 Acute Prospec- tive Tier	EFS 2023 Chros Pros tive
R ta al al cca su st at te aa ti su st al lcc ca ti al cca su st at te ad te al cca su su st at al cca su su su su su su su su su su su su su	true	true	true	true	true	true	true	true	true	true
A co fc su st au th ri sz ti ir su st co vo si	false	false	true	true	false	false	true	true	true	true
F d p ca su st al lc ca ti ir ca si ti ti te ci	false	false	false	false	false	false	false	false	false	false

Table 3.423 - continued from previous page

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N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFSA 2022 Acute Tier II	EFSA 2022 Chronic Tier II	EFSA 2023 Acute Prospec- tive Tier II	EFS 202: Chros tive II
U ez tr o- la ti rı	true	true	true	true	false	false	false	false	false	false
T ol fc ez tr o- la ti	10	10	10	10						
R st ez tr o- la ti tc ec N	true	true	true	true						
R st ez tr o- la ti tc au tt ri u	true	true	true	true						
Ir pi w te co co tr	true	true	true	true	true	true	true	true	true	true

Table 3.423 - continued from previous page

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFSA 2022 Acute Tier II	EFSA 2022 Chronic Tier II	EFSA 2023 Acute Prospec- tive Tier II	EFS 202 Chro Pros tive
W te ce til va (µ	0.1	0.1	0.05	0.05	0.1	0.1	0.05	0.05	0.05	0.05
R st w te in p ta tiv to th fr m to su st	true	true	true	true	true	true	true	true	true	true
R stt w te in pu ta ti to au th ri	false	false	false	false	false	false	false	false	false	false
R st w te in pl ta tiv to aj pl st	false	false	false	false	true	true	true	true	true	true

Table 3.423 - continued from previous page

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	N EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFSA 2022 Acute Tier II	EFSA 2022 Chronic Tier II	EFSA 2023 Acute Prospec- tive Tier II	EFS 202: Chros Pros tive
1 1 1 1 1 1	A true pl pl co in fa tc	true	true	true	true	true	true	true	true	true
1 (1 1	U false di tr bi ti	false	false	false	false	false	false	false	false	false
1 1 1 1 1 1 1 1 1	Ig false pi co fa fa le th 1	false	false	false	false	false	false	false	false	false
1 3 1	U BetaDis- va tribution al it		BetaDis- tribution		BetaDis- tribution		BetaDis- tribution		BetaDis- tribution	
] 1 1 1	E Realistic ti m ni tu		Realistic		Realistic		Realistic		Realistic	
	U Variabili- va tyFactor al it pa ra e- te		Variabili- tyFactor		Variabili- tyFactor		Variabili- tyFactor		Variabili- tyFactor	
	A false pl ci re pi te pi ci ag	false	true	true	false	false	true	true	true	true

Table 3.423 - continued from previous page

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFSA 2022 Acute Tier II	EFSA 2022 Chronic Tier II	EFSA 2023 Acute Prospec- tive Tier II	EFS 202 Chro Pros tive II
T ge le	External	External	External	External	External	External	External	External	External	Exte
C pr cu m la ti ri	true	true	true	true	true	true	true	true	true	true
H ef fe	Risk	Risk	Risk	Risk	Risk	Risk	Risk	Risk	Risk	Risk
R cl ac te i- sa ti ra	Hazard- Exposur- eRatio	Hazard- Exposur- eRatio	Hazard- Exposur- eRatio	Hazard- Exposur- eRatio	Hazard- Exposur- eRatio	Hazard- Exposur- eRatio	Hazard- Exposur- eRatio	Hazard- Exposur- eRatio	Hazard- Exposur- eRatio	Haza Expo eRat
R th ol	100	100	100	100	100	100	100	100	100	100
U in ve di tr bi tc ca ca ca ca ca ca ca ca ca ca ca ca ca	true	true	true	true	true	true	true	true	true	true
E po su ca ci la tio m	Modelled- Concen- tration	Modelled- Concen- tration	Modelled- Concen- tration	Modelled- Concen- tration	Modelled- Concen- tration	Modelled- Concen- tration	Modelled- Concen- tration	Modelled- Concen- tration	Modelled- Concen- tration	Mod Con tratic

Table 3.423 - continued from previous page

N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFSA 2022 Acute Tier II	EFSA 2022 Chronic Tier II	EFSA 2023 Acute Prospec- tive Tier II	EFS 202 Chros tive II
In cl si p cu ag v a al it it it te	99.8	99.8	99.8	99.8	99.8	99.8	99.8	99.8	99.8	99.8
C su ti o tt sa d c c fr fr tt sa sa p	false	false	false	false	false	false	false	false	false	false
U u v al it	true		true		true		true		true	
N th	false	false	false	false	false	false	false	false	false	false
C Va at m el	false	false	false	false	false	false	false	false	false	false
It en at su v	false	false	false	false	false	false	false	false	false	false

Table 3.423 - continued from previous page

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N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFSA 2022 Acute Tier II	EFSA 2022 Chronic Tier II	EFSA 2023 Acute Prospec- tive Tier II	EFS 202 Chro Pros tive
R P c c s t i an e : P f s t i an e : P f s t i an e : P f s t i an e : P f s t i an e : S t i s t i an e : S t i s t i s t i s t i s t i s t i s t i s t i s t i s t i s t i s t i s t i s t i s s t i s s t i s s t i s s t i s s t i s s t i s s t i s s t i s s t i s s s t i s s s t i s s s s	false	false	false	false	false	false	false	false	false	false
D fa co tr ti n	Empirical	Empirical	Empirical	Empirical	Empirical	Empirical	Empirical	Empirical	Empirical	Emp
In cl M fa ba n	false	false	false	false	false	false	false	false	false	false
R stt L ir P ta ti tc au th ri u	false	false	false	false	false	false	false	false	false	false

Table 3.423 - continued from previous page

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	N EC 2018 Acute Tier I	B EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFSA 2022 Acute Tier II	EFSA 2022 Chronic Tier II	EFSA 2023 Acute Prospec- tive Tier II	EFS 2023 Chros Pros tive II
	C Replace- sc ByLOR va u re pl m	Replace- ByLOR	Replace- ByLOR	Replace- ByLOR	Replace- ByLOR	Replace- ByLOR	Replace- ByLOR	Replace- ByLOR	Replace- ByLOR	Repl ByL
	F 0.5 tc f (f x L oi f x L oi f x L oi f x t L - L	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5	0.5
	S true pl b	true	true	true	true	true	true	true	true	true

Table 3.423 - continued from previous page

N EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFSA 2022 Acute Tier II	EFSA 2022 Chronic Tier II	EFSA 2023 Acute Prospec- tive Tier II	EFS 202 Chro Pros tive
Ir true p r r v v u u f r a v a u u u u u u u u u u u u u	true	true	true	true	true	true	true	true	true
C true re la ir p va u w sa p p p te	true	false	false	true	true	false	false	false	false
U true o c r f f f c i f c i f c i t z t i t	true	true	true	true	true	true	true	true	true

Table 3.423 - continued from previous page
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N	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFSA 2022 Acute Tier II	EFSA 2022 Chronic Tier II	EFSA 2023 Acute Prospec- tive Tier II	EFS 2023 Chros Pros tive II
P m ri u ce ta	false	false	false	false	false	false	false	false	false	false
N (r b) of di ir st v		2		2		2		2		2
M ty		OIM		OIM		OIM		OIM		OIM
S uj u fr q tc 1			true	true			true	true	true	true
R st us p c c a a t t t t t t us			true	true			true	true	true	true
C ce tr ti lii fi te ez ce fa									2	2

Table 3.423 - continued from previous page

N EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFSA 2022 Acute Tier II	EFSA 2022 Chronic Tier II	EFSA 2023 Acute Prospec- tive Tier II	EFS 2023 Chros Pros tive II
Ir cl fc ca ca m it it ca ca tr tr ti								true	true
F ca ca n n it su st co ca re p ca ag								20	20
A ju m fa tc fa tc fa ca fa ca tr ti								1	1

Table 3.423 - continued from previous page

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1	EC 2018 Acute Tier I	EC 2018 Chronic - Tier I	EC 2018 Acute Tier II	EC 2018 Chronic Tier II	EFSA 2022 Acute Tier I	EFSA 2022 Chronic Tier I	EFSA 2022 Acute Tier II	EFSA 2022 Chronic Tier II	EFSA 2023 Acute Prospec- tive Tier II	EFS 202 Chro Pros tive II
Ucctriiiittissccfffcccrrii	U de r s s s s t c c c c c c c c c c c c c c c								true	true
II cc cc tt ff ff ii r r r r r r r r r	Ir El Do I I I I I I I I I I I I I I I I I I									

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Retrospective dietary CRA (EC 2018) - Acute / Tier I

Name	Setting	From input tier	In module
Single value risk calculation method	FromIndi- vidual- Risks		
Percentage for percentile	0.1		
Ignore sampling weights	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Consump- tions
Exclude individuals with less than N days	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations

Table 3.424: Tier definition for Retrospective dietary CRA (EC 2018) - Acute / Tier I.

Name	Setting	From input tier	ln module
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Account for substance authorisations in substance conversions	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Use extrapolation rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Threshold for extrapolation	10	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Restrict extrapolations to equal MRLs	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Restrict extrapolations to authorised uses	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations

	Table	3.424 -	continued	from	previous	page
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Name	Setting	From input tier	In module
Impute water concentrations	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Water concentration value (µg/kg)	0.1	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Restrict water imputation to approved substances	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Processing factors

	Table	3.424 -	continued	from	previous	page
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Name	Setting	From input tier	ln module
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Processing factors
Unit variability model	BetaDis- tribution	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Unit variability factors
Estimates nature	Realistic	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Unit variability factors
Unit variability parameter	Variabili- tyFactor	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Unit variability factors
Apply occurrence pattern percentages	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Occur- rence patterns
Target level	External	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Hazard character- isations
Compute cumulative risks	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Risks

Table 3.424 – continued from p	previous pa	ige
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Name	Setting	From input tier	ln module
Health effect type	Risk	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Risks
Risk characterisation ratio	Hazard- Exposur- eRatio	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Risks
Risk threshold	100	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Risks
Use inverse distribution to calculate percentile	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Risks
Exposure calculation method	Modelled- Concen- tration	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Risks
Inclusion percentage variability interval	99.8	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Risks
Consumptions on the same day come from the same sample	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Dietary exposures

Table 3.424 - continued from previous page

Name	Setting	From input tier	In module
Use unit variability	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Dietary exposures
Model-then-add	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Dietary exposures
Covariate modelling	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Dietary exposures
Iterate survey	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Dietary exposures
Default concentration model	Empirical	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models

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Name	Setting	From input tier	ln module
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Sample based	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Correlate imputed values with sample potency	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models

Table 3.424 - continued from previous page

Name	Setting	From input tier	ln module
Parametric uncertainty	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier I	Concen- tration models

Table 3.424 - continued from previous page

Retrospective dietary CRA (EC 2018) - Chronic / Tier I

Table 3.425:	Tier	definition	for	Retrospective	dietary	CRA	(EC	2018) -
Chronic / Tie	er I.							

Name	Setting	From input tier	ln module
Single value risk calculation method	FromIndi- vidual- Risks		
Percentage for percentile	0.1		
Ignore sampling weights	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Consump- tions
Exclude individuals with less than N days	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Consump- tions
N (number of days in survey)	2	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Food con- versions

Name	Setting	From input tier	In module
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Account for substance authorisations in substance conversions	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Use extrapolation rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations

Table 3.425 - continued from previous page

Name	Setting	From input tier	In module
Threshold for extrapolation	10	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Restrict extrapolations to equal MRLs	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Restrict extrapolations to authorised uses	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Impute water concentrations	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Water concentration value (µg/kg)	0.1	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations

Table 3.425 - continued from previous page

Name	Setting	From input tier	In module
Restrict water imputation to approved substances	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Processing factors
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Processing factors
Apply occurrence pattern percentages	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Occur- rence patterns
Target level	External	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Hazard character- isations
Compute cumulative risks	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Risks

Table 3.425 - continued from previous page

Name	Setting	From input tier	ln module
Health effect type	Risk	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Risks
Risk characterisation ratio	Hazard- Exposur- eRatio	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Risks
Risk threshold	100	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Risks
Use inverse distribution to calculate percentile	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Risks
Exposure calculation method	Modelled- Concen- tration	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Risks
Inclusion percentage variability interval	99.8	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Risks
Consumptions on the same day come from the same sample	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Dietary exposures

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Name	Setting	From input tier	In module
Model type	OIM	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Dietary exposures
Model-then-add	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Dietary exposures
Covariate modelling	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Dietary exposures
Iterate survey	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Dietary exposures
Default concentration model	Empirical	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models

Table 3.425 - continued from previous page

Name	Setting	From input tier	ln module
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Sample based	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Correlate imputed values with sample potency	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models

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Name	Setting	From input tier	ln module
Parametric uncertainty	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier I	Concen- tration models

Table 3.425 - continued from previous page

Retrospective dietary CRA (EC 2018) - Acute / Tier II

Table 3.426: Tier definition for Retrospective dietary CRA (EC 2018) - Acute / Tier II.

Name	Setting	From input tier	ln module
Single value risk calculation method	FromIndi- vidual- Risks		
Percentage for percentile	0.1		
Ignore sampling weights	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Consump- tions
Exclude individuals with less than N days	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations

Name	Setting	From input tier	ln module
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Use extrapolation rules	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Threshold for extrapolation	10	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations

Table 3.426 – continued from previous page

Name	Setting	From input tier	In module
Restrict extrapolations to equal MRLs	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Restrict extrapolations to authorised uses	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Impute water concentrations	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations
Restrict water imputation to approved substances	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- trations

Table	3.426 -	continued	from	previous	page

Name	Setting	From input tier	In module
Apply processing factors	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Processing factors
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Processing factors
Unit variability model	BetaDis- tribution	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Unit variability factors
Estimates nature	Realistic	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Unit variability factors
Unit variability parameter	Variabili- tyFactor	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Unit variability factors
Apply occurrence pattern percentages	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Occur- rence patterns

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Name	Setting	From input tier	In module
Scale up use frequency to 100%	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Occur- rence patterns
Target level	External	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Hazard character- isations
Compute cumulative risks	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Risks
Health effect type	Risk	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Risks
Risk characterisation ratio	Hazard- Exposur- eRatio	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Risks
Risk threshold	100	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Risks

Table	3.426 -	continued	from	previous	page

Name	Setting	From input tier	ln module
Use inverse distribution to calculate percentile	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Risks
Exposure calculation method	Modelled- Concen- tration	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Risks
Inclusion percentage variability interval	99.8	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Risks
Consumptions on the same day come from the same sample	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Dietary exposures
Use unit variability	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Dietary exposures
Model-then-add	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Dietary exposures
Covariate modelling	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Dietary exposures

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Name	Setting	From input tier	ln module
Iterate survey	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Dietary exposures
Default concentration model	Empirical	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models

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Name	Setting	From input tier	ln module
Sample based	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Correlate imputed values with sample potency	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models
Parametric uncertainty	false	Retrospec- tive dietary CRA (EC 2018) - Acute / Tier II	Concen- tration models

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Retrospective dietary CRA (EC 2018) - Chronic / Tier II

Table 3.427: Tier definition for Retrospective dietary CRA (EC 2018) - Chronic / Tier II.

Name	Setting	From input tier	ln module
Single value risk calculation method	FromIndi- vidual- Risks		
Percentage for percentile	0.1		

Name	Setting	From input tier	ln module
Ignore sampling weights	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Consump- tions
Exclude individuals with less than N days	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Consump- tions
N (number of days in survey)	2	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations

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Name	Setting	From	In
		input tier	module
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Use extrapolation rules	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Threshold for extrapolation	10	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Restrict extrapolations to equal MRLs	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Restrict extrapolations to authorised uses	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations

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Name	Setting	From input tier	ln module
Impute water concentrations	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Restrict water imputation to approved substances	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Processing factors

Table 3.427 – continued from previous page

Name	Setting	From	In module
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Processing factors
Apply occurrence pattern percentages	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Occur- rence patterns
Target level	External	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Hazard character- isations
Compute cumulative risks	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Risks
Health effect type	Risk	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Risks

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3.8. Risk modules

Name	Setting	From input tier	In module
Risk characterisation ratio	Hazard- Exposur- eRatio	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Risks
Risk threshold	100	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Risks
Use inverse distribution to calculate percentile	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Risks
Exposure calculation method	Modelled- Concen- tration	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Risks
Inclusion percentage variability interval	99.8	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Risks
Consumptions on the same day come from the same sample	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Dietary exposures
Model type	OIM	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Dietary exposures

Table 3.427 - continued from previous page

Name	Setting	From input tier	ln module
Model-then-add	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Dietary exposures
Covariate modelling	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Dietary exposures
Iterate survey	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Dietary exposures
Default concentration model	Empirical	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models

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Name	Setting	From input tier	In module
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Sample based	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Correlate imputed values with sample potency	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models
Parametric uncertainty	false	Retrospec- tive dietary CRA (EC 2018) - Chronic / Tier II	Concen- tration models

Table 3.427 - continued from previous page

Retrospective dietary CRA (EFSA 2022) - Acute / Tier I

Name	Setting	From input tier	In module
Single value risk calculation method	FromIndi- vidual- Risks		
Percentage for percentile	0.1		
Ignore sampling weights	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Consump- tions
Exclude individuals with less than N days	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations

Table 3.428: Tier definition for Retrospective dietary CRA (EFSA 2022) - Acute / Tier I.

Name	Setting	From input tier	ln module
Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Account for substance authorisations in substance conversions	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Use extrapolation rules	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Impute water concentrations	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations

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Name	Setting	From input tier	In module
Water concentration value (µg/kg)	0.1	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Restrict water imputation to approved substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Processing factors

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Name	Setting	From input tier	ln module
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Processing factors
Unit variability model	BetaDis- tribution	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Unit variability factors
Estimates nature	Realistic	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Unit variability factors
Unit variability parameter	Variabili- tyFactor	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Unit variability factors
Apply occurrence pattern percentages	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Occur- rence patterns
Target level	External	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Hazard character- isations

Table 3.428 - continued from previous page
Name	Setting	From input tier	In module
Compute cumulative risks	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Risks
Health effect type	Risk	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Risks
Risk characterisation ratio	Hazard- Exposur- eRatio	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Risks
Risk threshold	100	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Risks
Use inverse distribution to calculate percentile	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Risks
Exposure calculation method	Modelled- Concen- tration	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Risks

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Name	Setting	From input tier	In module
Inclusion percentage variability interval	99.8	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Risks
Consumptions on the same day come from the same sample	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Dietary exposures
Use unit variability	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Dietary exposures
Model-then-add	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Dietary exposures
Covariate modelling	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Dietary exposures
Iterate survey	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Dietary exposures

Table 3.428 - continued from previous page

Name	Setting	From input tier	In module
Report consumptions and exposures per individual instead of per kg body weight	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Dietary exposures
Default concentration model	Empirical	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models

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Name	Setting	From input tier	ln module
Sample based	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Correlate imputed values with sample potency	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models
Parametric uncertainty	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier I	Concen- tration models

Table 3.428 - continued from previous page

Retrospective dietary CRA (EFSA 2022) - Chronic / Tier I

Table 3.429: Tier definition for Retrospective dietary CRA (EFSA 2022) - Chronic / Tier I.

Name	Setting	From input tier	In module
Single value risk calculation method	FromIndi- vidual- Risks		
Percentage for percentile	0.1		

Name	Setting	From input tier	In module
Ignore sampling weights	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Consump- tions
Exclude individuals with less than N days	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Consump- tions
N (number of days in survey)	2	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations

Table 3.429 – continued from previou	us pade
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Name	Setting	From input tier	ln module
Substance conversion method	UseMost- Toxic	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Account for substance authorisations in substance conversions	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Use extrapolation rules	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Impute water concentrations	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations

Table 3.429 - continued from previous page

Name	Setting	From input tier	ln module
Water concentration value (µg/kg)	0.1	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Restrict water imputation to approved substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Processing factors

Table 3.429 – continued from previous page
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Name	Setting	From input tier	ln module
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Processing factors
Apply occurrence pattern percentages	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Occur- rence patterns
Target level	External	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Hazard character- isations
Compute cumulative risks	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Risks
Health effect type	Risk	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Risks
Risk characterisation ratio	Hazard- Exposur- eRatio	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Risks

Table 3.429 - continued from previous page

Name	Setting	From input tier	In module
Risk threshold	100	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Risks
Use inverse distribution to calculate percentile	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Risks
Exposure calculation method	Modelled- Concen- tration	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Risks
Inclusion percentage variability interval	99.8	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Risks
Consumptions on the same day come from the same sample	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Dietary exposures
Model type	OIM	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Dietary exposures

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Name	Setting	From input tier	In module
Model-then-add	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Dietary exposures
Covariate modelling	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Dietary exposures
Iterate survey	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Dietary exposures
Default concentration model	Empirical	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models

Table 3.429 - continued from previous page

Name	Setting	From input tier	In module
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Sample based	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Correlate imputed values with sample potency	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models

Table 3.429 – continued from previous pa	bade
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Name	Setting	From input tier	ln module
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models
Parametric uncertainty	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier I	Concen- tration models

Table 3.429 – continued from previous page

Retrospective dietary CRA (EFSA 2022) - Acute / Tier II

Table 3.430:	Tier	definition	for Ret	rospective	dietary	CRA	(EFSA	2022)
- Acute / Tier	II.							

Name	Setting	From input tier	In module
Single value risk calculation method	FromIndi- vidual- Risks		
Percentage for percentile	0.1		
Ignore sampling weights	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Consump- tions
Exclude individuals with less than N days	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Food con- versions

Name	Setting	From input tier	In module
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations

Table 3.430 - continued from previous page

Name	Setting	From input tier	ln module
Use extrapolation rules	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Impute water concentrations	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations
Restrict water imputation to approved substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- trations

Table 3.430 - continued from previous page

Name	Setting	From input tier	ln module
Apply processing factors	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Processing factors
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Processing factors
Unit variability model	BetaDis- tribution	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Unit variability factors
Estimates nature	Realistic	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Unit variability factors
Unit variability parameter	Variabili- tyFactor	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Unit variability factors

Table 3.430 – continued from previous page

Name	Setting	From input tier	In module
Apply occurrence pattern percentages	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Occur- rence patterns
Target level	External	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Hazard character- isations
Compute cumulative risks	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Risks
Health effect type	Risk	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Risks

Table 3.430 - continued from previous page

Name	Setting	From input tier	ln module
Risk characterisation ratio	Hazard- Exposur- eRatio	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Risks
Risk threshold	100	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Risks
Use inverse distribution to calculate percentile	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Risks
Exposure calculation method	Modelled- Concen- tration	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Risks
Inclusion percentage variability interval	99.8	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Risks
Consumptions on the same day come from the same sample	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Dietary exposures

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Name	Setting	From input tier	ln module
Use unit variability	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Dietary exposures
Model-then-add	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Dietary exposures
Covariate modelling	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Dietary exposures
Iterate survey	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Dietary exposures
Default concentration model	Empirical	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models

Table 3.430 - continued from previous page

Name	Setting	From input tier	ln module
Include MRL fallback model	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Sample based	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models

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Name	Setting	From input tier	In module
Correlate imputed values with sample potency	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models
Parametric uncertainty	false	Retrospec- tive dietary CRA (EFSA 2022) - Acute / Tier II	Concen- tration models

Table 3.430 - continued from previous page

Retrospective dietary CRA (EFSA 2022) - Chronic / Tier II

Table 3.431:	Tier defi	inition for	Retrospective	dietary	CRA	(EFSA	2022)
- Chronic / T	ier II.						

Name	Setting	From input tier	ln module
Single value risk calculation method	FromIndi- vidual- Risks		
Percentage for percentile	0.1		
Ignore sampling weights	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Consump- tions
Exclude individuals with less than N days	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Consump- tions

Name	Setting	From input tier	ln module
N (number of days in survey)	2	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Consump- tions
Total diet study concentration data	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Food con- versions
Filter samples exceeding the concentration limits	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Use substance conversion rules	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations

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Name	Setting	From input tier	ln module
Account for substance authorisations in substance conversions	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Use extrapolation rules	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Impute water concentrations	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations

Table 3.431 - continued from previous page

Name	Setting	From input tier	In module
Restrict water imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Restrict water imputation to approved substances	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- trations
Apply processing factors	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Processing factors
Use distribution	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Processing factors
Ignore processing factors less than 1	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Processing factors
Apply occurrence pattern percentages	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Occur- rence patterns

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Name	Setting	From input tier	ln module
Scale up use frequency to 100%	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Occur- rence patterns
Target level	External	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Hazard character- isations
Compute cumulative risks	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Risks
Health effect type	Risk	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Risks
Risk characterisation ratio	Hazard- Exposur- eRatio	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Risks

Table 3.431 - continued from previous page

Name	Setting	From input tier	In module
Risk threshold	100	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Risks
Use inverse distribution to calculate percentile	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Risks
Exposure calculation method	Modelled- Concen- tration	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Risks
Inclusion percentage variability interval	99.8	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Risks
Consumptions on the same day come from the same sample	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Dietary exposures
Model type	OIM	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Dietary exposures

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Name	Setting	From input tier	ln module
Model-then-add	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Dietary exposures
Covariate modelling	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Dietary exposures
Iterate survey	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Dietary exposures
Default concentration model	Empirical	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Include MRL fallback model	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models

Table 3.431 - continued from previous page

Name	Setting	From input tier	In module
Restrict LOR imputation to authorised uses	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Sample based	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Correlate imputed values with sample potency	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models

Name	Setting	From input tier	ln module
Use occurrence frequencies for imputation	true	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models
Parametric uncertainty	false	Retrospec- tive dietary CRA (EFSA 2022) - Chronic / Tier II	Concen- tration models

Table 3.431 – continued from previous page

Prospective dietary CRA (EFSA 2023) - Acute / Tier II

Table 3.432: Tier definition for Prospective dietary CRA (EFSA 2023) - Acute / Tier II.

Name	Setting	From input tier	ln module
Single value risk calculation method	FromIndi- vidual- Risks		
Percentage for percentile	0.1		
Ignore sampling weights	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Consump- tions
Exclude individuals with less than N days	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Consump- tions
Total diet study concentration data	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Food con- versions

Name	Setting	From input tier	In module
Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Fix duplicate substance allocation inconsistencies	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Use extrapolation rules	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Impute water concentrations	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations

Table 3.432 - continued from previous page

Name	Setting	From input tier	ln module
Restrict water imputation to approved substances	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Include focal commodity concentrations	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Focal commodity substance occurrence percentage	20	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Adjustment factor for the focal food/substance concentration	1	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Use deterministic substance conversions for focal commodity	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- trations
Apply processing factors	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Processing factors

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Name	Setting	From input tier	In module
Use distribution	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Processing factors
Ignore processing factors less than 1	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Processing factors
Unit variability model	BetaDis- tribution	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Unit variability factors
Estimates nature	Realistic	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Unit variability factors
Unit variability parameter	Variabili- tyFactor	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Unit variability factors
Apply occurrence pattern percentages	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Occur- rence patterns

Table 3.432 - continued from previous page

Name	Setting	From input tier	ln module
Scale up use frequency to 100%	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Occur- rence patterns
Target level	External	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Hazard character- isations
Compute cumulative risks	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Risks
Health effect type	Risk	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Risks
Risk characterisation ratio	Hazard- Exposur- eRatio	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Risks

Table 3.432 – continued from previous pa	ade
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Name	Setting	From input tier	In module
Risk threshold	100	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Risks
Use inverse distribution to calculate percentile	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Risks
Exposure calculation method	Modelled- Concen- tration	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Risks
Cumulation setting	RPFWeighte	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Risks
Inclusion percentage variability interval	99.8	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Risks
Consumptions on the same day come from the same sample	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Dietary exposures

Table 3.432 - continued from previous page

Name	Setting	From input tier	In module
Use unit variability	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Dietary exposures
Model-then-add	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Dietary exposures
Covariate modelling	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Dietary exposures
Iterate survey	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Dietary exposures
Default concentration model	Empirical	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models

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Name	Setting	From input tier	ln module
Include MRL fallback model	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Sample based	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models

Table 3.432 - continued from previous page
Name	Setting	From input tier	In module
Correlate imputed values with sample potency	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models
Parametric uncertainty	false	Prospec- tive dietary CRA (EFSA 2023) - Acute / Tier II	Concen- tration models

Table 3.432 - continued from previous page

Prospective dietary CRA (EFSA 2023) - Chronic / Tier II

Table 3.433: Tier definition for Prospective dietary CRA (EFSA 2023) - Chronic / Tier II.

Name	Setting	From input tier	In module
Single value risk calculation method	FromIndi- vidual- Risks		
Percentage for percentile	0.1		
Ignore sampling weights	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Consump- tions
Exclude individuals with less than N days	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Consump- tions

Name	Setting	From input tier	In module
N (number of days in survey)	2	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Consump- tions
Total diet study concentration data	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Food con- versions
Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations

Table 3.433 - continued from previous page

Name	Setting	From input tier	In module
Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Use extrapolation rules	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Impute water concentrations	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations

Table 3.433 – continued from previous pa	ade
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Name	Setting	From input tier	In module
Restrict water imputation to the five most toxic substances	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Restrict water imputation to approved substances	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Include focal commodity concentrations	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Focal commodity substance occurrence percentage	20	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Adjustment factor for the focal food/substance concentration	1	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Use deterministic substance conversions for focal commodity	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- trations
Apply processing factors	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Processing factors
Use distribution	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Processing factors
Ignore processing factors less than 1	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Processing factors
Apply occurrence pattern percentages	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Occur- rence patterns

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Name	Setting	From input tier	In module
Restrict use percentage up-scaling to authorised uses	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Occur- rence patterns
Target level	External	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Hazard character- isations
Compute cumulative risks	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Risks
Health effect type	Risk	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Risks
Risk characterisation ratio	Hazard- Exposur- eRatio	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Risks
Risk threshold	100	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Risks

Table 3.433 - continued from previous page

Name	Setting	From input tier	ln module
Use inverse distribution to calculate percentile	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Risks
Exposure calculation method	Modelled- Concen- tration	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Risks
Cumulation setting	RPFWeight	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Risks
Inclusion percentage variability interval	99.8	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Risks
Consumptions on the same day come from the same sample	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Dietary exposures
Model type	OIM	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Dietary exposures

Table 3.433 – continued from previous page

Name	Setting	From input tier	ln module
Model-then-add	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Dietary exposures
Covariate modelling	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	<i>Dietary</i> <i>exposures</i>
Iterate survey	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Dietary exposures
Default concentration model	Empirical	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Include MRL fallback model	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models

Table 3.433 - continued from previous page

Name	Setting	From input tier	In module
Restrict LOR imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Sample based	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Correlate imputed values with sample potency	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models

	Table	3.433 -	continued	from	previous	page
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Name	Setting	From input tier	ln module
Use occurrence frequencies for imputation	true	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models
Parametric uncertainty	false	Prospec- tive dietary CRA (EFSA 2023) - Chronic / Tier II	Concen- tration models

Table 3.433 – continued from previous page

Prospective dietary CRA (EFSA 2024) - Acute / Tier II

Table 3.434: Tier definition for Prospective dietary CRA (EFSA 2024) - Acute / Tier II.

Name	Setting	From input tier	In module
Single value risk calculation method	FromIndi- vidual- Risks		
Percentage for percentile	0.1		
Ignore sampling weights	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Consump- tions
Exclude individuals with less than N days	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Consump- tions
Total diet study concentration data	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Food con- versions

Name	Setting	From input tier	In module
Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Fix duplicate substance allocation inconsistencies	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Use extrapolation rules	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Impute water concentrations	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Water concentration value (µg/kg)	0.05	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations

Table 3.434 - continued from previous page

Name	Setting	From input tier	In module
Restrict water imputation to approved substances	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Include focal commodity concentrations	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Focal commodity substance occurrence percentage	20	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Adjustment factor for the focal food/substance concentration	1	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Use deterministic substance conversions for focal commodity	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations
Include processed derivatives of the focal food in measurement replacement	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- trations

Table 3.434 – continued from previous pace
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Name	Setting	From input tier	In module
Apply processing factors	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Processing factors
Use distribution	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Processing factors
Ignore processing factors less than 1	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Processing factors
Unit variability model	BetaDis- tribution	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Unit variability factors
Estimates nature	Realistic	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Unit variability factors
Unit variability parameter	Variabili- tyFactor	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Unit variability factors

Table 3.434 - continued from previous page

Name	Setting	From	ln module
Apply occurrence pattern percentages	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Occur- rence patterns
Target level	External	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Hazard character- isations
Compute cumulative risks	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Risks
Health effect type	Risk	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Risks

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Name	Setting	From input tier	In module
Risk characterisation ratio	Hazard- Exposur- eRatio	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Risks
Risk threshold	100	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Risks
Use inverse distribution to calculate percentile	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Risks
Exposure calculation method	Modelled- Concen- tration	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Risks
Cumulation setting	RPFWeighte	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Risks
Inclusion percentage variability interval	99.8	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Risks

Table 3.434 - continued from previous page

Name	Setting	From input tier	In module
Consumptions on the same day come from the same sample	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Dietary exposures
Use unit variability	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	<i>Dietary</i> <i>exposures</i>
Model-then-add	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Dietary exposures
Covariate modelling	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Dietary exposures
Iterate survey	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Dietary exposures

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Name	Setting	From input tier	In module
Default concentration model	Empirical	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Include MRL fallback model	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Sample based	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models

Table 3.434 – continued from previous page
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Name	Setting	From input tier	In module
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Correlate imputed values with sample potency	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models
Parametric uncertainty	false	Prospec- tive dietary CRA (EFSA 2024) - Acute / Tier II	Concen- tration models

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Prospective dietary CRA (EFSA 2024) - Chronic / Tier II

Name	Setting	From input tier	ln module
Single value risk calculation method	FromIndi- vidual- Risks		
Percentage for percentile	0.1		
Ignore sampling weights	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Consump- tions

Table 3.435: Tier definition for Prospective dietary CRA (EFSA 2024) - Chronic / Tier II.

Name	Setting	From input tier	ln module
Exclude individuals with less than N days	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Consump- tions
N (number of days in survey)	2	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Consump- tions
Total diet study concentration data	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Food con- versions
Filter samples exceeding the concentration limits	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Concentration limit filter exceedance factor	2	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Use substance conversion rules	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations

Table 3.435 - continued from previous page

Name	Setting	From input tier	ln module
Substance conversion method	DrawRan- dom	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Retain all allocated substances after active substance allocation	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Account for substance authorisations in substance conversions	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Fix duplicate substance allocation inconsistencies	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Use extrapolation rules	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Impute water concentrations	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations

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Name	Setting	From input tier	In module
Water concentration value (µg/kg)	0.05	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Restrict water imputation to the five most toxic substances	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Restrict water imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Restrict water imputation to approved substances	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Include focal commodity concentrations	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Focal commodity substance occurrence percentage	20	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations

Table 3.435 - continued from previous page

Name	Setting	From input tier	ln module
Adjustment factor for the focal food/substance concentration	1	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Use deterministic substance conversions for focal commodity	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Include processed derivatives of the focal food in measurement replacement	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- trations
Apply processing factors	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Processing factors
Use distribution	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Processing factors
Ignore processing factors less than 1	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Processing factors

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Name	Setting	From input tier	In module
Apply occurrence pattern percentages	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Occur- rence patterns
Scale up use frequency to 100%	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Occur- rence patterns
Restrict use percentage up-scaling to authorised uses	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Occur- rence patterns
Target level	External	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Hazard character- isations
Compute cumulative risks	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Risks
Health effect type	Risk	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Risks

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Name	Setting	From input tier	In module
Risk characterisation ratio	Hazard- Exposur- eRatio	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Risks
Risk threshold	100	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Risks
Use inverse distribution to calculate percentile	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Risks
Exposure calculation method	Modelled- Concen- tration	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Risks
Cumulation setting	RPFWeighte	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Risks
Inclusion percentage variability interval	99.8	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Risks

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Name	Setting	From input tier	ln module
Consumptions on the same day come from the same sample	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Dietary exposures
Model type	OIM	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Dietary exposures
Model-then-add	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Dietary exposures
Covariate modelling	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Dietary exposures
Iterate survey	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Dietary exposures
Report consumptions and exposures per individual instead of per kg body weight	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Dietary exposures

Table 3.435 - continued from previous page

Name	Setting	From input tier	ln module
Default concentration model	Empirical	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Include MRL fallback model	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Restrict LOR imputation to authorised uses	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Censored values replacement	Replace- ByLOR	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Factor f (f x LOR or f x LOD or LOD + f x (LOQ - LOD)	0.5	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Sample based	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models

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Name	Setting	From input tier	In module
Impute missing values from available values (if unchecked, missing values are imputed with 0)	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Correlate imputed values with sample potency	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Use occurrence frequencies for imputation	true	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models
Parametric uncertainty	false	Prospec- tive dietary CRA (EFSA 2024) - Chronic / Tier II	Concen- tration models

Table 3.435 - continued from previous page

Calculation of single value risks

Single value risk can be computed by route and substance in the form of the risk characterisation ratio hazard/exposure or exposure/hazard. Single value risks are risk estimates obtained from combining single value exposures with single value hazard characterisations or as a percentile from a risk distribution. Optionally, the exposure and hazard behind the percentile calculation can be adjusted with fixed adjustment factors or factors drawn from parametric uncertainty distributions.

• Single value risks calculation

Inputs used: Single value dietary exposures Hazard characterisations Risks

Settings used

• Calculation Settings

842

Category	Module	Inputs	Used by	Description
	Та	ble 3.436: Over	rview of MCRA m	nodules.
Category	Module	Inputs	Used by	Description
Primary entity	Foods		Consump- tions, Singlevalue con-sumptions,Marketshares, Foodrecipes, Con- centrations,Concentra- tiondistributions,Single valueconcentra- 	Foods are uniquely defined sources of dietary exposure to chemical substances. Foods may refer to 1) foods as eaten, foods as coded in food consumption data (e.g. pizza); 2) modelled foods, foods as coded in concentration data (e.g. wheat, tomato); 3) any other type of food (e.g. ingredients like flour, tomato sauce).
			Single value	
			dietary exposures	
			Internal	

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Category	Module	Inputs	Used by	Description
	Non-dietary exposure sources			Non-dietary exposure sources are the sources containing chemical substances to which individuals in a population are exposed via any of three non-dietary routes: dermal, inhalation or oral, per day.

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Category	Module	Inputs	Used by	Description
	Substances		Concentra-	Substances are chemical
			tions,	entities that can refer to: 1)
			Concentra-	active substances such as
			tion	investigated in toxicology; 2)
			distributions,	measured substances such as
			Single value	defined in specific analytical
			concentra-	methods. MCRA assessments
			tions,	can have one or more
			Processing	substances as the scope. When
			factors, Unit	more than one substance is
			variability	specified, there is an option to
			factors,	perform a cumulative
			Occurrence	assessment. In that case one of
			patterns,	the substances has to be
			Occurrence	indicated as the index/reference
			frequencies,	substance, and results will be
			Substance	expressed in equivalents of the
			authorisa-	index substance.
			tions,	
			Substance	
			approvals,	
			Substance	
			conversions,	
			Deterministic	
			substance	
			conversion	
			factors, Con-	
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			models,	
			Modelled foods Food	
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			Dietary	
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			Single value	Obenter O. M. de
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Category	Module	Inputs	Used by	Description
	Effects		Concentra- tion models, High exposure food- substance combina- tions, Dietary exposures, Exposure mixtures, QSAR membership models, Molecular docking models, Active substances, Relative potency factors, Hazard characteri- sations, Points of departure, Effect repre- sentations, Inter-species conversions, Intra species factors, AOP networks, Exposure response functions, Burden of disease, Risks, Single value risks.	Effects are biological or toxicological consequences for human health, that may result from chemical exposure and are the focus of hazard or risk assessment.

Table	3.436 -	continued	from	previous	page
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Category	Module	Inputs	Used by	Description
	Populations		Consump- tions, Single value con- sumptions, Concentra- tions, Consump- tions by modelled food, Dietary exposures, Single value dietary exposures, Single value non-dietary exposures, Non-dietary exposures, Internal exposures, HBM con- centrations, HBM con- centrations, HBM analysis, Biological matrix con- centration comparisons, Dust exposures, Individuals, Consumer product use frequencies, Consumer product use frequencies, Consumer product exposures, Har consumer product exposures, Consumer product exposures, Consumer product exposures, Consumer product exposures, Consumer product exposures, Consumer product exposures, Consumer product exposures, Consumer product exposures, Consumer product exposures, Consumer product exposures, Consumer product exposures, Consumer product exposures, Consumer product exposures, Consumer product exposures, Consumer product exposures, Consumer product exposures, Consumer product exposures, Consumer product expos	Populations are groups of human individuals that are the scope of exposure or risk assessments. Optional descriptors of populations are location (e.g. a country), time period (with a start and end date), age range (with a minimum and maximum age) and gender. Example: the French population in 2005-2007 (= time period) of women (= gender) of child-bearing age 18-45 yr (= age range).

Table 3.436 - continued from previous page

Category	Module	Inputs	Used by	Description
	Test systems		Responses, Dose response models, Dose response data.	Test systems are biological or artificial systems used for assessing hazard in relation to chemical exposure from substances in varying doses. Test systems may refer to 1) in-vivo test systems (e.g. a rat 90-day study, a human biomonitoring study); 2) in-vitro test systems (e.g. HepaRG cells).
	Responses	Test systems.	Dose response models, Dose response data, Effect representa- tions.	Responses are measurable entities in test systems. Responses are used to represent effects (see effect representations) and their measured values are collected in dose response data.
	<i>Consumer</i> products		Consumer product con- centrations, Internal exposures, Consumer product use frequencies, Consumer product exposure de- terminants, Consumer product exposures	Consumer products are uniquely defined sources of (personal care) products.
Consumption	Consump- tions	Populations, Foods.	Food conversions, Consump- tions by modelled food.	Consumptions data are the amounts of foods consumed on specific days by individuals in a food consumption survey. For acute exposure assessments, the interest is in a population of person-days, so one day per individual may be sufficient. For chronic exposure assessments, the interest is in a population of persons, so preferably two or more days per individual are needed.
	Single value consump- tions	Consump- tions by modelled food.	Single value dietary exposures.	Single value consumption data are the single value amounts (Large Portion, Mean Consumption, p97.5Consumption) of modelled foods (foods-as-measured) consumed in a population.

Category	Module	Inputs	Used by	Description
	Market shares	Foods.	Food conversions.	Market shares data specify for a given food, percentages of more specific foods (subfoods, e.g. brands) representing their share in a market. Market shares are used when consumption data are available at a more generalised level than concentration data.
	Food recipes	Foods.	Food conversions.	Food recipes data specify the composition of specific foods (typically: foods-as-eaten) in terms of other foods (intermediate foods or modelled foods) by specifying proportions in the form of a percentage.
Occurrence	Concentra- tions	Foods, Substances, Populations, Focal food concentra- tions, Food extrapola- tions, Substance conversions, Deterministic substance conversion factors, Relative potency factors, Substance authorisa- tions, Active substances, Concentra- tion limits, Substance approvals, Processing factors.	Single value concentra- tions, Occurrence patterns, Concentra- tion models, Modelled foods.	Concentrations data are analytical measurements of chemical substances occurring in food samples. In their simplest form, concentration data can just be used as provided by datasets. Optionally, concentrations data can be manipulated for active substances, extrapolated to other foods, and/or default values can be added for water.
	Concentra- tion distributions	Foods, Substances.	Concentra- tion models, Dietary exposures.	Concentration distributions describe substance concentrations on foods in the form of summary statistics.

Table 3	3.436 -	continued	from	previous page
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Category	Module	Inputs	Used by	Description
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	Single value concentra- tions	Active substances, Concentra- tions, Concentra- tion limits, Deterministic substance conversion factors.	Modelled foods, Single value dietary exposures.	Single value concentrations data are the single value estimates (High Residue, Maximum Residue Limit, Supervised Trials Median Residue) of residue concentrations on modelled foods.
	Processing factors	Foods, Substances.	Concentra- tions, Food conversions, Dietary exposures, Single value dietary exposures.	Processing factors are multiplication factors to derive the concentration in a processed food from the concentration in an unprocessed food and can be specified for identified processing types (e.g., cooking, washing, drying). Processing factors are primarily used in dietary exposure assessments to correct for the effect of processing on substance concentrations in dietary exposure calculations.
	Unit variability factors	Foods, Substances.	Dietary exposures, Single value dietary exposures.	Unit variability factors specify the variation in concentrations between single units of the same food, which have been put together in a mixture sample on which the concentration measurements have been made. Unit variability factors are used to account for the fact that concentration data often relate to composite samples, whereas an acute risk may result from single food units.

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Category	Module	Inputs	Used by	Description
	<i>Occurrence patterns</i>	Substance authorisa- tions, Active substances, Concentra- tions.	Occurrence frequencies, Dietary exposures.	Occurrence patterns (OPs) are the combinations (or mixtures) of substances that occur together on foods and the frequencies of these mixtures occurring per food, expressed in percentages. In the context of pesticides, occurrence patterns are associated with agricultural use percentages. Occurrence patterns are relevant to account for co-occurrence of active substances in exposed individuals. Occurrence patterns may be specified as data or modelled based on observed patterns of positive concentrations.
	Occurrence frequencies	Active substances, Occurrence patterns.	Concentra- tion models, Single value dietary exposures.	Occurrence frequencies specify how often substances occur on foods. Frequencies are expressed as percentages.
	Substance authorisa- tions	Foods, Substances.	Concentra- tions, Occurrence patterns, Concentra- tion models.	Substance authorisations specify which food/substance combinations are authorised for (agricultural) use. If substance authorisations are used, then only the food/substance combinations that are specified in the data are assumed to be authorised and all other combinations are assumed to be not authorised. This information may, for instance, be used to determine whether concentration measurements below the LOQ or LOD could be assumed true zeros. I.e., if a food/substance combinations is assumed to be unauthorised, then the LOQ, LOD may be assumed to be a zero.
	Substance approvals	Substances.	Concentra- tions.	Substance approvals specify which substances are approved within the definition under regulation (EC) No 1107/2009. This information may, for instance, be used to to restrict water imputation to approved substances only.

Table 3.436 - continued from previous page

Category	Module	Inputs	Used by	Description
	Substance conversions	Substances, Active substances.	Concentra- tions.	Substance conversions specify how measured substances are converted into active substances, which are the substances assumed to cause health effects. In pesticide legislation such measured substances and the substance conversion rules are known as residue definitions.
	Deterministic substance conversion factors	Substances, Foods.	Concentra- tions, Single value con- centrations.	Data module for deterministic substance conversion factors. These are (possibly food specific) conversion factors for expressing measured substance concentrations to equivalent active substance concentrations.
	Concentra- tion limits	Foods, Substances.	Concentra- tions, Single value con- centrations, Concentra- tion models, Modelled foods.	Concentration limits specify (legal) limit values for substance concentrations on foods and are sometimes used as conservative values for concentration data. In the framework of pesticides the legal Maximum Residue Limit (MRL) is the best known example.
	Concentra- tion models	Concentra- tions, Concentra- tion limits, Active substances, Modelled foods, Substance authorisa- tions, Occurrence frequencies, Relative potency factors, Con- centration distributions, Total diet study sample composi- tions.	High exposure food- substance combina- tions, Dietary exposures.	Concentration models are distributional models of substance concentrations on foods. They describe both the substance presence (yes/no, with no representing an absolute zero concentration) and the substance concentrations. Concentration models are specified per food/substance combination.

Table	3.436 -	continued	from	previous	page

Category	Module	Inputs	Used by	Description
	Modelled foods	Concentra- tions, Single value con- centrations, Concentra- tion limits.	Concentra- tion models, Food conversions.	Modelled foods are foods within the foods scope for which concentration data or MRLs of substances are available (or expected).
	Focal food concentra- tions	Foods, Substances.	Concentra- tions.	In some cases the attention in an assessment is on a specific food (focal food), against the background of other foods. Focal food concentrations are separate concentration data for one or more focal food commodities, that will take the place of any other concentration data for the focal food in the ordinary concentration data.
	Total diet study sample compositions	Foods.	Concentra- tion models, Food conversions.	Total diet study sample compositions specify the composition of mixed food samples, such as used in a total diet study (TDS), in terms of their constituting foods.
	Food extrap- olations	Foods.	Concentra- tions, Food conversions.	Food extrapolations data specify which foods (data rich foods) can be used to impute concentration data for other foods with insufficient data (data poor foods).
	Exposure biomarker conversions	Substances.	HBM analysis.	Occasionally, the biomarker of interest (substance) is not measured. Exposure biomarker conversions specify how measured biomarkers are converted to the relevant biomarkers (substances). This type of conversion is within biological matrices.
	Consumer product con- centrations	Substances, Consumer products.	Consumer product exposures.	Consumer product concentration (distributions) data are analytical measurements of chemical substances occurring in PCPs. In their simplest form, concentration data can just be used as provided by datasets.

Table 3.436 - continued from previous page

Category	Module	Inputs	Used by	Description
Exposure	Food conversions	Consump- tions, Modelled foods, Processing factors, Food recipes, Market shares, Food extrapola- tions, Total diet study sample com- positions, Active substances.	Consump- tions by modelled food, Dietary exposures.	Food conversions relate foods-as-eaten, as found in the consumption data, to modelled foods (foods-as-measured), which are the foods for which concentration data are available. A food-as-eaten can be linked to one, or multiple modelled foods using various conversion steps (e.g., using food recipes to translate a composite food into its ingredients). There are several types of conversion steps, and a conversion path may comprise multiple conversion steps between a food-as-eaten and a modelled food.
	Consump- tions by modelled food	Consump- tions, Food conversions.	Single value consump- tions, High exposure food- substance combina- tions, Dietary exposures.	Consumptions by modelled food are consumptions of individuals expressed on the level of the foods for which concentration data are available (i.e., the modelled-foods). These are calculated from consumptions of foods-as-eaten and food conversions that link the foods-as-eaten amounts to modelled-foods amounts.
	High exposure food- substance combina- tions	Consump- tions by modelled food, Con- centration models, Active substances, Relative potency factors.	Dietary exposures.	Identification of food-as-eaten/modelled food/substance combinations that have the highest expected contribution to exposure based on a simple screening model.

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Category	Module	Inputs	Used by	Description
	Dietary exposures	Consump- tions by modelled food, Con- centration models, Processing factors, Unit variability factors, High exposure food- substance combina- tions, Active substances, Occurrence patterns, Relative potency factors, Food conversions, Concentra- tion distributions.	Internal exposures, Exposure mixtures, Dust exposures, Soil exposures, Air exposures, Risks, Envi- ronmental burden of disease.	Dietary exposures are the amounts of substances, expressed per kg bodyweight or per individual, to which individuals in a population are exposed from their diet per day. Depending on the exposure type, dietary exposures can be short-term/acute exposures and then contain exposures for individual-days, or they can be long-term/chronic exposures, in which case they represent the average exposure per day over an unspecified longer time period.
	Single value dietary exposures	Single value consump- tions, Single value con- centrations, Processing factors, Unit variability factors, Occurrence freauencies.	Single value risks.	Single value dietary exposures are based on the single value concentrations of substances, expressed per standard (kg) bodyweight and/or single value amounts of consumed modelled food. Depending on the exposure type, dietary exposures can be short-term/acute exposures.
	Single value non-dietary exposures			Single value non-dietary exposures are based on the single value concentrations or amounts of substances, as opposed to the distribution-based exposures of individuals in the non-dietary exposure module. Exposures are via any of the non-dietary routes: dermal, inhalation, or oral. Depending on the exposure type, single value non-dietary exposures can be short-term/acute or long term/chronic exposures. The exposures can be modelled as external exposures or internal exposures.

Table 3.436 – continued from previous page

Category	Module	Inputs	Used by	Description
	Non-dietary exposures	Populations, Substances, Active substances.	Internal exposures.	Non-dietary exposures are the amounts of substances to which individuals in a population are exposed via any of three non-dietary routes: dermal, inhalation or oral, per day.
	Internal exposures	Dietary exposures, Non-dietary exposures, Dust exposures, Soil exposures, Active substances, Relative potency factors, Absorption factors, Kinetic conversion factors, PBK model parametrisa- tions, Air exposures, Consumer product exposures.	Exposure mixtures, Biological matrix con- centration comparisons, Risks, Envi- ronmental burden of disease.	Internal doses and concentrations are amounts of substances, typically expressed per mass unit and per day, to which individuals in a population are exposed at the chosen target level. This target level is an internal dose or an internal concentration (expressed per unit [organ] weight). Internal exposures may be aggregated from dietary and non-dietary exposures using either absorption factors for systemic doses or kinetic conversion factors or PBK models to translate the external exposures to internal concentrations at the specific target or tissue. Internal exposures can be short-term/acute exposures and then contain exposures for individual-days, or they can be long-term/chronic exposures, in which case they represent the average exposure per day over an unspecified longer time period
	Exposure mixtures	Dietary exposures, Internal exposures, Relative potency factors, HBM analysis.		Exposure mixtures will select sets of co-occurring substances (one or more) that contribute most to the overall exposure patterns.
	HBM con- centrations	Populations, Substances.	HBM analysis.	Human (bio)monitoring data quantify substance concentrations found in humans collected in human monitoring surveys.

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Category	Module	Inputs	Used by	Description
	HBM analysis	HBM con- centrations, Active substances, Relative potency factors, Kinetic conversion factors, Exposure biomarker conversions.	Exposure mixtures, Biological matrix con- centration comparisons, Risks, Envi- ronmental burden of disease.	Human biomonitoring concentrations are substance concentration estimates for a biological matrix (e.g., urine or blood) derived from data obtained from human monitoring studies.
	Biological matrix con- centration comparisons	HBM analysis, Internal exposures.		Substances in the human body are absorbed, excreted without transformation, excreted after metabolization or stored in various tissues, bones or body fluids. The term biological matrix refers to all human specimens where concentratrions of a chemical can be measured like bodily fluids, such as blood, urine, saliva, breast milk, sweat, and other specimens, such as faeces, hair, teeth, and nails. Biological matrix concentration comparisons compares observed human monitoring data with predictions made for the same population of individuals from dietary survey data, concentration data and (optionally) non-dietary exposure data.
	Dust concen- tration distributions	Substances.	Dust exposures.	Dust concentration distributions data are analytical measurements of chemical substances occurring in dust samples. In their simplest form, concentration data can just be used as provided by datasets.
	Dust exposure determinants	Substances.	Dust exposures.	Dust exposure determinants specify the factors that determine the exposure of individuals to dust.

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Category	Module	Inputs	Used by	Description
	Dust exposures	Active substances, Dietary exposures, Dust concen- tration distributions, Dust exposure de- terminants, Individuals.	Internal exposures.	Dust exposures are the amounts of substances, expressed per kg bodyweight, to which individuals in a population are exposed from dust per day. Dust exposures are long-term/chronic exposures, they represent the average exposure per day over an unspecified longer time period.
	Soil concen- tration distributions	Substances.	Soil exposures.	Soil concentration distributions data are analytical measurements of chemical substances occurring in soil samples. In their simplest form, concentration data can just be used as provided by datasets.
	Soil exposure determinants	Substances.	Soil exposures.	Soil exposure determinants specify the factors that determine the exposure of individuals to soil.
	Soil exposures	Active substances, Dietary exposures, Soil concen- tration distributions, Soil exposure determi- nants, Individuals.	Internal exposures.	Soil exposures are the amounts of substances, expressed per kg bodyweight, to which individuals in a population are exposed from soil per day. Soil exposures are long-term/chronic exposures, they represent the average exposure per day over an unspecified longer time period.
	Indoor air concentra- tions	Substances.	Air exposures.	Indoor air concentrations data are analytical measurements of chemical substances occurring in indoor air samples. In their simplest form, concentration data can just be used as provided by datasets.
	Outdoor air concentra- tions	Substances.	Air exposures.	Outdoor air concentrations data are analytical measurements of chemical substances occurring in outdoor air samples. In their simplest form, concentration data can just be used as provided by datasets.
	Air exposure determinants	Substances.	Air exposures.	Air exposure determinant data are distributions of chemical substances occurring in air samples. The distributions can differ per age group.

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Category	Module	Inputs	Used by	Description
	<i>Air</i> <i>exposures</i>	Active substances, Dietary exposures, Indoor air concentra- tions, Outdoor air concentra- tions, Air exposure de- terminants, Individuals.	Internal exposures.	Air exposures are the amounts of substances, expressed per kg bodyweight, to which individuals in a population are exposed from air per m3. Air exposures are long-term/chronic exposures, they represent the average exposure per m3 over an unspecified longer time period.
	Individuals	Populations.	Dust exposures, Soil exposures, Air exposures.	Simulates individuals with properties like age, body weight, sex, and body surface area.
	Consumer product use frequencies	Populations, Consumer products.	Consumer product exposures.	Consumer product use frequencies specify the frequency of use for a consumer product (personal care, do-it-yourself, cleaning product) per day.
	Consumer product exposure determinants	Substances, Consumer products.	Consumer product exposures.	Consumer product exposure determinant data are distributions of chemical substances occurring in consumer products. The distributions can differ per age group.
	Consumer product exposures	Active substances, Consumer product con- centrations, Consumer product use frequencies, Consumer product exposure de- terminants.	Internal exposures.	Consumer product exposures are the amounts of substances, expressed per kg bodyweight, to which individuals in a population are exposed from average consumer product usage per individual. Consumer product exposures are long-term/chronic exposures, they represent the average exposure per day over an unspecified longer time period.
In-silico	QSAR membership models	Substances, Effects, AOP networks.	Active substances.	QSAR membership models specify assessment group memberships for active substances related to a specific health effect (adverse outcome). Memberships should be derived externally from Quantitative Structure-Activity Relationship (QSAR) models.

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Category	Module	Inputs	Used by	Description
	Molecular docking models	Substances, Effects, AOP networks.	Active substances.	Molecular docking models specify binding energies for substances in specific molecular docking models related to a specific health effect (adverse outcome).
Kinetic	Absorption factors	Substances, Active substances.	Internal exposures, Hazard characteri- sations.	Absorption factors are chemical- and route-specific values representing the fraction of external exposures passing the absorption barrier. They are used to derive internal systemic exposures from external exposures for some specified exposure route. The corresponding exposure is referred to as internal dose.
	PBK model parametrisa- tions	Substances, Active substances, PBK model definitions.	Internal exposures, Hazard characteri- sations.	The parametrisation of PBK models are specified as data. PBK models relate exposures or hazard characterisations from one or more external routes (oral, dermal, inhalation) to an internal target or biological matrix. PBK models are based on differential-equation. MCRA currently includes the EuroMix generic PBK model and the bisphenol model of ETHZ. The corresponding exposure in the biological matrix is referred to as internal concentration.
	Kinetic conversion factors	Substances, Active substances.	Internal exposures, HBM analysis, Hazard characteri- sations.	Kinetic conversion factors relate exposures or hazard characterisations from one or more external routes (dietary, non-dietary oral, dermal, inhalation) to an internal target or biological matrix. The corresponding exposure is referred to as internal concentration.
	PBK model definitions		PBK model parametrisa- tions.	PBK model definitions implement an uploadable version of PBK (SBML) models.

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Category	Module	Inputs	Used by	Description
Hazard	Active substances	AOP networks, Points of departure, Hazard characteri- sations, Molecular docking models, QSAR membership models.	Concentra- tions, Single value con- centrations, Occurrence patterns, Occurrence frequencies, Substance conversions, Non-dietary exposures, Dust exposures, Soil exposures, Air exposures, Air exposures, Consumer product exposures, Absorption factors, PBK model parametrisa- tions, Kinetic conversion factors, Relative potency factors, Relative potency factors, Hazard characteri- sations, Inter-species conversions, Inter-species factors, Con- centration models, Food conversions, High exposure food- substance combina- tions, Dietary exposures, Internal exposures, HBM analysis.	Active substances are substances that may lead (P>0) to a specific health effect (adverse outcome). Active substances are specified directly as data or calculated from POD presence, QSAR models or Molecular docking models. Active substances can have an assessment group membership 1 (crisp), or values in the range (0,1] (probabilistic).

Table 3.436 - continued from previous page

Category	Module	Inputs	Used by	Description
	Relative potency factors	Active substances, AOP networks, Hazard characteri- sations.	Concentra- tions, Concentra- tion models, High exposure food- substance combina- tions, Dietary exposures, Internal exposures, Exposure mixtures, HBM analysis, Risks.	Relative potency factors (RPFs) quantify potencies of substances with respect to a defined effect, relative to the potency of a chosen index substance. RPFs can be used to express combined exposures of multiple substances in terms of a the exposure value of the chosen index substance (i.e., in index substance equivalents). In MCRA, hazard characterisations, and therefore also RPFs are based on mass units (e.g., µg), and not on mol units. RPFs can be different for different levels of the human organism (external, internal, specific compartment). RPFs can be given as data or computed from hazard characterisations. RPFs can be specified with uncertainty. Computation from uncertain hazard characterisations allows to include correlations between uncertain RPFs which originate from using the same index substance.

Category	Module	Inputs	Used by	Description
	Hazard characteri- sations	AOP networks, Active substances, Points of departure, Dose response models, Effect repre- sentations, Inter-species conversions, Intra species factors, Absorption factors, Kinetic conversion factors, PBK model parametrisa- tions.	Active substances, Relative potency factors, Risks, Single value risks.	Hazard characterisations are reference exposure values for active substances at the chosen biological target level (external or internal). Hazard characterisations may be specified for specific effects or for the critical effect as defined in hazard characterisation. Hazard characterisations are specified as external values (e.g. human based guidance values, such as ADI or ARfD) or are based on points of departure, such as BMD(L)s from dose-response models or externally specified points of departure (NOAEL, LOAEL, MDS). The computation may involve assessment factors, e.g. for inter-species conversion, intra-species variation or additional sources of uncertainty. The calculation may also use kinetic conversion factors, PBK models or absorption factors to convert external doses to internal concentration/doses or vice versa.
	Points of departure	Substances, Effects, AOP networks.	Active substances, Hazard characteri- sations.	Externally specified points of departure can be used as an alternative to calculated BMDs from dose response models. Points of departure can be of various types, such as NOAEL, LOAEL or BMD. They can be used to construct the list of active substances, to derive relative potency factors, and to perform health impact assessments.
	Dose response models	Dose response data, Effect representa- tions.	Hazard characteri- sations.	Dose response models are models fitted to dose response data and can be provided as data or calculated using a local or remote version of PROAST. The main results for hazard and risk assessment are benchmark doses (BMDs, BMDLs), related to a specified substance, response, optionally covariate value, and the benchmark response (BMR).

Table 3.436 - continued from previous page

Category	Module	Inputs	Used by	Description
	Dose response data	Substances, Test systems, Responses.	Dose response models.	Dose response data are data on response values of test systems at specified doses of substances (or mixtures of substances) from dose response experiments.
	Effect repre- sentations	Effects, Responses, AOP networks.	Hazard characteri- sations, Dose response models.	Effect representations specify the responses that can be used to measure specified effects and which response levels, the benchmark response (BMR), define the hazard limits for the effects.
	<i>Inter-species</i> <i>conversions</i>	Substances, Effects, Active substances.	Hazard characteri- sations.	Inter-species conversions specify how to convert a hazard characterisation for a given species to a hazard characterisation for humans. In the simplest approach, this specifies a fixed inter-species factor. In a higher tier, this specifies a geometric mean (GM) and geometric standard deviation (GSD) for a lognormal uncertainty distribution of the interspecies factor. Inter-species conversion are specified per effect and can be general or substance-specific.

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Category	Module	Inputs	Used by	Description
	Intra species factors	Substances, Effects, Active substances.	Hazard characteri- sations.	Intra-species factors describe variation between individuals concerning their individual sensitivities to experience well-defined health effects. Traditionally the intraspecies factor is a fixed value, but the true distribution might be (very) uncertain. There is some support for assuming a lognormal distribution to describe the variability between individuals. In MCRA, intraspecies factors are sampled from a lognormal distribution, characterised by a geometric mean (GM) equal to 1 and a geometric standard deviation (GSD) thats needs to be given a value representing the intraspecies variability. GM is 1 by definition (50% of the population is assumed to be less sensitive than the average, 50% is mor sensitive) and has no uncertainty. On the other hand, there is uncertainty about the GSD. In MCRA it is assumed that this uncertainty is described by a Chi-square distribution with df degrees of freedom. By specifying a lower and upper bound for the p95 sensitive individual e.g. a lower value 2 and upper value 10 (meaning, the p95 individuals are between 2 and 10 times more sensible than the average human), a Chi-square distribution can be estimated where 1) the GSD specifies the variability and 2) the degrees of freedom specifies the uncertainty around the GSD.

Table 3.436 - continued from previous page

Category	Module	Inputs	Used by	Description
	<i>AOP</i> <i>networks</i>	<i>Effects.</i>	QSAR membership models, Molecular docking models, Active substances, Relative potency factors, Hazard characteri- sations, Points of departure, Effect repre- sentations.	Effects are related to each other using the toxicological concept of adverse outcome pathways (AOPs) and adverse outcome pathway networks (see https://aopwiki.org). Adverse Outcome Pathway (AOP) Networks specify how biological events (effects) can lead to an adverse outcome (AO) in a qualitative way through relations of upstream and downstream key events (KEs), starting from molecular initiating events (MIEs). Using AOPs, the adverse outcome (AO), e.g., liver steatosis, is linked to key events (KEs), e.g., triglyceride accumulation in the liver, and to molecular initiating events (MIEs), e.g., PPAR-alpha receptor antagonism. In general, multiple AOPs may lead to the same AO, and therefore AOP networks can be identified.
	Exposure response functions	Substances, Effects.	Environmen- tal burden of disease.	Exposure response functions define the relation between percentiles and percentile specific ratios for exposure effect related to environmental burden of disease.
	Burden of disease	Populations, Effects.	Environmen- tal burden of disease.	Burden of disease data specify the burden of disease information related to specific effects and burden of disease indicators in specific populations.
Risk	Risks	Dietary exposures, Internal exposures, Hazard characteri- sations, HBM analysis, Relative potency factors.	Single value risks.	Risks (health impacts) are defined as a function of exposure and hazard characterisation at a chosen biological level (external or internal). Risk metrics are either based on the ratio hazard/exposure (e.g., MOE(T)) or exposure/hazard (e.g., HI, HQ, and RPI).

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Category	Module	Inputs	Used by	Description
	Single value risks	Single value dietary exposures, Hazard characteri- sations, Risks.		Single value risks are risk estimates obtained from combining single value exposures with single value hazard characterisations or as a percentile from a risk distribution.
	Environmen- tal burden of disease	Dietary exposures, Internal exposures, HBM analysis, Exposure response functions, Burden of disease.		Environmental Burden of Disease (EBD) assessment.

Table 3.436 - continued from previous page

STANDARD ACTIONS

A standard action is a user friendly way to perform a complex probabilistic calculation. By using a standard action predefined settings are used and the user can set only a limited number of selections. All settings (pre-defined and set by the user) are visible in the output. As a result a short output is presented. More detailed output is still available.

4.1 Chronic mixture risk assessment of metals

This standard action is of type: Risks

In the context of the European projects HBM4EU and PARC, the National Institute for Public Health and the Environment of the Netherlands (RIVM) performed a case study on the risk assessment of the combined exposure of four metals relevant for chronic kidney disease, i.e. cadmium, lead, in-organic arsenic and inorganic mercury. The exposure assessment used chemical concentration in foods of 14 European countries over the years 2014-2018, obtained from the EFSA data warehouse and individual Dutch food consumption data. Since chronic kidney disease is relevant for the adult population, adults in the age of 18-64 years were selected in the standard action. Exposure estimates were obtained using the observed individual means (OIM) model implemented in MCRA. Using relative potency factors, the exposure to the metals was expressed as equivalents of cadmium and summed per individual in the food consumption database, yielding a distribution of summed cadmium-equivalents. The summed cadmium-equivalents per individual were divided by the reference point of cadmium, which resulted in a distribution of risk characterisation ratios (exposure/hazard) E/H (or better modified reference point indexes) from which the mean, P50 and P95 are obtained. A ratio E/H or modified reference point index > 1 means either a risk cannot be excluded or refinement of the assessment is needed, depending on the assesses uncertainties. The standard action is meant for training and demonstration purposes, to demonstrate assessment of the impact of setting (new) maximum limits concentrations (MLs).

Table Group	Name	Repository	Туре
AssessmentGroup- Memberships	Catalogues_DNT_NEF- metals.xlsx	Standard Actions/Chronic mixture risk assess- ment of metals	Fixed
Compounds	Catalogues_DNT_NEF- metals.xlsx	Standard Actions/Chronic mixture risk assess- ment of metals	Fixed
Effects	Catalogues_DNT_NEF- metals.xlsx	Standard Actions/Chronic mixture risk assessment of metals	Fixed
FoodTranslations	Catalogues_DNT_NEF- metals.xlsx	Standard Actions/Chronic mixture risk assess- ment of metals	Fixed
Foods	Catalogues_DNT_NEF- metals.xlsx	Standard Actions/Chronic mixture risk assess- ment of metals	Fixed
HazardCharacterisa- tions	Catalogues_DNT_NEF- metals.xlsx	Standard Actions/Chronic mixture risk assess- ment of metals	Fixed
Populations	Catalogues_DNT_NEF- metals.xlsx	Standard Actions/Chronic mixture risk assess- ment of metals	Fixed
RelativePotencyFactors	Catalogues_DNT_NEF- metals.xlsx	Standard Actions/Chronic mixture risk assess- ment of metals	Fixed
Concentrations	DNT-NEF-SSD-metals.xlsx	Standard Actions/Chronic mixture risk assess- ment of metals	Fixed
MaximumResidueLim- its	NEF-MLs-metals.xlsx	Standard Actions/Chronic mixture risk assess- ment of metals	Fixed
Survey	NL-FE1-FCS2016_Core_1- 79y.accdb	Standard Actions/Chronic mixture risk assess- ment of metals	Fixed

Table 4.1: Datasources for Chronic mixture risk assessment of metals.

References:

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- Sprong et al. Combined chronic dietary exposure to cadmium, lead, inorganic arsenic and inorganic mercury may pose a risk for nephrotoxicity in the adult population of ten European countries. manuscript in preparation.

4.2 Chronic cumulative exposure assessment PA

This standard action is of type: Risks

In 2017 EFSA published a statement *Risks for human health related to the presence of pyrrolizidine alkaloids in honey, tea, herbal infusions and food supplements*, see EFSA (2017b). Occurrence data used in this opinion was published in 2015 in the external scientific report *Occurrence of Pyrrolizidine Alkaloids in food*, see Mulder et al. (2015). The occurrence data in tea and herbal infusions is linked to the Consumption of 6 different population groups of the DNFCS 2012-2016. An lower bound (LB) and upper bound (UB) chronic cumulative exposure assessment can be calculated for different PAs. One scenario is assuming equipotency, another scenario is using provisional RPFs (Merz and Schrenk (2016)).

4.3 Chronic cumulative exposure assessment PFAS

This standard action is of type: Risks

This standard action can be used to calculate a chronic cumulative exposure for four PFAS. Assuming equipotency, and using proposed RPFs.

Table Group	Name	Repository	Туре
Survey	ConsumptionData-1-2yr.mdb	Standard Actions/Chronic cumulative expo- sure assessment PFAS	Fixed
Survey	ConsumptionData-10-17yr.mdb	Standard Actions/Chronic cumulative expo- sure assessment PFAS	Fixed
Survey	ConsumptionData-18-64yr.mdb	Standard Actions/Chronic cumulative expo- sure assessment PFAS	Fixed
Survey	ConsumptionData-3-9yr.mdb	Standard Actions/Chronic cumulative expo- sure assessment PFAS	Fixed
Survey	ConsumptionData-65-74yr.mdb	Standard Actions/Chronic cumulative expo- sure assessment PFAS	Fixed
Survey	ConsumptionData-75plusyr.mdb	Standard Actions/Chronic cumulative expo- sure assessment PFAS	Fixed
Concentrations	PFAS-Occurrencedata-EFSA- LB.mdb	Standard Actions/Chronic cumulative expo- sure assessment PFAS	Fixed
AssessmentGroup- Memberships	PFAS-OtherData.mdb	Standard Actions/Chronic cumulative expo- sure assessment PFAS	Fixed
Compounds	PFAS-OtherData.mdb	Standard Actions/Chronic cumulative expo- sure assessment PFAS	Fixed
Effects	PFAS-OtherData.mdb	Standard Actions/Chronic cumulative expo- sure assessment PFAS	Fixed
Foods	PFAS-OtherData.mdb	Standard Actions/Chronic cumulative expo- sure assessment PFAS	Fixed
HazardCharacterisa- tions	PFAS-OtherData.mdb	Standard Actions/Chronic cumulative expo- sure assessment PFAS	Fixed
RelativePotencyFac- tors	PFAS- RelativePotencyFactorsEquipotenc	Standard Actions/Chronic cumulative expo- sure assessment PFAS	Fixed

Table 4 2. Datasour	ces for Chronic o	cumulative exposu	re assessment PFAS
	cos for childine	cumulative exposu	

4.4 Demo acute cumulative risk assessment

This standard action is of type: Risks

In this demo with fictitious data, acute cumulative risk assessments can be performed following various calculation

methods (EFSA 2012 Optimistic and Pessimistic, EC 2018 Tier 1 and Tier 2). Here, also the effect of applying processing factors can be assessed.

Table Group	Name	Repository	Туре
MaximumResidue- Limits	DemoConcentrationLim- its.mdb	Standard Actions/Demo Acute Cumulative Risk Assessment	Fixed
Concentrations	DemoConcentrations.mdb	Standard Actions/Demo Acute Cumulative Risk Assessment	Fixed
Survey	DemoConsumptions.mdb	Standard Actions/Demo Acute Cumulative Risk Assessment	Fixed
Effects	DemoEffects.mdb	Standard Actions/Demo Acute Cumulative Risk Assessment	Fixed
FoodExtrapolations	DemoFoodExtrapola- tions.mdb	Standard Actions/Demo Acute Cumulative Risk Assessment	Fixed
FoodTranslations	DemoFoodRecipes.mdb	Standard Actions/Demo Acute Cumulative Risk Assessment	Fixed
Foods	DemoFoods.mdb	Standard Actions/Demo Acute Cumulative Risk Assessment	Fixed
HazardDoses	DemoPointsOfDepar- ture.mdb	Standard Actions/Demo Acute Cumulative Risk Assessment	Fixed
Processing	DemoProcessingfac- torOri.mdb	Standard Actions/Demo Acute Cumulative Risk Assessment	Fixed
AuthorisedUses	DemoSubstanceAuthorisa- tions.mdb	Standard Actions/Demo Acute Cumulative Risk Assessment	Fixed
ResidueDefinitions	DemoSubstanceConver- sions.mdb	Standard Actions/Demo Acute Cumulative Risk Assessment	Fixed
Compounds	DemoSubstances.mdb	Standard Actions/Demo Acute Cumulative Risk Assessment	Fixed
UnitVariability	DemoUnitVar36.mdb	Standard Actions/Demo Acute Cumulative Risk Assessment	Fixed
UnitVariability	DemoUnitVarPRIMo.mdb	Standard Actions/Demo Acute Cumulative Risk Assessment	Fixed

Table 4.3: Datasources for Demo acute cumulative risk assessment.

4.5 Acute single substance dietary exposure assessment of carbofuran or chlorpyrifos

This standard action is of type: Dietary exposures

This standard action allows you to perform probabilistic acute dietary exposure assessments of single substances in food. This standard actions contains examples for the two pesticides carbofuran and chlorpyrifos. The assessment is based on consumption data for two-years-old from the Dutch National Food Consumption Survey (DNFCS)-Young children (Ocké et al. (2008)). The consumption data are coded according to EFSAs coding system raw primary commodity (RPC) (EFSA (2019)). The 2014 concentration data of carbofuran and chlorpyrifos were extracted from EFSAs Data Warehouse and are organised according to the SSD data format. The standard action is developed for training purposes and results should not be regarded as representative outcomes of full dietary exposure assessments of carbofuran or chlorpyrifos.

In this standard action you can:

- Select the substance; carbofuran or chlorpyrifos,
- Choose the method for handling left-censored concentration measurements (lower bound or upper bound approach),
- Choose whether to include correction for processing,
- Choose whether to account for unit-to-unit variation using the Beta model,

• Run with or without uncertainty.

Table 4.4: Datasources for Acute single substance dietary exposure assess
ment of carbofuran or chlorpyrifos.

Table Group	Name	Repository	Туре
Com- pounds	Catalogues_WHO_Training_Acu NL2.xlsx	Standard Actions/Acute single substance dietary exposure assessment of carbofuran or chlorpyrifos	Fixed
Food- Transla- tions	Catalogues_WHO_Training_Acu NL2.xlsx	Standard Actions/Acute single substance dietary exposure assessment of carbofuran or chlorpyrifos	Fixed
Foods	Catalogues_WHO_Training_Act NL2.xlsx	Standard Actions/Acute single substance dietary exposure assessment of carbofuran or chlorpyrifos	Fixed
Popula- tions	Catalogues_WHO_Training_Act NL2.xlsx	Standard Actions/Acute single substance dietary exposure assessment of carbofuran or chlorpyrifos	Fixed
Process- ing	Catalogues_WHO_Training_Act NL2.xlsx	Standard Actions/Acute single substance dietary exposure assessment of carbofuran or chlorpyrifos	Fixed
UnitVari- ability	Catalogues_WHO_Training_Act NL2.xlsx	Standard Actions/Acute single substance dietary exposure assessment of carbofuran or chlorpyrifos	Fixed
Concen- trations	Concentra- tions_WHO_Carbofuran_Chlorp	Standard Actions/Acute single substance dietary exposure assessment of carbofuran or chlorpyrifos	Fixed
Survey	Consumptions_NL_2yr_VCP-2002-2006_RPC.xlsx	Standard Actions/Acute single substance dietary exposure assessment of carbofuran or chlorpyrifos	Fixed

4.6 Chronic single substance dietary exposure assessment of lead or atropine

This standard action is of type: Dietary exposures

This standard action enables you to perform probabilistic chronic dietary exposure assessments of single substances in food (Boon et al. (2017), Boon et al. (2022)). The assessment is based on consumption data of women aged 15-44 years from the FAO-WHO Gift survey of Bangladesh, coded according to EFSA's FoodEx2 classification. The atropine concentration data were extracted from the GEMS database. The lead concentration data for foods are artificial, manually generated around the means reported by EFSA in 2012 (EFSA 2012). The concentration data are organised according the SSD data format. The standard action is developed for training purposes and results should not be seen as a full dietary exposure assessment of atropine or lead

This example allows you to:

- Select the substance atropine or lead,
- Choose the method for handling left-censored concentration measurements (lower bound or upper bound approach),
- Select a method for modelling long term intakes (OIM/LNN),
- Run with or without uncertainty

Table Group	Name	Repository	Туре
Com- pounds	Catalogues_WHO_Training_ch BD.xlsx	Standard Actions/Chronic single substance dietary expo- sure assessment lead or atropine	Fixed
Food- Transla- tions	Catalogues_WHO_Training_ch BD.xlsx	Standard Actions/Chronic single substance dietary expo- sure assessment lead or atropine	Fixed
Foods	Catalogues_WHO_Training_ch BD.xlsx	Standard Actions/Chronic single substance dietary expo- sure assessment lead or atropine	Fixed
Popula- tions	Catalogues_WHO_Training_ch BD.xlsx	Standard Actions/Chronic single substance dietary expo- sure assessment lead or atropine	Fixed
Concentra- tions	Concentra- tions_WHO_Training_Pb_At.x	Standard Actions/Chronic single substance dietary expo- sure assessment lead or atropine	Fixed
Survey	Consump- tions_BD_GIFT_2022.xlsx	Standard Actions/Chronic single substance dietary expo- sure assessment lead or atropine	Fixed

Table 4.5: Datasources for Chronic single substance dietary exposure a	as-
sessment of lead or atropine.	

4.7 Demo Human Monitoring Analysis bisphenols

This standard action is of type: Biological matrix concentration comparisons

Human BioMonitoring (HBM) is a primary instrument to measure real-life exposure to chemicals. Because of the associated high costs, chemical levels in body fluids such as blood or urine would ideally be predictable from estimated external exposure levels, such as in the diet and/or from other non-dietary sources. It is needed to convert external exposures to internal concentrations by use of a kinetic (PBK) model or the application of a simple kinetic conversion factor.

This standard action provides a simple demonstration of this approach. It is based on the EuroMix biomonitoring study (Husøy et al. (2019), Karrer et al. (2019)) and considers three of the investigated chemical substances in this study, i.e. the bisphenols BPA, BPS and BPF, which could have adverse estrogenic effects and therefore require risk assessment. These substances were measured in the urine of 144 adult individuals on two days (in this demo, we only use the data of a single day). The study subjects also kept detailed diaries on their food consumption and use of personal care products.

In this demonstrator standard action, the dietary exposure and optionally the non-dietary exposure from personal care products and thermal paper is modelled. HBM measurement values below the limit of detection are imputed. The specific gravity of the urine samples was measured as well, which is used in the calculation to calculate adjusted urine concentrations as decribed in Husøy et al. (Husøy et al. (2019)). The dietary exposure is predicted from the consumption data for 226 modelled foods derived from the food diaries and food monitoring concentration data for BPA, BPS and BPF in these foods. Additionally, non-dietary exposure from personal care products and from handling thermal paper was modelled separately by Karrer et al. (Karrer et al. (2020)) and these exposures are available as data for three non-dietary routes, i.e. dermal, oral and inhalation. These non-dietary exposures can be aggregated with the dietary exposures at the individual level. In this demonstrator the conversion from external exposures to urine concentration can be done using a simple kinetic conversion factor approach or using a PBK model developed by Karrer et al., which is also available in MCRA (Karrer et al. (2018), Karrer et al. (2019), Karrer et al. (2020)).

In this demonstrator standard action several choices can be made:

- Exposure type: Select acute or chronic exposure. For acute, HBM data are taken per day and modelled exposures are obtained from 1000 simulated dietary exposures obtained from Monte Carlo integration of individual day consumption patterns and food concentrations and possibly combined/aggregated with matching non-dietary exposures of the simulated individual days. For chronic, HBM data and non-dietary exposure data are averaged per individual and dietary exposures are modelled using the observed individual means model.
- Exposure route: Derive modelled concentrations from dietary exposure only or aggregate with non-dietary exposure sources.

- **Censored valude handling method:** For the modelled concentrations, assume censored values in the food concentration data (measurements below the limit of reporting LOR) to be zero or impute them with a value 0.5 x LOR.
- **Kinetic conversion model:** Choose from two example data files with substance-specific absorption factors or use the PBK model developed by Karrer et al.

Table Group	Name	Repository	Туре
Compounds	Demo_HBM_CataloguesAndSecondaryData	Standard Actions/Demo Human Mon- itoring Analysis	Fixed
Effects	Demo_HBM_CataloguesAndSecondaryData	Standard Actions/Demo Human Mon- itoring Analysis	Fixed
FoodTranslations	Demo_HBM_CataloguesAndSecondaryData	Standard Actions/Demo Human Mon- itoring Analysis	Fixed
Foods	Demo_HBM_CataloguesAndSecondaryData	Standard Actions/Demo Human Mon- itoring Analysis	Fixed
RelativePotency- Factors	Demo_HBM_CataloguesAndSecondaryData	Standard Actions/Demo Human Mon- itoring Analysis	Fixed
Concentrations	Demo_HBM_ConcentrationsSSD.xlsx	Standard Actions/Demo Human Mon- itoring Analysis	Fixed
Survey	Demo_HBM_Consumptions.xlsx	Standard Actions/Demo Human Mon- itoring Analysis	Fixed
KineticConver- sionFactors	Demo_HBM_KineticConversionFactorsBisp	Standard Actions/Demo Human Mon- itoring Analysis	Fixed
PbkModels	Demo_HBM_KineticModelsBisphenols_KM	Standard Actions/Demo Human Mon- itoring Analysis	Fixed
NonDietary	Demo_HBM_NonDietaryExposures.xlsx	Standard Actions/Demo Human Mon- itoring Analysis	Fixed
HumanMonitor- ingData	Demo_HBM_PhenolsUrinePooledOneDay.x	Standard Actions/Demo Human Mon- itoring Analysis	Fixed

Table 4.6.	Datasources	for D	emo I	Human	Monit	toring	Analys	ie hier	henole
Table 4.0.	Datasources	101 D	enio	numan	WIOIII	loring	Analys.	is dis	menois.

4.8 TDS-based long term dietary exposure and risk assessment

This standard action is of type: Risks

This standard action provides an overview about probabilistic long-term dietary exposure assessments with TDS data in MCRA. Possibilities to calculate exposure for population (sub-)groups from different countries and for different substances are provided. In addition, options for regional, seasonal or production type specific exposure are demonstrated along with options to select for consumers of different foods. In order to highlight options of the MCRA TDS module, the standard actions provide a simple and condensed overview about possible settings in MCRA. Consult the documentation pages on *TDS-based exposure and risk assessment* if you want to learn more about running these types of assessments in MCRA.

4.9 Long-term dietary exposure and risk of nickel for the Belgian population

This standard action is of type: Risks

The risk assessment originally conducted by the European Food Safety Authority stated some concerns regarding the chronic exposure of the European population to nickel due to food intake. This study aimed to evaluate the extent of the Belgian population's exposure to nickel via intake of different foods/drinks available in their market.

This standard actions is using nickel concentration data and Belgian consumption data-2014 data, for training and demonstration purposes and is meant to demonstrate the average nickel exposure for total Belgian consumers, for differences Belgian sub-populations and specifying the exposure for different foods (LB and UB).

This standard action demonstrates the aggregated exposure assessment of the effects of nickel occurrence in food of Belgian population. It uses 2017 Nickel contamination data in foods available in Belgian market and Belgian consumption data 2014.

	for the Beigian population.		
Table Group	Name	Repository	Туре
Survey	2021-09- 17_BelgianConsumptions.xlsx	Standard Actions/Long-term dietary exposure and risk of nickel for Belgian age groups	Fixed
Compounds	Catalogues_FNS_Case_Study_BE_Ni_M	Standard Actions/Long-term dietary exposure and risk of nickel for Belgian age groups	Fixed
Effects	Catalogues_FNS_Case_Study_BE_Ni_M	Standard Actions/Long-term dietary exposure and risk of nickel for Belgian age groups	Fixed
Foods	Catalogues_FNS_Case_Study_BE_Ni_M	Standard Actions/Long-term dietary exposure and risk of nickel for Belgian age groups	Fixed
HazardChar- acterisations	Catalogues_FNS_Case_Study_BE_Ni_M	Standard Actions/Long-term dietary exposure and risk of nickel for Belgian age groups	Fixed
Populations	Catalogues_FNS_Case_Study_BE_Ni_M	Standard Actions/Long-term dietary exposure and risk of nickel for Belgian age groups	Fixed
Concentra- tions	Contaminants_BE_2017- 2019_INNIBEL_Ni_MCRA_FE2.xlsx	Standard Actions/Long-term dietary exposure and risk of nickel for Belgian age groups	Fixed
FoodTrans- lations	FoodTranslations_INNIBEL_Ni.xlsx	Standard Actions/Long-term dietary exposure and risk of nickel for Belgian age groups	Fixed

Table 4.7: Datasources for Long-term dietary exposure and risk of nickel for the Belgian population.

4.10 TDS-based long-term exposure and risk assessment of methylmercury for German children

This standard action is of type: Risks

Note

This standard action was developed within the FNS-Cloud project (https://www.fns-cloud.eu). Related consumption data and example conversion scripts for uploading data will be made available in the FNS Cloud (status 05/2022: in progress).

4.10.1 Introduction

This standard action performs *a TDS-based probabilistic risk assessment* regarding the exposure of German children to methylmercury. Total Diet Study (TDS) data are combined with food consumption data, and stratification of the calculations regarding region, season and production type (organic or conventional) can be considered. Total Diet Studies (TDS) are an appropriate method to representatively reflect background occurrence levels of substances in food in a population's diet. More than 90 % of the diet are covered, the foods are prepared as consumed and pooled to composite samples prior to chemical analysis. Next to the representation of almost the total diet in Germany, the German TDS (BfR MEAL Study) also allows to elaborate potential differences in occurrence of substances in different stratifications. The stratifications consider four different regions (North, East, South and West Germany), two different seasons (which are in this demo fictitious summer and winter season), and the production type (i.e. organic versus conventional production).

In this example, consumption data for children in Germany and fictive TDS occurrence data for methylmercury are used. The commonly used lower bound (LB) or upper bound (UB) scenarios are used for treatment of censored data. Exposure is calculated applying the Observed Individual Means model. The exposure is expressed as μg per kg body weight and day for the mean, and the 50th, 90th, 95th, 97.5th, 99th and 99.9th percentiles of the exposure

distribution and the relative contributions from modelled foods or main food groups to total exposure is displayed. For risk assessment, the exposure distribution is compared to the tolerable weekly intake (TWI) of 1.3 μ g per kg body weight and day set for methylmercury set by EFSA. The risk in indicated by the risk characterisation ratio exposure/hazard E/H quantified by exposure relative to the TWI.

4.10.2 Standard action options

In this standard action, potential differences between stratification related exposure assessments are demonstrated. The user is able to select and combine the above-mentioned stratifications and compare exposure outcomes and risks.

- By selection of a certain region the occurrence data from this region will be combined with the respective consumptions data from children living in this region.
- By selection of a certain season the occurrence data from this season will be combined with consumption events during this season. Summer season is defined from April to September and Winter season from October till March.
- By selection of a type of production occurrence data from organically or conventionally produced foods will be combined with the overall consumption, since information on consumption of organically or conventionally produced foods is not included in the consumption data.
- Exposure from foods that are not stratified according to the above-mentioned characteristics is included on an unspecific basis into the assessments (e.g., the item "pasta" is not sampled in different regions, therefore regional exposure will be calculated by combining regional consumption with the concentration of the unspecified sample).
- The assessments can be done either for all individuals included in the consumption survey or can be restricted to consumers only. The restriction will consider consumers of one or more selected TDS (modelled) foods but includes all consumptions of these individuals.
- The option to calculate uncertainties analysis results in 95 % confidence intervals (CIs) by resampling the individuals.

4.10.3 Standard action data

Table Group	Name	Repository	Туре
Compounds	Cata- logues ENS DE MeH(Standard Actions/TDS-based long-term exposure and risk as-	Fixed
Effects	Cata- logues FNS DE MeHs	Standard Actions/TDS-based long-term exposure and risk as- sessment of methylmercury for German children	Fixed
Foods	Cata- logues FNS DE MeH	Standard Actions/TDS-based long-term exposure and risk as- sessment of methylmercury for German children	Fixed
HazardChar- acterisations	Cata- logues_FNS_DE_MeH	Standard Actions/TDS-based long-term exposure and risk as- sessment of methylmercury for German children	Fixed
TotalDiet- Study	Cata- logues_FNS_DE_MeHs	Standard Actions/TDS-based long-term exposure and risk as- sessment of methylmercury for German children	Fixed
Concentra- tions	Concentra- tions_FNS_DE_MeHg_	Standard Actions/TDS-based long-term exposure and risk as- sessment of methylmercury for German children	Fixed
Survey	Consumptions_DE_VEl 2002	Standard Actions/TDS-based long-term exposure and risk as- sessment of methylmercury for German children	Fixed

Table 4.8: Datasources for TDS-based long-term exposure and risk assessment of methylmercury for German children.

4.11 Acute Cumulative Risk Assessment Craniofacial Alterations (EFSA 2022)

This standard action is of type: Single value risks

This standard action will enable you to run acute cumulative dietary risk assessment of craniofacial alterations by pesticide residues using the same methods and data as used by EFSA in 2022.

Table Group	Name	Repository	Туре
Compounds	Catalogues.xlsx	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Other data	Fixed
Effects	Catalogues.xlsx	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Other data	Fixed
Foods	Catalogues.xlsx	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Other data	Fixed
Concentrations	Concentra- tionsSSD_DAC.md	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Concentrations	Fixed
Survey	Consump- tionsBE.mdb	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Consumptions	Fixed
Survey	Consumption- sCZ.mdb	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Consumptions	Fixed
Survey	Consump- tionsDE.mdb	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Consumptions	Fixed
Survey	Consump- tionsDK.mdb	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Consumptions	Fixed
Survey	Consump- tionsES.mdb	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Consumptions	Fixed
Survey	Consump- tionsFI.mdb	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Consumptions	Fixed
Survey	Consump- tionsFR.mdb	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Consumptions	Fixed
Survey	Consumption- sHU.mdb	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Consumptions	Fixed
Survey	Consumption- sIE.mdb	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Consumptions	Fixed
Survey	Consumption- sIT.mdb	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Consumptions	Fixed
Survey	Consumption- sLT.mdb	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Consumptions	Fixed
Survey	Consumption- sNL.mdb	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Consumptions	Fixed
Survey	Consumption- sRO.mdb	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Consumptions	Fixed
Survey	Consumption- sSE.mdb	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Consumptions	Fixed
Assessment- GroupMember- ships	Secondary- Data.xlsx	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Other data	Fixed
AuthorisedUses	Secondary-	Standard Actions/Acute Cumulative Risk Assessment Cranio-	Fixed
FoodTranslations	Data.xlsx Secondary-	facial Alterations (EFSA 2022)/Other data Standard Actions/Acute Cumulative Risk Assessment Cranio-	Fixed
HazardDoses	Secondary-	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EESA 2022)/Other data	Fixed
MaximumResidu-	Secondary-	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EESA 2022)/Other data	Fixed
Processing	Secondary-	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EESA 2022)/Other data	Fixed
ResidueDefini-	Secondary-	Standard Actions/Acute Cumulative Risk Assessment Cranio-	Fixed
tions Substance Ap-	Data.xlsx Secondary-	facial Alterations (EFSA 2022)/Other data Standard Actions/Acute Cumulative Risk Assessment Cranic	Fixed
provals	Data.xlsx	facial Alterations (EFSA 2022)/Other data	TIXEU
UnitVariability	UnitVar36.xlsx	Standard Actions/Acute Cumulative Risk Assessment Cranio- facial Alterations (EFSA 2022)/Other data	Fixed

Table 4.9: Datasources for Acute Cumulative Risk Assessment Craniofa
cial Alterations (EFSA 2022).

4.12 EU acute cumulative exposure assessment (2018) Tier I and Tier II

This standard action is of type: Single value risks

This standard action is based on work done in 2018 (van Klaveren et al. (2019a)). In the context of the second framework partnership agreement between the National Institute for Public Health and the Environment of the Netherlands (RIVM) and the European Food Safety Authority (EFSA) acute cumulative dietary exposure assessments were performed for two cumulative assessment groups (CAGs) of pesticides that affect the nervous system: pesticides causing brain and/or erythrocyte AChE inhibition (CAG-NAN, 47 pesticides) and pesticides causing functional alterations of the motor division (CAG-NAM, 100 pesticides). The exposure assessments used monitoring data collected by the Netherlands under their official monitoring programmes in 2014, 2015 and 2016 and individual Dutch food consumption data. Exposure estimates were obtained for each group of pesticides using the MCRA software. The Standing Committee on Plants, Animals, Food and Feed (SC PAFF) discussed the scope of the assessment in 2018 and agreed on the parameters to be used for the cumulative exposure assessment. Based on that discussion, a very conservative tier I modelling approach and a refined, but still conservative tier II modelling approach were used. In these assessments, common risk assessment practice was followed and the cumulative exposure was expressed as the total riskcharacterisation ratio hazard/exposure H/E at the 50th, 90th, 95th, 99th and 99.9th percentile of the exposure distribution.

Table Group	Name	Repository	Туре
AssessmentGroup- Memberships	LeanCAGsAssessmentGroupMem- bership10%p999.xlsx	Standard Actions/EU 2018 Acute Cumula- tive Exposure Assessment	Fixed
AuthorisedUses	SecondaryInputData.mdb	Standard Actions/EU 2018 Acute Cumula- tive Exposure Assessment	Fixed
Compounds	SecondaryInputData.mdb	Standard Actions/EU 2018 Acute Cumula- tive Exposure Assessment	Fixed
Effects	SecondaryInputData.mdb	Standard Actions/EU 2018 Acute Cumula- tive Exposure Assessment	Fixed
FoodExtrapolations	SecondaryInputData.mdb	Standard Actions/EU 2018 Acute Cumula- tive Exposure Assessment	Fixed
FoodTranslations	SecondaryInputData.mdb	Standard Actions/EU 2018 Acute Cumula- tive Exposure Assessment	Fixed
Foods	SecondaryInputData.mdb	Standard Actions/EU 2018 Acute Cumula- tive Exposure Assessment	Fixed
HazardDoses	SecondaryInputData.mdb	Standard Actions/EU 2018 Acute Cumula- tive Exposure Assessment	Fixed
MaximumResidue- Limits	SecondaryInputData.mdb	Standard Actions/EU 2018 Acute Cumula- tive Exposure Assessment	Fixed
Processing	SecondaryInputData.mdb	Standard Actions/EU 2018 Acute Cumula- tive Exposure Assessment	Fixed
ResidueDefinitions	SecondaryInputData.mdb	Standard Actions/EU 2018 Acute Cumula- tive Exposure Assessment	Fixed
UnitVariability	UnitVar36.mdb	Standard Actions/EU 2018 Acute Cumula- tive Exposure Assessment	Fixed
Survey	a_ConsumptionsNL2.mdb	Standard Actions/EU 2018 Acute Cumula- tive Exposure Assessment	Fixed
Survey	a_ConsumptionsNL3_6.mdb	Standard Actions/EU 2018 Acute Cumula- tive Exposure Assessment	Fixed

Table 4.10: Datasources for EU acute cumulative exposure assessment (2018) Tier I and Tier II.

4.13 EU chronic cumulative exposure assessment (2018) Tier I and Tier II

This standard action is of type: Risks

This standard action is based on research done in 2018 van Klaveren et al. (2019b).

This standard action will enable you to reproduce the exposure assessment of chronic cumulative effects of pesticide residues in food affecting the thyroid. These are retrospective exposure assessments of the cumulative exposure for the thyroid using monitoring data from 2014, 2015 and 2016. In this standard action Dutch monitoring and consumption data are used. The results, data used and methodology are reported in a scientific report following published on the EFSA website in September 2019. The methodology fulfils the requirements set by the European Commission.

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Table Group	Name	Repository	Туре
AssessmentGroup-	LeanCAGsAssessmentGroup-	Standard Actions/EU 2018 Chronic Cumu-	Fixed
Memberships	Membership10%p999.xlsx	lative Exposure Assessment	
AuthorisedUses	SecondaryInputData.mdb	Standard Actions/EU 2018 Chronic Cumu- lative Exposure Assessment	Fixed
Compounds	SecondaryInputData.mdb	Standard Actions/EU 2018 Chronic Cumu- lative Exposure Assessment	Fixed
Effects	SecondaryInputData.mdb	Standard Actions/EU 2018 Chronic Cumu- lative Exposure Assessment	Fixed
FoodExtrapolations	SecondaryInputData.mdb	Standard Actions/EU 2018 Chronic Cumu- lative Exposure Assessment	Fixed
FoodTranslations	SecondaryInputData.mdb	Standard Actions/EU 2018 Chronic Cumu- lative Exposure Assessment	Fixed
Foods	SecondaryInputData.mdb	Standard Actions/EU 2018 Chronic Cumu- lative Exposure Assessment	Fixed
HazardDoses	SecondaryInputData.mdb	Standard Actions/EU 2018 Chronic Cumu- lative Exposure Assessment	Fixed
MaximumResidue- Limits	SecondaryInputData.mdb	Standard Actions/EU 2018 Chronic Cumu- lative Exposure Assessment	Fixed
Processing	SecondaryInputData.mdb	Standard Actions/EU 2018 Chronic Cumu- lative Exposure Assessment	Fixed
ResidueDefinitions	SecondaryInputData.mdb	Standard Actions/EU 2018 Chronic Cumu- lative Exposure Assessment	Fixed
Survey	c_ConsumptionsNL2.mdb	Standard Actions/EU 2018 Chronic Cumu- lative Exposure Assessment	Fixed
Survey	c_ConsumptionsNL3_6.mdb	Standard Actions/EU 2018 Chronic Cumu- lative Exposure Assessment	Fixed

Table 4.11: Datasources for EU chronic cumulative exposure assessment (2018) Tier I and Tier II.

4.14 EU Prospective Dietary Cumulative Risk Assessment (2023)

This standard action is of type: *Single value risks*

In the context of the third framework partnership agreement between the National Institute for Public Health and the Environment of the Netherlands (RIVM) and the European Food Safety Authority (EFSA) Standard Regulatory Actions are provided to allow users to perform prospective acute and chroniccumulative dietary exposure assessments.

A user can choose for different effects. For two cumulative assessment groups (CAGs) of pesticides that affect the nervous system: pesticides causing brain and/or erythrocyte AChE inhibition (CAG-NAN, 47 pesticides) and pesticides causing functional alterations of the motor division (CAG-NAM, 100 pesticides).

For two cumulative assessment groups (CAGs) of pesticides that affect craniofacial alterations: pesticides causing

craniofacial alterations due abnormal skeletal development (CAG-DAC, 41 pesticides) and pesticides causing head abnormalities not due to abnormal skeletal (CAG-DAH, 41 pesticides).

For chronic an assessment group of pesticides that affect the nervous system: pesticides causing brain and/or erythrocyte AChE inhibition (CAG-NCN, 47 pesticides. And the group a group of pesticides having an effect on the parafollicular (C-) cells or the calcitonin system (CAG-TCP, 18 substances) and pesticide residues affecting follicular cells and/or thyroid hormone (T3/T4) system (CAG TCF, 124 substances).

Occurrence datasets for the background are different 3-year cycle datasets containing occurrence data of all member states in the 3-year cycle, or it can be omitted (for comparison) so that only the background exposure is calculated. MRL exceedances can be removed taking into account measurement uncertainty. As an advanced option it is possible to adjust the 99.9th percentile of the exposure using uncertain adjustment factors from the EFSA cumulative risk assessment report 2020. The exposure is calculated probabilistically and the risk is expressed as margin of exposure (MOE) and is compared to a threshold value of 100. The user can upload own focal commodity/focal substance data (+ processing factors/conversion factors).

Table Group	Name	Repository
AssessmentGroupMemberships	EFSA-CRA-2023-CAGs level 2.zip	EFSA-CRA/CRA 2023
Effects	EFSA-CRA-2023-CAGs level 2.zip	EFSA-CRA/CRA 2023
HazardCharacterisations	EFSA-CRA-2023-CAGs level 2.zip	EFSA-CRA/CRA 2023
HazardDoses	EFSA-CRA-2023-CAGs level 2.zip	EFSA-CRA/CRA 2023
Concentrations	EFSA-CRA-2023-Concentrations (2019-2021).zip	EFSA-CRA/CRA 2023/Concentration
Survey	EFSA-CRA-2023-Consumptions AT-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions BE-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions BE-Other children.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions BG-Other children.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions BG-Toddlers.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions CZ-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions CZ-Other children.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions DE-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions DE-Other children.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions DE-Toddlers.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions DK-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions DK-Toddlers.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions ES-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions ES-Other children.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions FI-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions FI-Other children.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions FI-Toddlers.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions FR-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions FR-Other children.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions GR-Other children.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions HU-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions IE-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions IT-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions LV-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions NL-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions NL-Other children.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions NL-Toddlers.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions RO-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions SE-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions SE-Other children.zip	EFSA-CRA/CRA 2023/Consumption
AuthorisedUses	EFSA-CRA-2023-Secondary data.zip	EFSA-CRA/CRA 2023
Compounds	EFSA-CRA-2023-Secondary data.zip	EFSA-CRA/CRA 2023
FoodTranslations	EFSA-CRA-2023-Secondary data.zip	EFSA-CRA/CRA 2023

Table 4.12: Datasources for EU Prospective Dietary Cumulative Risk Assessment (2023).

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Table Group	Name	Repository
Foods	EFSA-CRA-2023-Secondary data.zip	EFSA-CRA/CRA 2023
MaximumResidueLimits	EFSA-CRA-2023-Secondary data.zip	EFSA-CRA/CRA 2023
Processing	EFSA-CRA-2023-Secondary data.zip	EFSA-CRA/CRA 2023
ResidueDefinitions	EFSA-CRA-2023-Secondary data.zip	EFSA-CRA/CRA 2023
SubstanceApprovals	EFSA-CRA-2023-Secondary data.zip	EFSA-CRA/CRA 2023
UnitVariability	EFSA-CRA-2023-Unit variability factors Tier 2.zip	EFSA-CRA/CRA 2023

Table 4.12 – continued from previous page

4.15 EU Prospective Dietary Cumulative Risk Assessment (2024)

This standard action is of type: Single value risks

This standard action allows you to run and compare the background and Tier II scenarios to probabilistic prospective risk assessment for a newly proposed use (focal substance/food) as defined by the EC working group. This is a risk assessment for four acute effects (NAN, NAM, DAC, DAH) and for three chronic effects (NCN, TCF and TCP) for different consumption surveys/populations. Occurrence datasets for the background are different 3-year cycle datasets containing occurrence data of all member states in the 3-year cycle, or it can be omitted (for comparison) so that only the background exposure is calculated. MRL exceedances can be removed taking into account measurement uncertainty. As an advanced option it is possible to adjust the 99.9th percentile of the exposure using uncertain adjustment factors from the EFSA cumulative risk assessment report 2020. The exposure is calculated probabilistically and the risk is expressed as margin of exposure (MOE) and is compared to a threshold value of 100. You can upload your own focal commodity/focal substance data (+ processing factors/conversion factors).

Table 4.13: Datasources for EU Prospective Dietary Cumulative Risk Assessment (2024).

Table Group	Name	Repository	Туре
AssessmentGroupMember- ships	EFSA-CRA-2024-CAGs level 2.zip	EFSA-CRA/CRA 2024	Fixed
Effects	EFSA-CRA-2024-CAGs level 2.zip	EFSA-CRA/CRA 2024	Fixed
HazardCharacterisations	EFSA-CRA-2024-CAGs level 2.zip	EFSA-CRA/CRA 2024	Fixed
HazardDoses	EFSA-CRA-2024-CAGs level 2.zip	EFSA-CRA/CRA 2024	Fixed
Compounds	EFSA-CRA-2024-Secondary data.zip	EFSA-CRA/CRA 2024	Fixed
FoodTranslations	EFSA-CRA-2024-Secondary data.zip	EFSA-CRA/CRA 2024	Fixed
Foods	EFSA-CRA-2024-Secondary data.zip	EFSA-CRA/CRA 2024	Fixed
Processing	EFSA-CRA-2024-Secondary data.zip	EFSA-CRA/CRA 2024	Fixed
ResidueDefinitions	EFSA-CRA-2024-Secondary data.zip	EFSA-CRA/CRA 2024	Fixed
UnitVariability	EFSA-CRA-2024-Unit variability factors Tier 2.zip	EFSA-CRA/CRA 2024	Fixed

4.16 EU Retrospective Dietary Cumulative Risk Assessment (2023)

This standard action is of type: Single value risks

This standard action will enable you to run retrospective cumulative dietary risk assessment of pesticide residues using the same methods and data as used by EFSA using data from the period 2019-2021.

	Assessment (2023).	
Table Group	Name	Repository
AssessmentGroupMemberships	EFSA-CRA-2023-CAGs level 2.zip	EFSA-CRA/CRA 2023
Effects	EFSA-CRA-2023-CAGs level 2.zip	EFSA-CRA/CRA 2023
HazardCharacterisations	EFSA-CRA-2023-CAGs level 2.zip	EFSA-CRA/CRA 2023
HazardDoses	EFSA-CRA-2023-CAGs level 2.zip	EFSA-CRA/CRA 2023
Concentrations	EFSA-CRA-2023-Concentrations (2019-2021).zip	EFSA-CRA/CRA 2023/Concentratio
Survey	EFSA-CRA-2023-Consumptions AT-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions BE-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions BE-Other children.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions BG-Other children.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions BG-Toddlers.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions CZ-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions CZ-Other children.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions DE-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions DE-Other children.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions DE-Toddlers.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions DK-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions DK-Toddlers.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions ES-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions ES-Other children.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions FI-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions FI-Other children.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions FI-Toddlers.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions FR-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions FR-Other children.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions GR-Other children.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions HU-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions IE-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions IT-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions LV-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions NL-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions NL-Other children.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions NL-Toddlers.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions RO-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions SE-Adults.zip	EFSA-CRA/CRA 2023/Consumption
Survey	EFSA-CRA-2023-Consumptions SE-Other children.zip	EFSA-CRA/CRA 2023/Consumption
AuthorisedUses	EFSA-CRA-2023-Secondary data.zip	EFSA-CRA/CRA 2023
Compounds	EFSA-CRA-2023-Secondary data.zip	EFSA-CRA/CRA 2023
FoodTranslations	EFSA-CRA-2023-Secondary data.zip	EFSA-CRA/CRA 2023
Foods	EFSA-CRA-2023-Secondary data.zip	EFSA-CRA/CRA 2023
MaximumResidueLimits	EFSA-CRA-2023-Secondary data.zip	EFSA-CRA/CRA 2023
Processing	EFSA-CRA-2023-Secondary data.zip	EFSA-CRA/CRA 2023
ResidueDefinitions	EFSA-CRA-2023-Secondary data.zip	EFSA-CRA/CRA 2023
SubstanceApprovals	EFSA-CRA-2023-Secondary data zin	EFSA-CRA/CRA 2023
UnitVariability	EFSA-CRA-2023-Unit variability factors Tier 1 zin	EFSA-CRA/CRA 2023

Table 4.14: Datasources for EU Retrospective Dietary Cumulative Risk Assessment (2023).

4.17 EU Retrospective Dietary Cumulative Risk Assessment (2024)

This standard action is of type: Single value risks

This standard action will enable you to run retrospective cumulative dietary risk assessment of pesticide residues using the same methods and data as used by EFSA using data from the period 2019-2021.

Table Group	Name	Repository	Туре
AssessmentGroupMemberships	EFSA-CRA-2024-CAGs level 2.zip	EFSA-CRA/CRA 2024	Fixed
Effects	EFSA-CRA-2024-CAGs level 2.zip	EFSA-CRA/CRA 2024	Fixed
HazardCharacterisations	EFSA-CRA-2024-CAGs level 2.zip	EFSA-CRA/CRA 2024	Fixed
HazardDoses	EFSA-CRA-2024-CAGs level 2.zip	EFSA-CRA/CRA 2024	Fixed
Compounds	EFSA-CRA-2024-Secondary data.zip	EFSA-CRA/CRA 2024	Fixed
FoodTranslations	EFSA-CRA-2024-Secondary data.zip	EFSA-CRA/CRA 2024	Fixed
Foods	EFSA-CRA-2024-Secondary data.zip	EFSA-CRA/CRA 2024	Fixed
Processing	EFSA-CRA-2024-Secondary data.zip	EFSA-CRA/CRA 2024	Fixed
ResidueDefinitions	EFSA-CRA-2024-Secondary data.zip	EFSA-CRA/CRA 2024	Fixed

Table 4.15: Datasources for EU Retrospective Dietary Cumulative Risk Assessment (2024).

4.18 Risk steatosis from imazalil

This standard action is of type: Risks

Traditional risk assessment often uses animal data to evaluate toxicological limit values, such as the acceptable daily intake (ADI). However, in-vitro tests may help to reduce the use of test animals. In the standard action described here such a traditional risk calculation can be compared to a similar calculation based on in-vitro data. For example, a risk calculation of steatosis as health effect and imazalil as chemical substance can be compared to the toxicological hazard characterisation based on in-vitro data from AdipoRed measurements in human liver cells. In-vitro concentrations are assumed to represent internal liver concentrations. A human physiologically based kinetic model for imazalil is used to extrapolate from in-vitro to in-vivo doses.

In this standard action, risk for steatosis from exposure to imazalil is estimated from in-vivo or in-vitro based hazard characterisations.

Two parameter sets are available:

- 1. estimates based on QSAR models only, and
- 2. estimates based on QSAR models and in-vitro experiments.

The latter parameter set shows that in the long term concentration levels of imazalil in the liver are stationary.

Table Group	Name	Repository	Туре
AdverseOutcomePath- wayNetworks	AOPN-Effects-EffectRelations- 181017.xlsx	Standard Actions/Risk steatosis from imazalil	Fixed
Effects	AOPN-Effects-EffectRelations- 181017.xlsx	Standard Actions/Risk steatosis from imazalil	Fixed
DoseResponseData	BfR-HepaRG-AdipoRed-Single.xlsx	Standard Actions/Risk steatosis from imazalil	Fixed
Concentrations	ConcentrationsSSD_20190129.zip	Standard Actions/Risk steatosis from imazalil	Fixed
Compounds	EuroMix Substances Inventory (v8) (PPPs).zip	Standard Actions/Risk steatosis from imazalil	Fixed
PbkModels	Euromix-Cosmos_Tebby et al ParamB_QSAR_vitro.xlsx	Standard Actions/Risk steatosis from imazalil	Fixed
UnitVariability	Foods coded in FoodEx1.mdb	Standard Actions/Risk steatosis from imazalil	Fixed
HazardCharacterisations	HazardCharacterisation Imazalil.xlsx	Standard Actions/Risk steatosis from imazalil	Fixed
Survey	NL-VCP-RPC 2005-2006 2-6yr.mdb	Standard Actions/Risk steatosis from imazalil	Fixed
Survey	NL-VCP-RPC 2007-2010 7-69yr.mdb	Standard Actions/Risk steatosis from imazalil	Fixed
Survey	NL-VCP-RPC 2010-2012 70+yr.mdb	Standard Actions/Risk steatosis from imazalil	Fixed
FoodTranslations	NL-VCP-RPC Foods.mdb	Standard Actions/Risk steatosis from imazalil	Fixed
Foods	NL-VCP-RPC Foods.mdb	Standard Actions/Risk steatosis from imazalil	Fixed
EffectRepresentations	TestSystems-Responses- EffectRepresentations-181026.xlsx	Standard Actions/Risk steatosis from imazalil	Fixed
Responses	TestSystems-Responses- EffectRepresentations-181026.xlsx	Standard Actions/Risk steatosis from imazalil	Fixed
TestSystems	TestSystems-Responses- EffectRepresentations-181026.xlsx	Standard Actions/Risk steatosis from imazalil	Fixed

Table 4.16: Datasources for Risk steatosis from imazalil.

4.19 Training prospective risk assessment acute Tier II

This standard action is of type: Single value risks

This standard action allows you to run and compare the background and different Tier II approaches to probabilistic prospective risk assessment for a newly proposed use (focal substance/food) as defined by the EC working group. This is an acute risk assessment of a neurological effect (functional alternations of the motor division) for the Dutch toddlers, the Dutch other children, the Bulgarian other children and Italian adults populations. The focal food and substance combination that can be selected are emamectin with peach, acrinathrin with wheat (fictitious data) or pirimicarb with lettuces (fictitious data). The focal exposure can be based on a MRL, a GAP or an actual exposure is calculated. The 99.9th percentile of the exposure has been adjusted using uncertain adjustment factors from the EFSA cumulative risk assessment report 2020. The exposure is calculated probabilistically and the risk is expressed as margin of exposure (MOE) and is compared to a threshold value of 100.
Table Group	Name	Repository	Туре
FocalFoods	FocalConcentrations- MrlScenario.mdb	Standard Actions/Training Acute Prospective Risk Assessment Tier II	Fixed
AssessmentGroupMember- ships	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Acute Prospective Risk Assessment Tier II	Fixed
AuthorisedUses	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Acute Prospective Risk Assessment Tier II	Fixed
Compounds	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Acute Prospective Risk Assessment Tier II	Fixed
Effects	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Acute Prospective Risk Assessment Tier II	Fixed
FoodExtrapolations	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Acute Prospective Risk Assessment Tier II	Fixed
FoodTranslations	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Acute Prospective Risk Assessment Tier II	Fixed
Foods	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Acute Prospective Risk Assessment Tier II	Fixed
HazardDoses	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Acute Prospective Risk Assessment Tier II	Fixed
MaximumResidueLimits	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Acute Prospective Risk Assessment Tier II	Fixed
Processing	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Acute Prospective Risk Assessment Tier II	Fixed
ResidueDefinitions	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Acute Prospective Risk Assessment Tier II	Fixed
DeterministicSubstance- ConversionFactors	SingleValueCalcula- tions.mdb	Standard Actions/Training Acute Prospective Risk Assessment Tier II	Fixed
UnitVariability	UnitVar36.mdb	Standard Actions/Training Acute Prospective Risk Assessment Tier II	Fixed
Concentrations	a_ConcentrationsSSD_NA	Standard Actions/Training Acute Prospective Risk Assessment Tier II	Fixed
Survey	a_ConsumptionsBU.mdb	Standard Actions/Training Acute Prospective Risk Assessment Tier II	Fixed
Survey	a_ConsumptionsIT.mdb	Standard Actions/Training Acute Prospective Risk Assessment Tier II	Fixed
Survey	a_ConsumptionsNL2.mdb	Standard Actions/Training Acute Prospective Risk Assessment Tier II	Fixed
Survey	a_ConsumptionsNL3_6.mc	Standard Actions/Training Acute Prospective Risk Assessment Tier II	Fixed

Table 4.17: Datasources for Training prospective risk assessment acute Tier II.

4.20 Training prospective risk assessment chronic Tier II

This standard action is of type: Single value risks

This standard action allows you to run and compare the background and different Tier II approaches to probabilistic chronic prospective risk assessment for a newly proposed use (focal substance/food) as defined by the EC working group. This is a chronic risk assessment of an effect (hypothyroidism) for the Dutch toddlers, Dutch other children and German adults populations. The focal food and substance combination that can be selected are cyprodinil with apples (fictitious data), valifenalate with lettuce or valifenalate with aubergines. The focal exposure can be based on a GAP or actual exposure scenario as defined by the working group, or it can be omitted (for comparison) so that only the background exposure is calculated. The 99.9th percentile of the exposure has been adjusted using uncertain adjustment factors from the EFSA cumulative risk assessment report 2020. The exposure is calculated

probabilistically and. the risk is expressed as margin of exposure (MOE) and is compared to a threshold value of 100.

Table Group	Name	Repository	Туре
FocalFoods	FocalConcentrations- GAPScenario.mdb	Standard Actions/Training Chronic Prospec- tive Risk Assessment Tier II	Fixed
AssessmentGroupMem- berships	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Chronic Prospec- tive Risk Assessment Tier II	Fixed
AuthorisedUses	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Chronic Prospec- tive Risk Assessment Tier II	Fixed
Compounds	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Chronic Prospec- tive Risk Assessment Tier II	Fixed
Effects	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Chronic Prospec- tive Risk Assessment Tier II	Fixed
FoodExtrapolations	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Chronic Prospec- tive Risk Assessment Tier II	Fixed
FoodTranslations	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Chronic Prospec- tive Risk Assessment Tier II	Fixed
Foods	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Chronic Prospec- tive Risk Assessment Tier II	Fixed
HazardDoses	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Chronic Prospec- tive Risk Assessment Tier II	Fixed
MaximumResidueLimits	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Chronic Prospec- tive Risk Assessment Tier II	Fixed
Processing	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Chronic Prospec- tive Risk Assessment Tier II	Fixed
ResidueDefinitions	SecondaryInput- DataMRL2016.mdb	Standard Actions/Training Chronic Prospec- tive Risk Assessment Tier II	Fixed
DeterministicSubstance- ConversionFactors	SingleValueCalcula- tions.mdb	Standard Actions/Training Chronic Prospec- tive Risk Assessment Tier II	Fixed
Concentrations	c_ConcentrationsSSD_TCF	Standard Actions/Training Chronic Prospec- tive Risk Assessment Tier II	Fixed
Survey	c_ConsumptionsDE.mdb	Standard Actions/Training Chronic Prospec- tive Risk Assessment Tier II	Fixed
Survey	c_ConsumptionsNL2.mdb	Standard Actions/Training Chronic Prospec- tive Risk Assessment Tier II	Fixed
Survey	c_ConsumptionsNL3_6.md	Standard Actions/Training Chronic Prospec- tive Risk Assessment Tier II	Fixed

Table 4.18: Datasources for Training prospective risk assessment chronic Tier II.

4.21 Training substance prioritisation acute neuro

This standard action is of type: Risks

This standard action allows to prioritise substances to be included in a cumulative risk assessment by inspecting a list of (semi-)probabilistic hazard quotients, ranked from high to low. This is a risk assessment for either all neurotoxicological effects in the Dutch toddler population (level 1), or one of the effect subgroups (NAM or NAN) (level 2). The hazard quotient (termed hazard index in MCRA) compares a percentile from the dietary exposure distribution (P99 or P99.9 can be selected) to the ARfD, which a fixed value for the hazard characterisation (hence the designation as semi-probabilistic).

Table Group	Name	Repository	Туре
AssessmentGroup- Memberships	ActiveSubstances.xlsx	Standard Actions/Training Substance Prioritisa- tion Acute Neuro	Fixed
MaximumResidueLim- its	ConcentrationLimits.xlsx	Standard Actions/Training Substance Prioritisa- tion Acute Neuro	Fixed
Effects	Effects.xlsx	Standard Actions/Training Substance Prioritisa- tion Acute Neuro	Fixed
FoodExtrapolations	FoodExtrapolations.xlsx	Standard Actions/Training Substance Prioritisa- tion Acute Neuro	Fixed
FoodTranslations	FoodRecipes.xlsx	Standard Actions/Training Substance Prioritisa- tion Acute Neuro	Fixed
Foods	Foods.xlsx	Standard Actions/Training Substance Prioritisa- tion Acute Neuro	Fixed
HazardCharacterisa- tions	HazardCharacterisa- tions.xlsx	Standard Actions/Training Substance Prioritisa- tion Acute Neuro	Fixed
Processing	ProcessingFactors.xlsx	Standard Actions/Training Substance Prioritisa- tion Acute Neuro	Fixed
AuthorisedUses	SubstanceAuthorisa- tions.xlsx	Standard Actions/Training Substance Prioritisa- tion Acute Neuro	Fixed
ResidueDefinitions	SubstanceConver- sions.xlsx	Standard Actions/Training Substance Prioritisa- tion Acute Neuro	Fixed
Compounds	Substances.xlsx	Standard Actions/Training Substance Prioritisa- tion Acute Neuro	Fixed
UnitVariability	UnitVarPrimo.xlsx	Standard Actions/Training Substance Prioritisa- tion Acute Neuro	Fixed
Concentrations	a_ConcentrationsSSD_Neu	Standard Actions/Training Substance Prioritisa- tion Acute Neuro	Fixed
Survey	a_ConsumptionsNL2.mdb	Standard Actions/Training Substance Prioritisa- tion Acute Neuro	Fixed
Survey	a_ConsumptionsNL3_6.md	Standard Actions/Training Substance Prioritisa- tion Acute Neuro	Fixed

Table 4.19: Datasources for Training substance prioritisation acute neuro.

- Chronic mixture risk assessment of metals
- Chronic cumulative exposure assessment PA
- Chronic cumulative exposure assessment PFAS
- Demo acute cumulative risk assessment
- Acute single substance dietary exposure assessment of carbofuran or chlorpyrifos
- Chronic single substance dietary exposure assessment of lead or atropine
- Demo Human Monitoring Analysis bisphenols
- TDS-based long term dietary exposure and risk assessment
- Long-term dietary exposure and risk of nickel for the Belgian population
- TDS-based long-term exposure and risk assessment of methylmercury for German children
- Acute Cumulative Risk Assessment Craniofacial Alterations (EFSA 2022)
- EU acute cumulative exposure assessment (2018) Tier I and Tier II
- EU chronic cumulative exposure assessment (2018) Tier I and Tier II
- EU Prospective Dietary Cumulative Risk Assessment (2023)
- EU Prospective Dietary Cumulative Risk Assessment (2024)
- EU Retrospective Dietary Cumulative Risk Assessment (2023)

- EU Retrospective Dietary Cumulative Risk Assessment (2024)
- Risk steatosis from imazalil
- Training prospective risk assessment acute Tier II
- Training prospective risk assessment chronic Tier II
- Training substance prioritisation acute neuro

TYPE AND UNIT DEFINITIONS

5.1 Adjustment factor distribution method types

Accepted justment factor distribution method types.

Name	Short name	Aliases	Description
No adjustment factor	None		No adjustment factor.
Fixed adjustment factor	Fixed		Fixed adjustment factor.
LogNormal	LogNormal		Lognormal distribution with parameters a and b and offset c (default $c = 0$).
LogStudents_t	LogStudents_t		Log Students-t distribution with parameters a, b and c and offset d (default $d = 0$).
Beta	Beta		Beta distribution with shape parameters a and b on interval [c, d], (default = $0, 1$).
Gamma	Gamma		Gamma distribution with shape parameter a and rate parameter b with offset = c (default = 0).

Table 5.1.	Unit	definition	for	Adjustment	factor	distribution	method	tunes
	Unit	deminition	101	Aujustinent	lacioi	distribution	memou	types

5.2 Age alignment method

Age alignment method used for the selected exposure source.

Name	Short name	Aliases	Description
Fit to closest available age	Closest	FitClosest	Fit to closest available age.
User defined bins (in years)	Bins	AgeBins	User defined bins.

Table 5.2: Unit definition for Age alignment method.

5.3 Application amount unit

Accepted units for consumer product application amounts.

Table 5.3:	Unit definition	for Application	amount unit.
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Name	Short name	Aliases
kilogram	kg	kg, kilograms, kilogr, G167A
gram	g	g, grams, gr, G148A
milligram	mg	mg, milligrams, milligram

5.4 Assessment group membership calculation methods

Accepted Assessment group membership calculation methods.

Name	Short name	Aliases	Description
Any (crisp)	Any (crisp)		Assign the highest membership value as membership. For crisp memberships, assign positive substance membership if any model indicates positive membership, and negative membership otherwise.
Majority	Majority		Assign positive substance membership if the
(crisp)	(crisp)		Assign positive substance membership if the majority of the membership models indicates positive membership, otherwise, the substance is considered not to be in the assessment group.
Ratio	Ratio		
(probabilistic)	(probabilistic)		Express substance membership as a probability ranging from zero (certainly out) to one (certainly in), computed as the average membership score.
Bayesian (probabilistic)	Bayesian (probabilistic)		Express substance memberships as a probability with values ranging from zero (certainly out) to one (certainly in) computed using a Bayesian approach.

Table 5.4: Unit definition for Assessment group membership calculation methods.

5.5 Benchmark response type

Accepted benchmark response types.

Name	Short name	Aliases	Description
Fraction change	Fraction change	Fraction- Change, FactorChange	The benchmark response is defined as a fraction change of the background response (i.e., defined for both increase and decrease). E.g., for a factor of 0.1, the benchmark response is at +/- 10% of background response.
Percentage change	Percentage change	Percent- ageChange	The benchmark response is defined as a percentage change of the background response (i.e., defined for both increase and decrease). E.g., for a percentage of 10, the benchmark response is at +/- 10% of background response.
Fraction of background response	Fraction of background	Factor, Fraction, Facto- rOfBackground	The benchmark response is defined as a fraction of the background response. E.g., for a factor of 0.9, the benchmark response is at 0.9 times the background response (i.e., a decrease).
Percentage of background response	Percentage of background	Percentage, PercentageOf- Background	The benchmark response is defined as a percentage of the background response. E.g., for a percentage of 90, the benchmark response is at 90% of the background response (i.e., a decrease).
Extra risk	ER	ExtraRisk	For quantal response types. The benchmark dose is defined as the dose that corresponding with an extra risk of a factor times the background risk. A factor of 0.05 corresponds with 5% extra risk.
Additional risk	AR	AdditionalRisk	For quantal response types. The benchmark dose is defined as the dose that corresponding with an additional risk of a factor times the background risk. A factor of 0.05 corresponds with 5% additional risk.
ED50	ED50	ED50	For quantal response types. The benchmark dose is defined as the dose that corresponds with an estimated risk of 50% (ED50).
Absolute threshold value	Threshold value	Absolute	The benchmark dose is defined as an absolute threshold value.
Absolute difference	Absolute difference	Difference	The benchmark dose is defined an absolute difference with the background risk.

Table 5.5: Unit definition for Benchmark response type.

5.6 Biological matrix

Accepted types of biological matrices.

Name	Short name	Aliases	Description
Whole body	Whole body	Body, Whole body, WholeBody, CTotal	Use whole body when the results apply to the whole body.
Blood	Blood	Blood, Whole blood, WholeBlood, Full blood, FullBlood, BWB, CPlasmaOut	The complete blood from a standard blood donation.
Blood serum	Serum	Serum, BloodSerum, Blood serum, BS	The portion of blood plasma that excludes clotting factors.
Blood plasma	Plasma	Plasma, BloodPlasma, Blood plasma, O_CP, BP	The liquid component of blood, in which erythrocytes are suspended.
Cord blood	Cord blood	CordBlood, Cord blood, CBWB	Blood that remains in the placenta and in the attached umbilical cord after childbirth.
Cord blood serum	Cord blood serum	CordBlood- Serum, Cord blood serum, CBS	The portion of cord blood plasma that excludes clotting factors.
Cord blood plasma	Cord blood plasma	CordBlood- Plasma, Cord blood plasma, CBP	The liquid component of cord blood, in which erythrocytes are suspended.
Venous blood	Venous blood	VenousBlood, Venous blood, O_CV, CVen	The deoxygenated blood which travels from the peripheral blood vessels.
Venous plasma	Venous plasma	VenousPlasma	Plasma of the deoxygenated blood which travels from the peripheral blood vessels.
Arterial blood	Arterial blood	ArterialBlood, Arterial blood, CArt, O_CA	The oxygenated blood in the circulatory system found in the pulmonary vein.
Arterial plasma	Arterial plasma	ArterialPlasma	Plasma of the oxygenated blood in the circulatory system found in the pulmonary vein.
Brain blood	Brain blood	BrainBlood, Brain blood, O_CBrb	The blood present in the brain.
Urine	Urine	Urine, AurinebpaOut, AurinegOut, AurineTotalOut	A non-invasive matrix definition for urine.

Table 5.6: Unit definition for Biological matrix.

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Name	Short name	Aliases	Description
Excreta	Excreta	Excreta	Human waste (i.e., both urine and feces).
Saliva	Saliva	Saliva	The extracellular fluid produced and secreted by salivary glands in the mouth.
Semen	Semen	Semen	The fluid that is emitted from the male reproductive tract and that contains sperm cells.
Breath	Breath	Breath	The condensate of exhaled breath.
Red blood cells	Red blood cells	RedBloodCells	The red blood cells.
Breast milk	Breast milk	BreastMilk	The human milk as a biological system.
Fat (adipose) tissue	Body fat	AdiposeTissue, Adipose Tissue, Body fat, BodyFat, FatTissue, Fat, CFat, O_CF	Adipose tissue, body fat, or simply fat is the main reservoir of fat in the body beneath the skin.
Hair	Hair	Hair	The human hair as a non-invasive matrix for biomarkers of exposure.
Toenails	Toenails	ToeNails	The layer of cells situated at the base of the fingernail or the toenail.
Big toenails	Big toenails	BigToeNails	The layer of cells situated at the base of the big fingernail or the big toenail
Outer skin	Outer skin	OuterSkin	The extracellular connective tissue matrix of the skin.
Amniotic fluid	Amniotic fluid	AmnioticFluid	The amniotic fluid is the protective liquid contained by the amniotic sac, serves as a cushion for the growing fetus.
Placenta tissue	Placenta	Placenta, PlacentaTissue	The tissues which support fetal development.
Uterus tissue	Uterus tissue	UterusTissue, O_CU	The uterus or womb is the organ in the reproductive system that accommodates the fetal development until birth.
Uterus	Uterus	Uterus	The uterus or womb (organ) as biological matrix.
Liver	Liver	Liver, CLiver, O_CL	The liver as biological matrix.
Muscle tissue	Muscle tissue	Muscle, Muscles, MuscleTissue, CPoor, O_CM	Perfused (muscle) tissue.
Viscera tissue	Viscera	Viscera, CRich	Perfused tissue of the viscera, the soft internal organs of the body, especially the intestines.
Skin	Skin	Skin	Skin of the body.
Viable epidermis exposed skin	Viable epidermis exposed skin	ViableEpider- misExposed- Skin, SkinViableEx- posed, ViableSkinEx- posed, CSkin_e	Viable epidermis of skin exposed to air.

Table 5.6 – continued from previous page

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Name	Short name	Aliases	Description
Viable epidermis unexposed skin	Viable epidermis unexposed skin	ViableEpider- misUnexposed- Skin, SkinViableUn- exposed, ViableSkinUn- exposed, CSkin_u	Viable epidermis of skin not exposed to air.
Stratum corneum of exposed skin	Stratum corneum of exposed skin	Stratum- CorneumEx- posed, Stratum- CorneumEx- posedSkin, CSkin_sc_e	The outermost layer of the epidermis, exposed to the open air.
Stratum corneum of unexposed skin	Stratum corneum of unexposed skin	StratumCorneu- mUnexposed, StratumCorneu- mUnexposed- Skin, CSkin_sc_u	The outermost layer of the epidermis, not exposed to the open air.
Kidney	Kidney	Kidney, Kidneys	The kidney organ as biological matix.
Adrenal gland	Adrenal gland	Adrenal, Adrenal gland, Adrenal glands	The adrenal gland as biological matix.
Thyroid gland	Thyroid gland	Thyroid, Thyroid gland	The thyroid gland as biological matix.
Heart	Heart	Heart, O CH	The heart as biological matix.
Lung	Lung	Lung, Lungs, O_CLu	The lung as biological matix.
Brain tissue	Brain tissue	BrainTissue	Brain tissue.
Brain	Brain	Brain	The brain (organ) as biological matix.
Eyes	Eyes	Eye, Eyes, Ocular	The eyes as biological matix.
Gonad tissue	Gonad	Gonad, Gonads, GonadTissue, CGonadOut, O_CBr	The biological matrix of the reproductive system that produces and releases eggs (ovary) or sperm (testicle/testis).
Slowly perfused tissue	Slowly perfused tissue	SlowlyPer- fusedTissue, PoorlyPer- fusedTissue, Slowly perfused tissue, O_CS	The biological matrix of poorly perfused tissue include skin and subcutaneous tissue, and resting muscle.
Richly perfused tissue	Richly perfused tissue	RichlyPer- fusedTissue, Richly perfused tissue, O_CR	The biological matrix of richly perfused tissue include liver, heart, lungs, kidneys, and brain.
Feces	Feces	Excrement, Poop	Bodily waste discharged through the anus.

Table 5.6 - continued from previous page

5.7 Biological organisation type

Accepted biological organisation types.

Name	Short name	Aliases	Description
Molecular	Molecular	Molecular	Molecular level
Cellular	Cellular	Cellular	Cellular level
Tissue	Tissue	Tissue	Tissue level
Organ	Organ	Organelle, Organ	Organ level
Individual	Individual	Individual	Whole body level
Population	Population	Population	Population level

Table 5 7.	Unit	definition	for	Biological	organisation	type
Table 5.7.	Unit	ucinition	101	Diological	organisation	type.

5.8 Biomarker conversion distribution type

Biomarker conversion distribution types.

Name	Short name	Aliases	Description
LogNormal	LogNormal	LogNormal	Lognormal distribution parameterised by a mean and upper p95 percentile.
Uniform	Uniform	Uniform	Uniform distribution specified by a mean/median and an upper boundary.
Beta	Beta	Beta	Beta distribution parameterised by a mean and a variance.

Table 5.8: Unit definition for Biomarker conversion distribution type.

5.9 BoD Indicator

Different metrics in which burden of disease can be expressed.

Name	Short name	Aliases	Description
DALY	DALY	DALY	Disability-adjusted life year.
Deaths	Deaths	Deaths	Number of deaths.
Cases	Cases	Cases	Number of cases.

Table 5.9: Unit definition for BoD Indicator.

5.10 Body weight unit

Accepted units for person body weights.

Table 5.10:	Unit definition	on for Body	weight unit.

Name	Short name	Aliases
kilogram	kg	kg, kilograms, kilogr, 3, G167A
gram	g	g, grams, gr, 0, G148A

5.11 Boolean type

Accepted boolean types.

Name	Short name	Aliases
True	True	True, Yes, T, Y
False	False	False, F, No, N

Table 5.11: Unit definition for Boolean type.

5.12 Burden of disease approach

Methodologial approach used for estimating the burden of disease attributable to a chemical exposure.

Name	Short name	Aliases	Description
Top-down	Top-down		Top-down approach starting with an estimate of the total burden in the population. A population attributable fraction (PAF) is calculated by applying an exposure response function (ERF) along with observed exposures, thereby estimating the proportion of the overall burden attributable to the chemical exposure. The attributable burden follows from multiplication of the total burden by the PAF.
Bottom-up	Bottom-up		Bottom-up approach starting either by estimating the number of new or excess cases based on observed exposures and the exposure response function (ERF), or by evaluating the expected shift in individual responses (e.g. changes in IQ) resulting from exposure. These are translated into a burden measure – often by multiplying by factors such as duration and disability weight (or a monetary value) to calculate the attributable BoD for indicators like DALYs.

	Table 5.12:	Unit definition	for Burden	of disease	approach.
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5.13 Cluster method type

Accepted cluster method types.

Name	Short name	Aliases	Description
Component selection (SNMU)	NoClustering		Only component selection is performed.
Component selection and population subgrouping (SNMU + k-means clustering)	Kmeans		Component selection followed by K-Means clustering of individuals based on their component exposure. K-means classifies individuals in multiple groups (i.e., clusters), such that individuals within the same cluster are as similar as possible (i.e., high intra-class similarity), whereas individuals from different clusters are as dissimilar as possible (i.e., low inter-class similarity). In k-means clustering, each cluster is represented by its center (i.e, centroid) which corresponds to the mean of points assigned to the cluster.
Component selection and population subgrouping (SNMU + hierarchical clustering)	Hierarchical		Component selection followed by hierarchical (Ward's) clustering of individuals based on their component exposure. Hierachical clustering builds a hierarchy from the bottom-up, and doesn't require to specify the number of clusters beforehand. Hierarchical clustering produces a tree-based representation of the observations known as a dendrogram.

Table 5.13:	Unit definition f	for Cluster	method type.
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5.14 Combination method membership info and PoD presence types

Accepted Combination method membership info and PoD presence types.

Name	Short name	Aliases	Description
Consider active if POD/HC AND (in-silico) memberships indicate active	Intersection		Consider active if POD/HC AND (in-silico) memberships indicate active.
Consider active if POD/HC OR (in-silico) memberships indicates active	Union		Consider active if POD/HC OR (in-silico) memberships indicates active.

Table 5.14: Unit definition for Combination method membership info and PoD presence types.

5.15 Concentration limit value type

Accepted concentration limit value types.

		51
Name	Short name	Aliases
Maximum residue limit	MRL	MRL, MaximumResidueLimit
Proposed maximum residue limit	Proposed-MRL	ProposedMrl, ProposedMaximumResidueLimit

Table 5.15: Unit definition for Concentration limit value type.

5.16 Concentration model types

Accepted Concentration model types.

Name	Short name	Aliases	Description
Empirical	Empirical		Residues are sampled from the empirical distribution. Fallback: zero.
Censored value Spike LogNormal	CVSpike- LogN		A lognormal model (logarithmic transformed values, with parameters mu and sigma^2) is fitted to the positive residues values. LOR information is not used. Fallback (if number of positives less than 2): Empirical, but Maximum Residu Limit for pessimistic assessments.
Censored Spike Truncated LogNormal	CVSpike- TruncLogN		A truncated lognormal model (with parameters mu and sigma^2) is fitted to the positive residues values. The LOR is used to estimate the truncated left tail of the distribution. Fallback: Lognormal non-detect spike.
Censored LogNormal	CensLogN		Advanced. A censored lognormal model (with parameters mu and sigma^2) is fitted to the censored and positives residue values. Note, this model is not available when agricultural use information is used. Fallback: Lognormal non-detect spike.
Zero Spike Censored LogNormal	ZeroSpike- CensLogN		Advanced. A mixture model with zero spike (p0) and censored lognormal model (with parameters mu and sigma^2) is fitted to the censored and positives residue values. Note, this model is not available when agricultural use information is used. Fallback: Censored lognormal.
Censored Spike Maximum Residue Limit	CVSpike- MRL		Censored Spike Maximum Residue Limit.
Summary statistics	Summary statistic		Summary statistics.
LogNormal	LogN		Lognormal model.

Table 5.16: Un	it definition for	Concentration	model types.
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5.17 Concentration unit

Accepted units for substance concentrations.

Name	Short name	Aliases
milligram/kilogram	mg/kg	mg/kg, milligram/kilogram, milligram/kg, milligr/kg, -6, G049A, G061A
kilogram/kilogram	kg/kg	kg/kg, kilogram/kilogram, kilogram/kg, 0, G063A
gram/kilogram	g/kg	g/kg, gram/kilogram, gram/kg, gr/kg, -3, G015A, G060A, G191A
micro- gram/kilogram	µg/kg	μg/kg, ug/kg, microgram/kilogram, microgram/kg, microgr/kg, -9, G050A, G076A
nanogram/kilogram	ng/kg	ng/kg, nanogram/kilogram, nanogram/kg, nanogr/kg, -12, G077A, G080A
picogram/kilogram	pg/kg	pg/kg, picogram/kilogram, picogram/kg, picogr/kg, -15, G081A
kilogram/liter	kg/L	kg/l, kg/L, kilogram/liter, kilogram/litre, G017A
gram/liter	g/L	g/l, g/L, gram/liter, gram/litre, gr/l, gr/L, G016A
milligram/liter	mg/L	mg/l, mg/L, milligram/liter, milligram/litre, milligr/l, milligr/L, G052A, G062A
microgram/liter	µg/L	μg/l, ug/L, microgram/liter, microgram/litre, microgr/l, microgr/L, G051A, G079A
nanogram/liter	ng/L	ng/l, ng/L, nanogram/liter, nanogram/litre, nanogr/l, nanogr/L, G078A
picogram/liter	pg/L	pg/l, pg/L, picogram/liter, picogram/litre, picogr/l, picogr/L
micro- gram/milliliter	µg/mL	µg/ml, ug/mL, microgram/milliliter, microgram/millilitre, microgr/ml, microgr/mL
nanogram/milliliter	ng/mL	ng/ml, ng/mL, nanogram/milliliter, nanogram/millilitre, nanogr/ml, nanogr/mL
milligram/deciliter	mg/dL	mg/dl, milligram/deciliter, milligr/dL, milligr/dl
microgram/gram	µg/g	µg/g, µgram/gram, µg/gr, ug/g, ugram/gram, ug/gr
nanogram/gram	ng/g	ng/g, ngram/gram, ngr/gr

Table 5.17:	Unit	definition	for	Concentration	unit.
10010 5.17.	om	acimition	101	concentration	unit.

5.18 Concentration value type

Accepted concentration value type.

Name	Short name	Aliases	Description
Mean concentration	MC	MeanConcen- tration, Concentration- Mean, MC	Mean value from the residue trials.
Median concentration	MR	MedianConcen- tration, MR, STMR, Super- visedTrialMedi- anResidue	Median concentration / residue value of the positive measurements of the residue trials.
Highest concentration	HR	HighestConcen- tration, HighestResidue, HR	Highest measured residue / concentration value.
Concentration percentile	СР	Percentile	
Limit of quantification	LOQ	LOQ	
Maximum residue limit	MRL	MRL	

Table 5.18: Unit definition for Concentration value type.

5.19 Concentration volume unit

Accepted units for substance concentration volumes.

Table 5 19.	Unit definition	for Concentration	volume unit
14010 5.17.	onn acminion	for concentration	volume unit.

Name	Short name	Aliases
micro- gram/cubimeter	ug/m3	ug/m3, ugram/m3, ugr/m3
microgram/liter	ug/L	ug/L, ugram/L, ugr/L

5.20 Consumer product application amount distribution types.

Consumer product application amount uncertainty distribution types.

trouton types			
Name	Short name	Aliases	Description
LogNormal	LogNormal	LogNormal	Lognormal distribution parameterised by a mean and a coefficient of variation.
Uniform	Uniform	Uniform	Uniform distribution specified by a mean/median and a coefficient of variation.

Table 5.20: Unit definition for Consumer product application amount distribution types..

5.21 Consumer product concentration uncertainty distribution types.

Consumer product concentration uncertainty distribution types.

Table 5.21: Unit definition for Consumer product concentration uncertainty distribution types..

Name	Short name	Aliases	Description
LogNormal	LogNormal	LogNormal	Lognormal distribution parameterised by a mean and a coefficient of variation.
Uniform	Uniform	Uniform	Uniform distribution specified by a mean/median and a coefficient of variation.

5.22 Consumption intake unit

Accepted units for consumption intake amounts.

Table 5.22:	Unit definition	for Consump	tion intake unit.
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Name	Short name	Aliases
gram/kilogram bodyweight/day	g/kg bw/day	g/kg bw, gram/kg bw, g/kg bw/day, gram/kg bw/day, gr/kg bw/day, G212A
gram/day	g/day	gram, grams, g/day, g/day, gram/day, gr/day

5.23 Consumption unit

Accepted units for consumption amounts.

Table 5.23: Unit definition for Consumption unit.

Name	Short name	Aliases
kilogram	kg	kg, kilograms, kilogr, 3, G167A
gram	g	g, grams, gr, 0, G148A

5.24 Consumption value type

Accepted consumption value types.

Name	Short name	Aliases
Large portion	LP	LP, LargePortion
Mean consumption	MC	MC, MeanConsumption
Percentile	Percentile	Percentile, P

Table 5.24: Unit definition for Consumption value type.

5.25 Covariate model types

Accepted Covariate model types.

Name	Short name	Aliases	Description
Only constant	Constant		No relation between exposure and e.g. age or gender.
Only covariable	Covariable		Exposure depends on the covariable, e.g. age.
Only cofactor	Cofactor		Exposure depends on the level of the cofactor, e.g. gender.
Both covariable and cofactor	CovariableCo- factor		Exposure depends on both covariable and cofactor (additive model).
Both covariable and cofactor and interaction	CovariableCo- factorInterac- tion		Exposure depends on both covariable and cofactor and the effect of the covariable differs for different levels of the cofactor (multiplicative model).

Table 5.25: Unit definition for Covariate model types.

5.26 Dietary exposures details level types

Accepted ietary exposures details level types.

Name	Short name	Aliases	Description
Full	Full		Show all details.
Restrict to risk-drivers (dietary exposures screening)	On- lyRiskDrivers		Restrict to detailed output for risk-drivers identified by dietary exposures screening.
Omit foods-as-eaten details	OmitFood- sAsEaten		Restrict to detailed output for modelled foods and substances. Omit foods-as-eaten details.

Table 5.26: Unit definition for Dietary exposures details level types.

5.27 Dose response model type

Accepted dose response model types.

Exp-m1Exp-m1Expm1Exponential model 1Exp-m2Exp-m2Expm2Exponential model 2Exp-m3Exp-m3Expm3Exponential model 3Exp-m4Exp-m4Expm4Exponential model 4Exp-m5Exp-m5Expm5Expm5
Exp-m2Expm2Expm2Exponential model 2Exp-m3Exp-m3Expm3Exponential model 3Exp-m4Exp-m4Expm4Exponential model 4Exp-m5Exp-m5Expm5Exponential model 5
Exp-m3Exp-m3Expm3Exponential model 3Exp-m4Exp-m4Expm4Exponential model 4Exp-m5Exp-m5Expm5Expm5
Exp-m4Expm4Expm4Exponential model 4Exp-m5Exp-m5Expm5Exponential model 5
Exp-m5 Expm5 Expm5 Exponential model 5
Hill-m1 Hillm1 Hillm1 Hill model 1
Hill-m2 Hill-m2 Hillm2 Hill model 2
Hill-m3 Hill-m3 Hillm3 Hill model 3
Hill-m4 Hillm4 Hillm4 Hill model 4
Hill-m5 Hill-m5 Hillm5 Hill model 5
TwoStage TwoStage TwoStage
LogLogist LogLogist
Weibull Weibull
LogProb LogProb LogProb
Gamma Gamma
Logistic Logistic Logistic
Probit Probit Probit
LVM Exp m2 LVM Exp m2 LVM Exp m2
LVM Exp m3 LVM Exp m3 LVM_Exp_M3
LVM Exp m4 LVM Exp m4 LVM Exp M4
$\frac{1}{1} \frac{1}{1} \frac{1}$
$\frac{1}{1} \frac{1}{1} \frac{1}$
$\frac{1}{1} \frac{1}{1} \frac{1}$
LVM Hill m5 LVM Hill m5 LVM Hill m5

Table 5.27: Unit definition for Dose response model type.

5.28 Dose unit

Accepted units for substance doses.

Name	Short name	Aliases
gram/kilogram bodyweight/day	g/kg bw/day	g/kg bw/day, g/kg/day, gram/kg bw/day, gram/kg/day, gr/kg bw/day, G212A
milligram/kilogram bodyweight/day	mg/kg bw/day	mg/kg bw/day, mg/kg/day, milligram/kg bw/day, milligr/kg bw/day, G211A
micro- gram/kilogram bodyweight/day	µg/kg bw/day	μg/kg bw/day, μg/kg/day, microgram/kg bw/day, microgram/kg/day, microgr/kg bw/day, microgr/kg/day, G210A
nanogram/kilogram bodyweight/day	ng/kg bw/day	ng/kg bw/day, ng/kg/day, nanogram/kg bw/day, nanogram/kg/day, nanogr/kg bw/day, nanogr/kg/day, G214A
picogram/kilogram bodyweight/day	pg/kg bw/day	pg/kg bw/day, pg/kg/day, picogram/kg bw/day, picogram/kg/day, picogr/kg bw/day, picogr/kg/day
fem- togram/kilogram bodyweight/day	fg/kg bw/day	fg/kg bw/day, fg/kg/day, femtogram/kg bw/day, femtogram/kg/day, femtogr/kg bw/day, femtogr/kg/day
gram/gram bodyweight/day	g/g bw/day	g/g bw/day, g/g/day, gram/g bw/day, gram/g/day, gr/g bw/day, gr/g/day
milligram/gram bodyweight/day	mg/g bw/day	mg/g bw/day, mg/g/day, milligram/g bw/day, milligram/g/day, milligr/g bw/day, milligr/g/day
microgram/gram bodyweight/day	µg/g bw/day	μg/g bw/day, μg/g/day, microgram/g bw/day, microgram/g/day, microgr/g bw/day, microgr/g/day
nanogram/gram bodyweight/day	ng/g bw/day	ng/g bw/day, ng/g/day, nanogram/g bw/day, nanogram/g/day, nanogr/g bw/day, nanogr/g/day
picogram/gram bodyweight/day	pg/g bw/day	pg/g bw/day, pg/g/day, picogram/g bw/day, picogram/g/day, picogr/g bw/day, picogr/g/day
femtogram/gram bodyweight/day	fg/g bw/day	fg/g bw/day, fg/g/day, femtogram/g bw/day, femtogram/g/day, femtogr/g bw/day, femtogr/g/day
kilogram/day	kg/day	kg/day, kilogram/day, kilogr/day
gram/day	g/day	g/day, gram/day, gr/day
milligram/day	mg/day	mg/day, milligram/day, milligr/day
microgram/day	µg/day	μg/day, microgram/day, microgr/day
nanogram/day	ng/day	ng/day, nanogram/day, nanogr/day
picogram/day	pg/day	pg/day, picogram/day, picogr/day
femtogram/day	fg/day	fg/day, femtogram/day, femtogr/day
kilogram/kilogram	kg/kg	kg/kg, kilogram/kilogram, kilogram/kg, kg/kg bw
gram/kilogram	g/kg	g/kg, gram/kilogram, gram/kg, gr/kg, g/kg bw
milligram/kilogram	mg/kg	mg/kg, milligram/kilogram, milligram/kg, milligr/kg, mg/kg bw, G225A
micro- gram/kilogram	µg/kg	μg/kg, microgram/kilogram, microgram/kg, microgr/kg, μg/kg bw
nanogram/kilogram	ng/kg	ng/kg, nanogram/kilogram, nanogram/kg, nanogr/kg, ng/kg bw
picogram/kilogram	pg/kg	pg/kg, picogram/kilogram, picogram/kg, picogr/kg, pg/kg bw
molar	М	M, mol/L
millimolar	mM	mM, mmol/L
micromolar	μΜ	uM, μ M, umol/L
nanomolar	nM	nM, nmol/L
moles	moles	moles, Moles
millimoles	mmoles	mmoles, mMoles
micromoles	µmoles	umoles, uMoles
nanomoles	nmoles	nmol, nmoles, nMoles
gram/kilogram bodyweight/week	g/kg bw/week	g/kg bw/week, gram/kg bw/week, gr/kg bw/week, G218A

Table 5.28: Unit definition for Dose unit.

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Name	Short name	Aliases
milligram/kilogram bodyweight/week	mg/kg bw/week	mg/kg bw/week, milligram/kg bw/week, milligr/kg bw/week, G217A
micro- gram/kilogram bodyweight/week	µg/kg bw/week	μg/kg bw/week, microgram/kg bw/week, microgr/kg bw/week, G216A
nanogram/kilogram bodyweight/week	ng/kg bw/week	ng/kg bw/week, nanogram/kg bw/week, nanogr/kg bw/week, G215A
picogram/kilogram bodyweight/week	pg/kg bw/week	pg/kg bw/week, picogram/kg bw/week, picogr/kg bw/week
fem- togram/kilogram bodyweight/week	fg/kg bw/week	fg/kg bw/week, femtogram/kg bw/week, femtogr/kg bw/week
gram/gram bodyweight/week	g/g bw/week	g/g bw/week, gram/g bw/week, gr/g bw/week
milligram/gram bodyweight/week	mg/g bw/week	mg/g bw/week, milligram/g bw/week, milligr/g bw/week
microgram/gram bodyweight/week	µg/g bw/week	µg/g bw/week, microgram/g bw/week, microgr/g bw/week
nanogram/gram bodyweight/week	ng/g bw/week	ng/g bw/week, nanogram/g bw/week, nanogr/g bw/week
picogram/gram bodyweight/week	pg/g bw/week	pg/g bw/week, picogram/g bw/week, picogr/g bw/week
femtogram/gram bodyweight/week	fg/g bw/week	fg/g bw/week, femtogram/g bw/week, femtogr/g bw/week
kilogram/week	kg/week	kg/week, kilogram/week, kilogr/week
gram/week	g/week	g/week, gram/week, gr/week
milligram/week	mg/week	mg/week, milligram/week, milligr/week
microgram/week	µg/week	µg/week, microgram/week, microgr/week
nanogram/week	ng/week	ng/week, nanogram/week, nanogr/week
picogram/week	pg/week	pg/week, picogram/week, picogr/week
femtogram/week	fg/week	fg/week, femtogram/week, femtogr/week
kilogram/liter	kg/L	kg/l, kg/L, kilogram/liter, kilogram/litre, G017A
gram/liter	g/L	g/l, g/L, gram/liter, gram/litre, gr/l, gr/L, G016A
milligram/liter	mg/L	mg/l, mg/L, milligram/liter, milligram/litre, milligr/l, milligr/L, G052A, G062A
microgram/liter	µg/L	μg/l, ug/L, microgram/liter, microgram/litre, microgr/l, microgr/L, G051A, G079A
nanogram/liter	ng/L	ng/l, ng/L, nanogram/liter, nanogram/litre, nanogr/l, nanogr/L, G078A
picogram/liter	pg/L	pg/l, pg/L, picogram/liter, picogram/litre, picogr/l, picogr/L
milligram/gram	mg/g	mg/g, milligram/gram
microgram/gram	µg/g	µg/g, microgram/gram
nanogram/gram	ng/g	ng/g, nanogram/gram
picogram/gram	pg/g	pg/g, picogram/gram

Table 5.28 - continued from previous page

5.29 Dust adherence amount uncertainty distribution types.

Dust adherence amount uncertainty distribution types.

Name	Short name	Aliases	Description
LogNormal	LogNormal	LogNormal	Lognormal distribution parameterised by a mean and a coefficient of variation.
Uniform	Uniform	Uniform	Uniform distribution specified by a mean/median and a coefficient of variation.

Table 5.29: Unit definition for Dust adherence amount uncertainty distribution types..

5.30 Dust availability fraction uncertainty distribution types.

Dust availability fraction uncertainty distribution types.

Table 5.30: Unit definition for Dust availability fraction uncertainty distribution types..

Name	Short name	Aliases	Description
LogNormal	LogNormal	LogNormal	Lognormal distribution parameterised by a mean and a coefficient of variation.
Uniform	Uniform	Uniform	Uniform distribution specified by a mean/median and a coefficient of variation.

5.31 Dust body exposure fraction uncertainty distribution types.

Dust body exposure fraction uncertainty distribution types.

Name	Short name	Aliases	Description
LogNormal	LogNormal	LogNormal	Lognormal distribution parameterised by a mean and a coefficient of variation.
Uniform	Uniform	Uniform	Uniform distribution specified by a mean/median and a coefficient of variation.

Table 5.31: Unit definition for Dust body exposure fraction uncertainty distribution types..

5.32 Dustingestion uncertainty distribution types.

Dust ingestion uncertainty distribution types.

Table 5.32: Unit definition for Dustingestion uncertainty distribution types..

Name	Short name	Aliases	Description
LogNormal	LogNormal	LogNormal	Lognormal distribution parameterised by a mean and a coefficient of variation.
Uniform	Uniform	Uniform	Uniform distribution specified by a mean/median and a coefficient of variation.

5.33 Effect metric

Accepted types of effect metrics.

Name	Short name	Aliases	Description
Relative risk	RR	RelativeRisk, Relative, RR, Relative Risk	Use relative risk when the risk is expressed as a relative risk.
Odds ratio	OR	OddsRatio, Odds, OR, Odds Ratio	Use odds ratio when the risk is expressed as an odds ratio.
Negative shift	NegativeShift	NegativeShift, Negative Shift	Use negative shift when the risk is expressed as a negative shift in a continuous variable.
Positive shift	PositiveShift	PositiveShift, Positive Shift	Use positive shift when the risk is expressed as a positive shift in a continuous variable.

5.34 Estimates nature types

Accepted Estimates nature types.

Name	Short name	Aliases	Description
Realistic	Realistic		For lognormal: no censoring at the value of the composite sample concentration, no upper limit to the unit concentration. For Beta: no censoring at the value of the composite sample concentration, unit values are never higher than the number of units in composite sample * value of composite sample concentration.
Conservative	Conservative		For lognormal: unit values will be left-censored at the value of the composite sample concentration, no upper limit to the unit concentration. For Beta: unit values will be left-censored at the value of the value of composite sample concentration, unit are values never higher than the number of units in composite sample * value of composite sample concentration.

Table 5.34:	Unit definition	for Estimates	nature types.
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5.35 Exposure approach types

Accepted Exposure approach types.

Name	Short name	Aliases	Description
Risk based (RPFs)	Risk based (RPFs)		Exposures are multiplied by the RPF and thus exposures to different substances are on the same and comparable scale.
Standardised	Standardised		All substances are standardised to equal variance (selection of the components will work on patterns of correlation only).
Unstandard- ised (RPFs = 1)	ExposuresUW		Exposures as such are taken (unstandardised). This is equivalent to RPFs equal to 1. Thus exposures to different substances are not on the same and comparable scale anymore.

Table 5.35: Unit definition for Exposure approach types.

5.36 Exposure calculation method

Method types for obtaining exposure estimates.

Name	Short name	Aliases	Description
Modelled exposures	Modelled exposures		Compute risks from dietary and/or non-dietary routes, aggregated.
Human biomonitoring concentrations	Human biomonitoring concentrations		Compute risks based on exposure estimates obtained from human biomonitoring concentrations.

Table 5.36:	Unit definition	for Exposure	calculation	method
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5.37 Exposure events and dose generation

Exposure events generation methods for PBK model simulations.

Table 5.37:	Unit definition	for Exposure	events and dose	generation.
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Name	Short name	Aliases	Description
Use usual (average) exposures per route	Average daily exposure		Use usual (average) exposures per route, only one event per day.
Use (randomly selected) daily exposures per route	Random daily exposures		Use (randomly selected) daily exposures per route and specify events.

5.38 Exposure grouping method

Method used for grouping exposures in bins.

Name	Short name	Aliases	Description
Custom bins	Custom bins		Define custom bins by providing bin boundaries.
ERF-defined bins	ERF-defined bins		Use bins defined by categories in the exposure response function.

Table 5.38: Unit definition for Exposure grouping method.

5.39 Exposure method types

Accepted Exposure method types.

Name	Short name	Aliases	Description
Manual	Manual		Exposure levels are determined by explicit specification.
Automatic	Automatic		Exposure levels are generated automatically based on the estimated exposure distribution.

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14010 0.07	· Om	actinition	101	Enposare	memou	c peo

5.40 Exposure path type

Accepted paths of exposure to chemicals. A path is a combination of a route and the class of exposure sources (i.e., dietary or non-dietary).

Name	Short name	Aliases	Description
Dietary exposure	Dietary	Dietary	Dietary exposure.
Oral exposure	Oral	Oral	Oral exposure (both from dietary and non-dietary sources).
Dermal exposure	Dermal	Dermal	Dermal exposure.
Inhalation exposure	Inhalation	Inhalation	Inhalation exposure.
At target	At target	AtTarget	Exposures directly at the target (organ).

Table 5.40: Unit definition for Exposure path type.

5.41 Exposure response type

Accepted exposure response types.

Name	Short name	Aliases	Description
Function	Function	Function	Exposure response expressed as function.
PerDoubling	PerDoubling	PerDoubling,	Exposure response expressed as increase per
		Doubling	doubling of exposure.
PerUnit	PerUnit	PerUnit, Unit	Exposure response expressed as increase per increase of 1 unit of exposure
Constant	Constant	Constant	Exposure response expressed as constant
Constant	Consum	Constant	value.

Table 5.41: Unit definition for Exposure response type.

5.42 Exposure route

Accepted routes of exposure to chemicals (i.e., oral, dermal, or inhalation).

Name	Short name	Aliases	Description
Oral	Oral	Oral, Dietary	Contact to a chemical via ingestion.
Dermal	Dermal	Dermal	Exposure via contact between a chemical and the skin.
Inhalation	Inhalation	Inhalation	Exposure via contact between a chemical and the respiratory system.

Table 5.42: Unit definition for Exposure route.

5.43 Exposure source

Accepted types of exposure sources, i.e., the origin of the substance(s) for the purpose of an exposure or risk assessment.

Name	Short name	Aliases	Description
Diet	Diet	Diet	Dietary exposure sources.
Air	Air	Air	Air exposure sources.
Dust	Dust	Dust	Dust exposure sources.
Soil	Soil	Soil	Soil exposure sources.
Consumer products	Consumer products	ConsumerProd- ucts	Consumer product exposure sources.
Other (non-diet)	Other (non-diet)	Other non-dietary	Other non-dietary exposure sources.

Table 5.43: Unit definition for Exposure source.

5.44 Exposure type

Accepted exposure types.

Table 5.44:	Unit	definition	for	Exposure type	e.
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Name	Short name	Aliases	Description
Acute	Acute	Acute	Acute exposure.
Chronic	Chronic	Chronic	Chronic exposure.

5.45 Expression type

Expression types define the way in which a substance amount may be standardised, or possible other way expressed. For example, a substance in urine may be expressed in terms of a creatinine standardisation.

Name	Short name	Aliases	Description
None	None	None	No expression type.
Lipids	Lipids	Lipids	Standardise lipid-soluble substances by total lipid content.
Creatinine	Creat	Creatinine	Standardise substance concentrations by creatinine content.
Specific gravity	SG	SpecificGravity, SpecificGravi- tyNormalised, Specific gravity, Specific gravity normalised, SG, SGNormalised, SG normalised	Normalise substance concentrations to specific gravity.

Table 5.45: Unit definition for Expression type.

5.46 External exposure unit

Accepted units for external exposures.

Name	Short name	Aliases
gram/kilogram bodyweight/day	g/kg bw/day	g/kg bw/day, g/kg/day, gram/kg bw/day, gr/kg bw/day, G212A
milligram/kilogram bodyweight/day	mg/kg bw/day	mg/kg bw/day, mg/kg/day, milligram/kg bw/day, milligr/kg bw/day, G211A
micro- gram/kilogram bodyweight/day	µg/kg bw/day	μg/kg bw/day, μg/kg/day, microgram/kg bw/day, microgr/kg bw/day, G210A
nanogram/kilogram bodyweight/day	ng/kg bw/day	ng/kg bw/day, ng/kg/day, nanogram/kg bw/day, nanogr/kg bw/day, G214A
picogram/kilogram bodyweight/day	pg/kg bw/day	pg/kg bw/day, picogram/kg bw/day, picogr/kg bw/day
fem- togram/kilogram bodyweight/day	fg/kg bw/day	fg/kg bw/day, fg/kg/day, femtogram/kg bw/day, femtogr/kg bw/day
gram/gram bodyweight/day	g/g bw/day	g/g bw/day, g/g/day, gram/g bw/day, gr/g bw/day
milligram/gram bodyweight/day	mg/g bw/day	mg/g bw/day, mg/g/day, milligram/g bw/day, milligr/g bw/day
microgram/gram bodyweight/day	μg/g bw/day	μg/g bw/day, μg/g/day, microgram/g bw/day, microgr/g bw/day
nanogram/gram bodyweight/day	ng/g bw/day	ng/g bw/day, nanogram/g bw/day, nanogr/g bw/day
picogram/gram bodyweight/day	pg/g bw/day	pg/g bw/day, pg/g/day, picogram/g bw/day, picogr/g bw/day
femtogram/gram bodyweight/day	fg/g bw/day	fg/g bw/day, fg/g/day, femtogram/g bw/day, femtogr/g bw/day
kilogram/day	kg/day	kg/day, kilogram/day, kilogr/day
gram/day	g/day	g/day, gram/day, gr/day, g/d
milligram/day	mg/day	mg/day, milligram/day, milligr/day, mg/d
microgram/day	µg/day	µg/day, microgram/day, microgr/day, ug/d
nanogram/day	ng/day	ng/day, nanogram/day, nanogr/day, ng/d
picogram/day	pg/day	pg/day, picogram/day, picogr/day, pg/d
femtogram/day	fg/day	fg/day, femtogram/day, femtogr/day, fg/d
gram/kilogram	g/kg	g/kg, gram/kg, gr/kg, G015A
milligram/kilogram	mg/kg	mg/kg, milligram/kg, milligr/kg, G061A
micro-	µg/kg	µg/kg, microgram/kg, microgr/kg, G050A
gram/kilogram	0	
nanogram/kilogram	ng/kg	ng/kg, nanogram/kg, nanogr/kg, GU//A
picogram/kilogram	pg/kg	pg/kg, picogram/kg, picogr/kg, GU81A
iem-	Ig/кg	ig/kg, iemtogram/kg, iemtogr/kg
logram/kilogram		

Table 5.46: Unit definition for External exposure unit.

5.47 Focal commodity replacement method types

Accepted Focal commodity replacement method types.

Name	Short name	Aliases	Description
Replace samples with focal commodity samples	Replace samples with focal commodity samples		Replace all samples of the selected focal commodity/commodities.
Append focal commodity samples	Append focal commodity samples		Add the samples of the focal commodity/commodities to the background concentration data.
Replace measurements of focal food/substance combinations with measurements from focal commodity samples	Replace measurements of focal food/substance combinations with measurements from focal commodity samples		Replace the substance concentrations of the background concentrations by substance concentrations from the focal commodity concentration data.
Remove measurements of focal food/substance combinations	Remove measurements of focal food/substance combinations		Remove substance measurements for the selected focal food/substance combinations.
Replace measurements of focal food/substance combinations with proposed concentration limit value from data	Replace measurements of focal food/substance combinations with concentration limit value from data		Replace the substance concentrations of the background concentrations with a proposed concentration limit value from data.
Replace measurements of focal food/substance combinations with proposed concentration limit value	Replace measurements of focal food/substance combinations with proposed concentration limit value		Replace the substance concentrations of the background concentrations with a proposed concentration limit value (in mg/kg).

Table 5.47: Unit definition for Focal commodity replacement method types.

5.48 Function types

Accepted Function types.

Name	Short name	Aliases	Description
Polynomial	Polynomial		A polynomial regression fits a nonlinear relationship between the value of the independent variable (e.g. age) and the corresponding conditional mean of y (here the exposure). A polynomial with a degree of 0 is simply a constant function; with a degree of 1 is a line; with a degree of 2 is a quadratic; with a degree of 3 is a cubic, and so on.
Spline	Spline		A spline fits a nonlinear relationship between the value of the independent variable (e.g. age) and the corresponding conditional mean of y (here the exposure). A spline with a degree of 0 is simply a constant function; with a degree of 1 is a line; with a degree of 2 is a quadratic; with a degree of 3 is a cubic, and so on.

Table 5.48:	Unit definition	for Function types.
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5.49 Harvest application type

Accepted harvest application types.

Table 5.49:	Unit definition	for Harvest	application	type.

Name	Short name	Aliases	Description
Pre-harvest application	Pre-harvest	PreHarvest	Pre-harvest application
Post-harvest application	Post-harvest	PostHarvest	Post-harvest application

5.50 Hazard characterisation type

Accepted hazard characterisation types.

Name	Short name	Aliases
Benchmark dose	BMD	BMD
No observed adverse effect level	NOAEL	NOAEL
Lowest observed adverse effect level	LOAEL	LOAEL
Acceptable daily intake	ADI	ADI
Acute reference dose	ARfD	ARfD, ARfD (from ADI)
No observed effect level	NOEL	NOEL
Tolerable daily intake	TDI	TDI
Tolerable weekly intake	TWI	TWI
Benchmark dose lower confidence limit of 1%	BMDL01	BMDL01
Benchmark dose lower confidence limit of 10%	BMDL10	BMDL10
Human biomonitoring guidance values	HBMGV	HBMGV
Other	Other	Other

Table 5.50: Unit definition for Hazard characterisation type.

5.51 Hazard characterisations calculation method types

Calculation method types for computing hazard characterisations.

Name	Short name	Aliases	Description	
Compute from points of departure	InVivoPods		Compute from points of departure.	
Compute from dose response models	FromBmds		Compute from dose response models.	
Combine point of departure from index substance with RPFs from dose response models	CombineIn- VivoPodInVit- roDrms		In-vivo Points of Departures for index substance, others using RPFs from in-vitro dose response models	

Table 5.51: Unit definition for Hazard characterisations calculation method types.
5.52 Hazard dose imputation method types

Accepted Hazard dose imputation method types.

Name	Short name	Aliases	Description
Munro P5 (TTC approach)	Munro P5 (TTC approach)		Use the P5 of the Munro NOEL collection.
Munro central value	Munro central value		Use an unbiased nominal value from the Munro NOEL collection; draw randomly from this collection in the uncertainty runs.
Available hazard charac- terisations distribution P5	Available hazard charac- terisations distribution P5		Use the P5 of the available points of departure.
Available hazard charac- terisations distribution central value	Available hazard charac- terisations distribution central value		Use an unbiased nominal value from the collection of available points of departure; draw randomly from this collection in the uncertainty runs.

Table 5.52: Unit definition for Hazard dose imputation method types.

5.53 Health effect types

Accepted Health effect types.

Name	Short name	Aliases	Description
Risk	Risk		Health effect is negative (risk).
Benefit	Benefit		Health effect is positive (benefit).

Table 5.53: Unit definition for Health effect types.

5.54 Individual generation method

Accepted methods for generation of individuals for air exposure.

Name	Short name	Aliases	Description
Simulate individuals	Simulate		Simulate individuals based on population data.
Use dietary exposures	UseDietaryEx- posures		Use individuals from dietary exposures.

Table 5.54: Unit definition for Individual generation method.

5.55 Individual generation method

Accepted methods for generation of individuals for dust exposure.

Table 5 55	Unit definition	for Individual	generation method
Table 5.55.	Unit demition	101 marviauai	generation method.

Name	Short name	Aliases	Description
Simulate individuals	Simulate		Simulate individuals based on population data.
Use dietary exposures	UseDietaryEx- posures		Use individuals from dietary exposures.

5.56 Individual generation method

Accepted methods for generation of individuals for soil exposure.

Name	Short name	Aliases	Description
Simulate individuals	Simulate		Simulate individuals based on population data.
Use dietary exposures	UseDietaryEx- posures		Use individuals from dietary exposures.

Table 5.56: Unit definition for Individual generation method.

5.57 Individual property type

Accepted individual property types.

Name	Short name	Aliases	Description
Categorical	Categorical	Categorical	Categorical e.g. blood type A, B, AB, O or region East, West, North, South.
Boolean	Boolean	Boolean	Boolean e.g. yes, no, true, false. See Boolean types unit definitions.
Numeric	Numeric	Numeric	Numeric, real numbers.
Nonnegative	Nonnegative	Nonnegative	Nonnegative real numbers, positive or zero.
Integer	Integer	Integer	Integer, integer numbers.
Nonnega- tiveInteger	Nonnega- tiveInteger	NonnegativeIn- teger	NonnegativeInteger integer numbers, positive or zero.
Month	Month	Month	Month. See Month types unit definitions.
DateTime	DateTime	DateTime	DateTime, period.
Gender	Gender	Gender, Sex	Gender, sex or sexuality. See Gender types unit definitions.
Location	Location	Location	Location, country.
Isced	Isced	Isced, Education	ISCED, education level. See Isced types unit definitions.
JobTask	JobTask	JobTask, WorkTask	Job task. See JobTask types unit definitions.

5.58 Individual subset types

Methods for selecting/matching survey individuals with a specified/scoped population.

Name	Short name	Aliases	Description
Match individuals selection to population definition	Match to population definition		Match individuals selection to population definition.
Ignore population definition (use all individuals in survey)	Ignore population definition		Ignore population definition (use all individuals in survey).
Match individuals selection to population definition using selected properties only	Match using selected properties		Match individuals selection to population definition using selected properties only.

Table 5.58: Unit definition for Individual subset types.

5.59 Intake model types

Accepted Intake model types.

Name	Short name	Aliases	Description
Observed Individual Means	OIM		Observed Individual Means: just the empirical means over the observed days.
BetaBinomial Normal	BBN		BetaBinomial distribution for frequency of exposure + (transformed) Normal distribution for amounts (de Boer et al. 2009).
Logistic- Normal Normal	LNN0		Logistic-Normal distribution for frequency of exposure + (transformed) Normal distribution for amounts.
Logistic- Normal Normal with correlation	LNN		Logistic-Normal distribution for frequency of exposure + (transformed) Normal distribution for amounts. Both models are estimated taking into account the correlation between exposure frequency and amounts.
Iowa State University Foods model	ISUF		Iowa State University Foods model: semiparametric distribution for frequency of exposure + (transformed) Normal distribution for amounts (de Boer et al. 2009, Dodd (1996)).

5.60 Internal model type

Accepted internal model types. PBK model or conversion factor model.

Name	Short name	Aliases	Description
Use kinetic conversion factors	Conversion- FactorModel	ConversionFac- torModel, ConversionFac- tor	Use a low-tier model that uses constant conversion factors read form a data source.
Use PBK models with fallback on kinetic conversion factors	PBKModel	PBKModel, PBK	Use PBK models with fallback on kinetic conversion factors (internal concentrations).
Use PBK models only	PBKModel only	PBKMode- lOnly, PBKOnly	Use PBK models (internal concentrations).

Table 5.60: Unit definition for Internal model type.

5.61 Isced type

Accepted International Standard Classification of Education (ISCED) types.

Name	Short name	Aliases	Description
EarlyChild- hoodEducation	ECE	EarlyChild- hoodEducation, ECE	Early childhood education
PrimaryEdu- cation	PE	PrimaryEduca- tion, PE	Primary education
LowerSec- ondaryEduca- tion	LSE	LowerSec- ondaryEduca- tion, LSE	Lower secondary education
UpperSec- ondaryEduca- tion	USE	UpperSec- ondaryEduca- tion, USE	Upper secondary education
PostSec- ondaryNon- TertiaryEduca- tion	PSNTE	PostSec- ondaryNonTer- tiaryEducation, PSNTE	Post-secondary non-tertiary education
ShortCy- cleTertiaryEd- ucation	SCTE	ShortCycleTer- tiaryEducation, SCTE	Short-cycle tertiary education
BachelorsOrE- quivalentLevel	BOEL	BachelorsOrE- quivalentLevel, BOEL	Bachelor's or equivalent level
MastersOrE- quivalentLevel	MOEL	MastersOrE- quivalentLevel, MOEL	Master's or equivalent level
DocteralOrE- quivalentLevel	DOEL	DocteralOrE- quivalentLevel, DOEL	Docteral or equivalent level

5.62 Job task type

Accepted Job Task types.

Name	Short name	Aliases	Description
BathPlating	BP	BathPlating, BP	Bath plating
Chrome-	CPA	Chrome-	Chrome paint applications
PaintApplica-		PaintApplica-	
tions		tions, CPA	
Machining	MA	Machining, MA	Machining
Welding	WE	Welding, WE	Welding
ThermalSpray-	TS	ThermalSpray-	Thermal spraying
ing		ing, TS	
SteelProduc-	SP	SteelProduc-	Steel production
tion		tion, SP	
Maintenance-	MALW	Maintenance-	Maintenance and laboratory workers
AndLaborato-		AndLaboratory-	
ryWorkers		Workers,	
		MALW	

Table 5.62:	Unit d	efinition	for	Job	task	type.
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5.63 Kinetic conversion factor uncertainty distribution types.

Kinetic conversion factor uncertainty distribution types.

	7 1		
Name	Short name	Aliases	Description
LogNormal	LogNormal	LogNormal	Lognormal distribution parameterised by a mean and upper p95 percentile.
Uniform	Uniform	Uniform	Uniform distribution specified by a mean/median and an upper boundary.
InverseUni- form	InverseUni- form	InverseUniform	Inverse uniform distribution specified by a median and an upper boundary.

Table 5.63: Unit definition for Kinetic conversion factor uncertainty distribution types..

5.64 Left-censored data handling methods

Methods for handling left-censored data (i.e., imputation/substitution).

Name	Short name	Aliases	Description
Replace all left-censored values by zero	By zero		All left-censored measurements are assumed to be zero's (set to 0).
Replace all left-censored values by f * LOR	By f * LOR		All left-censored measurements are substituted by f * LOR where f is a constant. Here, the term LOR (limit-of-reporting) is a generic term for LOD and LOQ. If both LOD and LOQ are available, then LOQ is used, also for non-detects (i.e., measurements reported as <lod).< td=""></lod).<>
Replace non-detects by f * LOD and non- quantifications by LOD + f * (LOQ - LOD)	By f * LOD or by LOD + f * (LOQ - LOD)		Non-detects are replaced by f * LOD; non-quantifications are replaced by LOD + f * (LOQ - LOD), where f is a constant.
Replace non-detects by 0 and non- quantifications by f * LOQ	By 0 or by f * LOQ		Non-detects are replaced by 0; non-quantifications are replaced by f * LOQ, where f is a constant.

Table 5.64: Unit definition for Left-censored data handling methods.

5.65 Mean value correction types

Accepted Mean value correction types.

Table 5.65: Unit definition for 1	Mean value correction types.
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Name	Short name	Aliases	Description
Unbiased	Unbiased		The mean of the lognormal is unbiased (bias correction).
Biased	Biased		The mean of the lognormal is biased (no bias correction).

5.66 Measurement result type

Specifies the type of a measurement result. E.g., a positive value, a non-detect, or missing value.

Name	Short name	Aliases	Description
VAL	VAL	VAL	Positive measurement greater than zero.
LOD	LOD	LOD	Measurement below the limit of detection (LOD).
LOQ	LOQ	LOQ	Measurement below the limit of quantification (LOQ).
MV	MV	MV	Missing value (MV).

Table 5.66: Unit definition for Measurement result type.

5.67 Missing value imputation method types

Missing value imputation method types.

Name	Short name	Aliases	Description
By zero	Set zero		Set missing values to zero.
Impute from data	Impute from data		Replace missing measurements by random other measurements of the same substance, biological matrix and sampling type.
No missing value imputation	No missing value imputation		No missing value imputation, all missing values remain in the data set and samples with missing values will be removed before analysis.

5.68 Modelled foods calculation source types

Accepted Modelled foods calculation source types.

Name	Short name	Aliases	Description
Derive modelled foods from concentrations	DeriveMod- elledFoods- FromSample- BasedConcen- trations		Derive modelled foods from sample based concentration data.
Derive modelled foods from single value concentrations	DeriveMod- elledFoods- FromSingle- ValueConcen- trations		Derive modelled foods from single value concentrations.
Derive modelled foods from concentration limits	UseWorstCa- seValues		Derive modelled foods from concentration limits.

Table 5.68: Unit definition for Modelled foods calculation source types.

5.69 Month type

Accepted months types.

Table 5.69: Unit definition for Month type.

Name	Short name	Aliases
January	Jan	Jan, Januari, 1
February	Feb	Feb, Februari, 2
March	Mar	Mar, 3
April	Apr	Apr, 4
May	May	May, 5
June	Jun	Jun, June, 6
July	Jul	Jul, July, 7
August	Aug	Aug, 8
September	Sep	Sep, Sept, 9
October	Oct	Oct, 10
November	Nov	Nov, 11
December	Dec	Dec, 12

5.70 Multiple substance handling method types

Accepted Multiple substance handling method types.

Name	Short name	Aliases	Description
Combined assessment of selected substances	Combined		Combined assessment of selected substances.
Loop over selected substances	Loop		Loop over selected substances.

Table 5.70: Unit definition for Multiple substance handling method types.

5.71 Network analysis type

Accepted Network analysis types.

Name	Short name	Aliases	Description
No network analysis	No network analysis		No network analysis is applied.
Apply network analysis	Apply network analysis		Network analysis is applied on the substance x component (U) matrix.

Table 5.71: Unit definition for Network analysis type.

5.72 Nondetect imputation method types

Accepted nondetect imputation method types.

Table 5.72: Unit definition for Nondetect imputation method type	for Nondetect imputation method types.
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Name	Short name	Aliases	Description
Replace by LOR/LOQ/LOD	ReplaceLimit		Non-quantifications are replaced by $f * LOR$ or $f * LOD$ or by LOD + $f * (LOQ - LOD)$ where f is a constant.
Impute from censored lognormal distribution	Impute from censoredIn		Replace nondetect measurements by a random draw from the lower (left) tail of the censored lognormal distribution.

5.73 PBK model compartment type

PBK model compartment type definitions.

Name	Short name	Aliases
Other	Other	
alveolar air	alveolar air	
compartment	compartment	
arterial blood	arterial blood	
compartment	compartment	
arterial plasma	arterial plasma	
compartment	compartment	
blood plasma	blood plasma	
compartment	compartment	
fat compartment	fat compartment	
feces compartment	feces	
	compartment	
gut compartment	gut compartment	
gonads	gonads	
compartment	compartment	
heart compartment	heart	
1.1.	compartment	
Kidney	Kidney	
compartment	compartment	
nver compartment	liver	
lung commontmont	luna	
lung compartment	lung	
poorly perfused	poorly perfused	
tissue compartment	tissue	
tissue compartment	compartment	
rest-of-body	rest-of-body	
compartment	compartment	
richly perfused	richly perfused	
tissue compartment	tissue	
F	compartment	
skin compartment	skin	
1	compartment	
stratum corneum	stratum corneum	
exposed skin	exposed skin	
compartment	compartment	
stratum corneum	stratum corneum	
unexposed skin	unexposed skin	
compartment	compartment	
urine compartment	urine	
	compartment	
uterus	uterus	
compartment	compartment	
venous blood	venous blood	
compartment	compartment	
venous plasma	venous plasma	
compartment	compartment	
viable epidermis	viable epidermis	
exposed skin	exposed skin	
compartment	compartment	
viable epidermis	viable epidermis	
unexposed skin	unexposed skin	
compartment	compartment	

Table 5.73: Unit definition for PBK model compartment type.

5.74 PBK model parameter distribution types.

Probability distribution types for PBK model parameters.

Name	Short name	Aliases	Description
LogNormal	LogNormal	LogNormal	Lognormal distribution parameterised by a mean and two coefficients of variation (CVs), one representing variability around the mean and the other reprenting uncertainty of the mean.
Normal	Normal	Normal	Normal distribution parameterised by a mean and two standard deviations (SDs), one representing variability around the mean and the other reprenting uncertainty of the mean.
Logistic Normal	Logistic Normal	LogisticNormal	Logistic Normal distribution parameterised by a mean and two coefficients of variation (CVs), one representing variability around the mean and the other reprenting uncertainty of the mean.
Deterministic	Deterministic	Deterministic, Constant	Deterministic distribution.

Table 5.74: Unit definition for PBK model parameter distribution types..

5.75 PBK model parameter type

PBK model parameter type definitions.

Name	Short name	Aliases
Other	Other	
physiological	physiological	
parameter	parameter	
sex	sex	
age	age	
body weight	body weight	
body surface area	body surface area	
body mass index	body mass index	
height	height	
physicochemical	physicochemical	
parameter	parameter	
molecular weight	molecular weight	
metabolic	metabolic	
parameter	parameter	
partition coefficient	partition coefficient	

Table 5.75: Unit definition for PBK model parameter type.

5.76 PBK model species type

PBK model species type definitions.

Table 5.76:	Unit definition	for PBK	model	species	type.

Name	Short name	Aliases	Description
human	human	Human	Human
Mice	Mice	Mice	Mice

5.77 PBK simulation methods

PBK simulation methods.

Name	Short name	Aliases	Description
Standard PBK model simulation	Repeated		Standard PBK model simulation, starting at current age and run simulation for specified time period.
Lifetime model simulation from birth to current age	LifeTimeCur- rentAge		Run (lifetime) PBK model simulation from birth until current age.
Lifetime model simulation from birth until specified age	LifeTimeSpec- ifiedAge		Run (lifetime) PBK model simulation from birth until specified age.

Table 5.77: Unit definition for PBK simulation methods.

5.78 Point of departure type

Accepted point of departure types.

Name	Short name	Aliases
Benchmark dose	BMD	BMD
No observed adverse effect level	NOAEL	NOAEL
Lowest observed adverse effect level	LOAEL	LOAEL
No observed effect level	NOEL	NOEL
Median lethal dose	LD50	LD50
Benchmark dose lower confidence limit of 1%	BMDL01	BMDL01
Benchmark dose lower confidence limit of 10%	BMDL10	BMDL10

Table 5.78: Unit definition for Point of departure type.

5.79 Point of departure types

Accepted Point of departure types.

Name	Short name	Aliases	Description
Unspecified (no conversion to common expression type)	FromRefer- ence		Do not convert non-standard point of departures.
BMD (convert all hazard characterisa- tions as BMDs)	BMD		Convert all point of departures to bench mark doses.
NOAEL (convert all hazard charac- terisations as NOAELs)	NOAEL		Convert all point of departures to NOAELs.

Table 5.79: Unit definition for Point of departure types.

5.80 Population alignment method

The population alignment method used for the selected exposure source.

Name	Short name	Aliases	Description
Match completely at random	MatchRandom	MatchRandom	Match completely at random.
Match at individual level	MatchIndivid- ualID	MatchIndividu- alID, MatchIndividu- alId	Match at individual level (on individual ID).
Match at random based on cofactors (stratified at random)	MatchCofac- tors	MatchCofactors	Match at random based on cofactors (stratified at random).

Table 5.80:	Unit definition	for Population	alignment method.
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5.81 Population characteristic distribution types.

Distribution types used to described population characteristics.

Name	Short name	Aliases	Description
Normal	Normal	LogNormal	Normal distribution parameterised by a mean and a coefficient of variation.
LogNormal	LogNormal	LogNormal	Lognormal distribution parameterised by a mean and a coefficient of variation.
Uniform	Uniform	Uniform	Uniform distribution specified by a mean/median and a coefficient of variation.

Table 5.81: Unit definition for Population characteristic distribution types..

5.82 Population characteristic type

Different characteristic types that can be used to describe a population.

Table 5.82: Unit definition for Population characteristic type.

Name	Short name	Aliases	Description
IQ	IQ	IQ	Intelligence quotient.
BirthWeight	BirthWeight	BirthWeight	Body weight of a baby at their birth.

5.83 Processing distribution type

Accepted processing distribution types.

Name	Short name	Aliases	Description
Logistic Normal distribution	LogisticNor- mal	LogisticNormal, 1	Logisticnormal distribution.
Log Normal distribution	LogNormal	LogNormal, 2	Lognormal distribution.

Table 5.83: Unit definition for Processing distribution type.

5.84 Property level type

Accepted property level types.

Table 5.84:	Unit definition	for Property	level type.
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Name	Short name	Aliases	Description
Individual	Individual	Individual	Individual level.
IndividualDay	IndividualDay	IndividualDay	IndividualDay.

5.85 Response type

Accepted response types.

Name	Short name	Aliases	Description
Continuous multiplicative	СМ	Continuous- Multiplicative	Response values are positive real numbers, e.g., weight, size.
Continuous additive	CA	ContinuousAd- ditive	Response values are real numbers, e.g., weight change, temperature.
Binary	В	Binary	Response values have binary outcomes (yes/no, true/false, success/failure, 0/1, etc.).
Quantal	Q	Quantal, Binomial	Response is measured in terms of number of successes out of N possible.
Quantal group	QG	QuantalGroup	Individual responses are measured as binary values, which may be grouped to form a quantal response.
Count	С	Count	Number of items (cells, molecules, deaths, etc.) in given interval/area/volume.
Ordinal	0	Ordinal	Relative scores (or graded scores) useable only for ranking.

Table 5.85: Unit definition for Response type.

5.86 Risk characterisation ratio

The form of the ratio for expressing risk based on an exposure estimate and a hazard threshold value. This can be either exposure/hazard or hazard/exposure.

Name	Short name	Aliases	Description
haz- ard/exposure	H/E		Express risk as the ratio hazard/exposure (e.g., MOE(T)).
expo- sure/hazard	E/H		Express risk as the ratio exposure/hazard (e.g., HI,HQ, RPI).

Table 5.86: Unit definition for Risk characterisation ratio.

5.87 Riskmetric calculation types

Accepted Riskmetric calculation types.

Table 5.87: 1	Unit definition for Riskmetric calculation t	ypes.
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Name	Short name	Aliases	Description
RPF weighted	RPF Weighted		Calculates risk as a single ratio involving cumulative RPF-weighted exposures.
Sum of risk characterisa- tion ratios	Sum of ratios		Calculate risk as sum of ratios of exposure and hazard.

5.88 Selected exposure routes

Selected exposure routes for inclusion in exposure calculations.

Table 5.88: Unit definition for Selected exposure routes.

Name	Short name	Aliases	Description
Dermal exposure	Dermal		Dermal exposure.
Inhalation exposure	Inhalation		Inhalation exposure.

5.89 Sex type

Controlled vocabulary for biological sex (at birth).

Name	Short name	Aliases	Description
Female	F	Female, F, Femme, Frau	Female
Male	Μ	Male, M, Homme, Mann	Male

Table 5.89: Unit definition for Sex type.

5.90 Single value dietary exposures calculation method types

Accepted Single value dietary exposures calculation method types.

Table 5.90: Unit definition for Single value dietary exposures calculation method types.

Name	Short name	Aliases	Description
IESTI	IESTI		IESTI.
IESTI new	IESTI new		IESTI new.
TMDI	TMDI		Theoretical Maximum Daily Intake.
IEDI	IEDI		International Estimated Daily Intake.
Rees–Day model (I)	Rees-Day(I)		Rees-Day model (I).
Rees–Day model (II)	Rees–Day (II)		Rees-Day model (II).

5.91 Single value risk calculation method types

Accepted Single value risk calculation method types.

Table 5.91: Unit definition for Single value risk calculation method types.

Name	Short name	Aliases	Description
From single value dietary exposures	From single value dietary exposures		From single value dietary exposures and hazard characterisations.
As percentile from risks distribution	As percentile		As percentile from risks distribution.

5.92 Soilingestion uncertainty distribution types.

Soil ingestion uncertainty distribution types.

		U	5 51
Name	Short name	Aliases	Description
LogNormal	LogNormal	LogNormal	Lognormal distribution parameterised by a mean and a coefficient of variation.
Uniform	Uniform	Uniform	Uniform distribution specified by a mean/median and a coefficient of variation.

Table 5.92: Unit definition for Soilingestion uncertainty distribution types..

5.93 Standardise blood methods

Accepted Standardise blood methods.

Name	Short name	Aliases	Description
Standardise by total lipid measured via gravimetric analysis	Gravimetric		Standardise by total lipid measured via gravimetric analysis.
Standardise by total lipid measured via enzymatic summation	Enzymatic		Standardise by total lipid measured via enzymatic summation.
Standardise by derived total lipid content of Triglyc- erides/Cholestero (Bernert et al. 2007)	Bernert		Standardise by derived total lipid content (Bernert et al. 2007).

Table 5.93: Unit definition for Standardise blood methods.

5.94 Standardise/normalise urine concentration method

Accepted Standardise urine concentration methods based on creatinine.

Name	Short name	Aliases	Description
Normalise by specific gravity	SGNorm		Normalise by specific gravity. Levine and Fahy (1945).
Standardise by creatinine concentration	CreatStand		Standardise by creatinine concentration.
Normalise by specific gravity derived from creatinine, adults (Carrieri et al. 2001)	SGCreatAdj		A specific gravity adjustment is applied by multiplying a creatinine adjusted concentration with a factor (default 1.48 for adults 18 - 68 year).
Normalise by specific gravity derived from nonlinear modelling of creatinine, children 6 - 14 yr (Busgang et al. 2023, model 1)	SGCreatNon- linearMode- lOne		A specific gravity adjustment is applied by nonlinear modelling of a creatinine adjusted concentration, using model 1 of table 3 of Busgang et al. 2023 (children 6 - 14 year).
Normalise by specific gravity derived from nonlinear modelling of creatinine, children 6 - 14 yr, age and gender dependent (Busgang et al. 2023, model 2)	SGCreatNon- linearMod- elTwo		A specific gravity adjustment is applied by nonlinear modelling of a creatinine adjusted concentration, using the age and gender dependent model 2 of table 3 of Busgang et al. 2023 (children 6 - 14 year).

Table 5.94: Unit definition for Standardise/normalise urine concentration method.

5.95 Substance group selection method types

Accepted Substance group selection method types.

Name	Short name	Aliases	Description
All substances	IncludeAll		Include all substances of the substances table and use hazard characterisation imputation for missing hazard data.
Restrict to available hazard data	RestrictHaz- ardDoseRpf		Restrict to the substances with available hazard data (either in the form of dose response models or RPFs).
Restrict to available hazard data and possible membership	RestrictHaz- ardDoseRp- fAndProbable- Membership		Consider only the substances with available hazard data and non-zero membership (i.e., $P(AG) > 0$).
Restrict to available hazard data and certain membership	RestrictHaz- ardDoseRp- fAndCertain- Membership		Consider only substances with certain assessment group membership (i.e., $P(AG) = 1$) and for which a hazard characterisation is available.
Restrict to non-zero membership	RestrictProba- bleMember- ship		Consider all substances, use TTC based on the Cramer class for the substances for which no limit dose or RPF is defined.
Restrict to certain membership	RestrictCer- tainMember- ship		Consider only the substances with certain assessment group membership (i.e., $P(AG) = 1$).

Table 5.95: Unit definition for Substance group selection method types.

5.96 Substance translation allocation method types

Accepted Substance translation allocation method types.

Name	Short name	Aliases	Description
Random allocation	Random allocation		Random allocation.
Allocate most potent	Allocate most potent		Allocate most potent active substance.
Nominal estimate	Nominal estimate		Allocate nominal estimate (weighted average allocation).
Allocate to all	Allocate to all		Allocate for each active substance independently as if all concentrations were allocated to this active substance.

Table 5.96: Unit definition for Substance translation allocation method types.

5.97 Target dose selection method types

Accepted Target dose selection method types.

Table 5.97:	Unit definition	for Target	dose selection	method types.

Name	Short name	Aliases	Description
Select most toxic	MostToxic		Choose the most toxic (default).
Take aggregate	Aggregate		Choose an aggregated hazard characterisation when there there are multiple available candidates in nominal runs.
Random draw	Draw		Draw a random hazard characterisation.

5.98 Target level type

The exposure target level indicates whether an exposure/dose is assumed to be an internal (target) concentration, an external dose, or an internal (systemic) dose.

Name	Short name	Aliases	Description
External dose	External	External, ExternalDose	The administered or external dose, often referred to as exposure or intake.
Internal (systemic) dose	Systemic	Systemic, InternalDose	The absorbed dose is the amount of chemical that is systemically available and can be regarded as the fraction of the external dose that is absorbed and enters the general circulation.
Internal (target) concentration	Internal	Internal, InternalConcen- tration	Internal dose or target dose (tissue dose) is the amount of chemical present in a specific target of interest, i.c. a concentration.

Table 5 98.	Unit	definition	for Target	level type
1000 5.70.	Omt	actinition	101 Tuiget	iever type.

5.99 Test system type

Accepted test system types.

Table 5.99:	Unit definition	for Test system type.
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Name	Short name	Aliases	Description
In vivo	In vivo	InVivo	In vivo
Cell line	Cell line	CellLine	CellLine
Primary cells	Primary cells	PrimaryCells, StemCells, Embryo	PrimaryCells
Tissue	Tissue	Tissue	Tissue
Organ	Organ	Organ	Organ

5.100 Testing method types

Accepted xxx types.

Name	Short name	Aliases	Description
Backward	Backward		Backward selection starts with selecting a model with a function of the highest degree. Then, the degree of the function is decreased by one and the model is tested again. This process is repeated until decreasing the degree does not improve the model fit anymore.
Forward	Forward		Forward selections starts with selecting a model with a function of the lowest degree. Then, the degree of the function is increased by one and the model is tested again. This process is repeated until increasing the degree does not improve the model fit anymore.

Table 5.100: Unit definition for Testing method types.

5.101 Time unit

Supported time units.

Table 5.101:	Unit definition	for Time unit.
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Name	Short name	Aliases	Description
seconds	s	Seconds, sec	Seconds
minutes	min	Minutes, min	In minutes
hours	h	Hours, h	In hours
days	d	days, d	days

5.102 Transform types

Accepted Transform types.

Name	Short name	Aliases	Description
Logarithmic	Logarithmic		Exposure amounts are transformed to normality using a logarithmic transformation.
No transformation	No transformation		Exposure amounts are not transformed.
Power	Power		Exposure amounts are transformed to normality using a Box-Cox power transformation.

5.103 Uncertainty types

Accepted Uncertainty types.

Table 5.103:	Unit definition	for Uncertainty types.
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Name	Short name	Aliases	Description
Empirical	Empirical		Data are taken as such.
Parametric	Parametric		A parametric model is fitted to the data.

5.104 Unit variability correlation types

Accepted Unit variability correlation types.

Table 5.104: Unit definition for U	Unit variability correlation types.
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Name	Short name	Aliases	Description
No correlation	NoCorrelation		The unit residue values for unit portions (consumption amount/unitweight) are randomly drawn, explicitly ignoring any correlation between unit residues.
Full correlation	FullCorrela- tion		The unit residue values for unit portions (consumption amount/unitweight) are randomly drawn, explicitly introducing correlation between unit residues, e.g. high (small) values occur more frequently together.

5.105 Unit variability model types

Accepted Unit variability model types.

Name	Short name	Aliases	Description
Beta distribution	Beta distribution		Requires knowledge of the number of units in a composite sample, and of the variability between units (realistic or conservative estimates). Under the beta model, the simulated unit values are drawn from a bounded distribution on the interval.
Lognormal distribution	Lognormal distribution		Requires only knowledge of the variability between units (realistic or conservative estimates). The lognormal distribution is considered as an appropriate model for many empirical positive concentration distributions (unbounded distribution).
Bernoulli distribution	Bernoulli distribution		Requires only knowledge of the number of units in a composite sample (results are always conservative). The bernoulli model is a limiting case of the beta model, which can be used if no information on unit variability is available, but only the number of units in a composite sample is known.

Table 5.105: Unit definition for Unit variability model types.

5.106 Unit variability types

Accepted Unit variability types.

Name	Short name	Aliases	Description
Variation coefficient	Variation coefficient		Standard deviation divided by the mean.
Variability factor	Variability factor		Defined as 97.5th percentile divided by the mean.

Table 5.106: Unit definition for Unit variability types.

5.107 Unit weight value type

Accepted unit weight types.

Name	Short name	Aliases	Description
Unit weight RAC	RAC	RAC, UnitWeigh- tRAC, UnitWeigh- tRawAgricul- turalCommod- ity	Unit weight raw agricultural commodity (RAC).
Unit Weight EP	EP	EP, UnitWeightEP, UnitWeightEdi- blePortion	Unit weight edible portion (EP).

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Table $5.10/$:	Unit definitio	n for Unit	weight	value type.

5.108 Value qualifier

Supported value qualifiers.

racie stroot child actinition for that quantier.	Table 5.108:	Unit definition for	Value qualifier.
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Name	Short name	Aliases
=	=	=, Equals
<	<	lt, LessThan, <

5.109 Ventilatory flow rate variability distribution types.

Ventilatory flow rate variability distribution types.

Table 5.109: Unit definition for Ventilatory flow rate variability distribution types..

Name	Short name	Aliases	Description
LogNormal	LogNormal	LogNormal	Lognormal distribution parameterised by a mean and a coefficient of variation.

APPLICATION PROGRAMMING INTERFACE (API)

The entire API interface is described using the Swagger application.

COMMAND LINE INTERFACE (CLI)

7.1 Introduction

The MCRA core library comes with a command line interface (CLI) utility to run MCRA actions using input files and producing output files. Data files and specification of sequences of modules and settings are supplied in an action template. An action template is a local disk folder with a defined structure which is described in *Action template structure*. A new empty action template can be created by the CLI as demonstrated in *Create a new action template*. Alternatively, the template can also be a compressed (zip) file format. Running an action via the CLI is described in *Run an action*. This generates results in local disk output files that can be accessed via, for instance, a web browser as described in *Action template structure*.

The latest release of the CLI utility of MCRA Core is available as a download from GitHub (https://github.com/ rivm-syso/mcra-core/releases) or can be built from the sources.

7.2 Action template structure

The CLI can process either folders or zip files as input. This paragraph explains the structure of the files and subfolders inside the input folder or zip file which the CLI needs to run an MCRA action correctly. This structure is the same for both the contents of a zip file and the contents of a folder. The term base folder will be used to refer to either the zip file or the folder containing the settings and data of the action. The base folder must contain the following XML files:

- _ActionData.xml: contains the configuration of the input data for the action
- _ActionSettings.xml: contains the configuration for the action

The data for the action may be included as CSV files containing MCRA table data directly in the base folder together with the settings files. In this case the _ActionData.xml file is optional. The other option is to put the input data files in a subfolder named Data. In this case the following file formats are accepted:

- Excel files: files with an XLS or XLSX extension;
- Access files: files with an ACCDB or MDB extension;
- Zipped CSV: zip files containing the MCRA data in one CSV file per table.

The files mentioned above must adhere to the data formats of MCRA available in the documentation.

Action settings XML file structure

The model settings XML file must be named '_ActionSettings.xml' (with a leading underscore '_' character). A simple example, configuring a 'Foods' action, is:

```
<project>
    </name>ExampleFoodsAction</name>
    </ActionType>Foods</ActionType>
</Project>
```

The Project tag is the root element of the XML file; it is named Project for historic reasons. This configuration file is an example of a minimalistic configuration in which all other settings have been omitted. By using the *Create a new*

action template option, a configuration file will be automatically generated that contains all possible options for the action.

Data configuration

As mentioned before, MCRA accepts Excel, Access and zipped CSV data files as input when they are included in the Data subfolder. For simple actions there is also the option to include CSV files directly in the root of the folder or zip file. In this case the folder or zip file itself becomes the single input file for the action. The next section lists all possible options. Data for running an action with the CLI is accepted in the following ways: 1. As CSV (comma separated values) files directly in the root of the folder or zip file. In this case, the zip file or folder is itself the single data file, containing the MCRA tables. A data configuration XML file is optional, but it can still be used to filter the table groups that are loaded during the run. 2. A Data subfolder containing one or more of the accepted MCRA file types. In this case a data configuration XML file is necessary to link the MCRA table groups and data files to use.

Data configuration XML file structure

The data configuration XML file must be named _ActionData.xml (with the leading underscore '_' character) and must be placed in the root of the zip file or folder. It describes the links between the modules and data. The link is made based on a so-called MCRA table group definition. A data file can contain data for one or more table groups. An example of a simple data configuration file is as follows:

<pre><datasourceconfiguration xmlns:xsd="http://www.w3.org/2001/XMLSchema</pre"></datasourceconfiguration></pre>
<pre>xmlns:xsi="http://www.w3.org/2001/XMLSchema-instance"></pre>
<datasourcemappingrecords></datasourcemappingrecords>
<datasourcemappingrecord></datasourcemappingrecord>
<pre><sourcetablegroup>Foods</sourcetablegroup></pre>
<name>InputData.xls</name>
<datasourcemappingrecord></datasourcemappingrecord>
<pre><sourcetablegroup>Compounds</sourcetablegroup></pre>
<name>InputData.xls</name>
<datasourcemappingrecord></datasourcemappingrecord>
<pre><sourcetablegroup>Survey</sourcetablegroup></pre>
<name>Consumptions.mdb</name>
<datasourcemappingrecord></datasourcemappingrecord>
<sourcetablegroup>FoodConsumptions</sourcetablegroup>
<name>Consumptions.mdb</name>

The data source configuration contains one collection of DataSourceMappingRecords. Each DataSourceMappingRecord must contain at least the SourceTableGroup. The Name element refers to the file name in the Data subfolder when data files are used. If the zip file or folder contains CSV table files in the root itself, the name is optional. In this case the name of the zip file itself is used.

7.3 Create a new action template

The CLI can generate an empty action template for any type of action that is supported by MCRA. This generated template consists of the correct files and folder structure for the action, with all default settings, and Excel data files with empty tables as a starting point to fill in your own input data.

Use the following command to create a new action of a given type:

mcra.exe create '<name_of_new_action>' -a <action type>

where the option -a specifies the type of action to be created. For example, to create a basic skeleton for analysing your own single value risk estimate with the name MySingleValueRisk, run the command,

mcra.exe create MySingleValueRisk -a SingleValueRisk

The name of the action type, SingleValueRisk in the example above, is a reserved keyword and must match with one of the supported action types. To get a list of all supported action types, run the command,

mcra.exe create -u

This will print out a list of reserved names of all action types. After creating this template, you can start filling in the input data in the Excel files and adjusting the calculation settings in the _ActionSettings.xml file.

7.4 Run an action

The most basic use of the CLI is to specify a folder name or a zip file containing the settings and data for the action that you want to run (see section *Action template structure* for a detailed description). In a terminal window. e.g. PowerShell or the command prompt, run the command:

```
mcra.exe run '<path to action folder>' [options]
or:
mcra.exe run '<name_of_action>.zip' [options]
```

As an example, assume that an action folder has been composed for doing a calculation on the retrospective Tier 2 of cumulative exposure to pesticides. This model can be run by the command:

mcra.exe run 'EU acute cumulative exposure assessment (2018) Tier 2'

The name of the action in this example is equal to the name of the folder that contains the data and settings as input for the CLI, so in this example there is a folder with the name 'EU acute cumulative exposure assessment (2018) Tier 2'. To get an overview of the available options for running the command line interface, run the following command:

```
mcra.exe help run
```

The output of this command lists the options for the run command as shown below (exact details maybe different, depending on the version of the CLI utility).

Parameter	Description
task input name	(pos. 0, required) Name of the input zip file or base folder containing the simulation task to be processed.
-o,output	Base folder for output. Project output will be written to a subfolder, using the project name.
overwrite	(Default: false) Overwrite existing output. If set to false, the output will be written to a uniquely named folder, otherwise the project name is used as output folder
skipreport	(Default: false) Don't render the full output report.
skiptables	(Default: false) Don't generate CSV output tables.
skipcharts	(Default: false) Don't generate SVG charts.
keeptempfiles	(Default: false) Keep temporary (intermediate) files.
-r, randomseed	Use this value as the Monte Carlo random seed for the project.
-i,interactive	(Default: false) Set to true to run in interactive mode.
-s,silent	(Default: false) Set to true to run in silent mode.
dbType	(Default: Csv) Database type. Possible options: - csv: Intermediate data will be written to
	CSV files during the run
help	Display this help screen.
version	Display version information.

Table 7.1: command-line parameters for running an action

Output

By default the CLI utility creates the output files in a subfolder in the location where the CLI command is run, use the '--output' command line option to specify a different output folder.

7.5 Output files and folder structure

The output of an action performed with the CLI is saved to a folder, which by default has the name of the action suffixed with date and time to create a unique folder name. The output of the action consists of metadata, data and image (chart) files. The CLI options '--skipreport', '--skiptables' and/or '--skipcharts' are available to limit the output if desired. The contents of the output folder are as follows.

File	Description
Metadata files	
_CsvFileIn-	Tab delimited text file containing lookup data for the output CSV data files. It contains the
dex.txt	name of the file and the path to the file in the full MCRA output.
_MCRAVer-	Contains detailed MCRA version information of the CLI that was used.
sion.txt	
_TOC.txt	Contains a detailed internal table of contents of references to report sections.
_TOC-	Contains a detailed table of contents of the output charts (if any).
Charts.txt	
_TOC-	Contains a detailed table of contents of the output CSV files.
CsvData.txt	
_TOC-	Contains a detailed table of contents of any output data in XML format.
XmlData.txt	
ProjectOrigi-	A copy of the original action settings file (_ActionSettings.xml)
nalSettings.xml	
ProjectSimulat-	A fully populated XML file containing all settings that were used in the run. This includes the
edSettings.xml	defaults of all other settings that are available in MCRA.
Data files	
FoodsTable.csv	Foods output data table
Substances-	Substances output data table
Table.csv	
Reports	
_Report.html	The MCRA report in HTML format. This report contains all output sections with tables and charts in one page.
Tables/charts	The tables and charts in the HTML file are included in this folder and are referenced from the HTML file.

APPENDICES

8.1 NOEL collection of Munro et al.

This collection is from Munro et al. (1996) and can be downloaded here.

8.2 Box-Cox power transformation

The Box-Cox power transformation is a data transformation to achieve a better normality and to stabilize the variance. In MCRA, the transformation parameter p in $(y^p - 1)/p$ is determined by maximizing the log-likelihood function

$$l(p) = -\frac{n}{s} \log \left[\frac{1}{n} \sum_{i=1}^{n} (y_i^{(p)} - \overline{y^{(p)}})^2 \right] + (p-1) \sum_{i=1}^{n} \log y_i$$

where i indexes the n observations and

$$\overline{y^{(p)}} = \frac{1}{n}\sum_{i=1}^n y_i^{(p)}$$

is the average of the $y_i^{(p)}$, see Box and Cox (1964).

8.3 Gauss-Hermite Integration

8.3.1 One-dimensional Gauss-Hermite integration

Gauss-Hermite integration approximates a specific integral as follows

$$\int\limits_{-\infty}^{\infty} f(x) \exp(-x^2) \mathrm{d}x \approx \sum_{j=1}^{N} w_j f(x_j)$$

in which w_j and x_j are weights and abscissas for N-point Gauss-Hermite integration, see Abramowitz and Stegun (1972). N-point integration is exact for all polynomials f(x) of degree 2N-1, see Dahlquist and Bjorck (1974). This can for instance be used to approximate the mean of a function F(Y) of a normally distributed random variable Y with mean μ and variance σ^2 :

$$\begin{split} &\int_{-\infty}^{\infty} F(x) \frac{1}{\sqrt{2\pi\sigma}} \exp\left(-\frac{(y-\mu)^2}{2\sigma^2}\right) \mathrm{d}y \\ &= \int_{-\infty}^{\infty} F(\mu + \sqrt{2}\sigma x) \frac{1}{\sqrt{\pi}} \exp(-x^2) \mathrm{d}x \\ &= \frac{1}{\sqrt{\pi}} \sum_{j=1}^{N} w_j F(\mu + \sqrt{2}\sigma x_j) \end{split}$$

8.3.2 Two-dimensional Gauss-Hermite integration

One-dimensional Gauss-Hermite integration can readily be extended to two dimensions. The following principal result in two dimensions is more or less given in Jäckel (2005) for the standard bivariate normal distribution $\phi(x, y; \rho)$ with correlation parameter ρ :

$$\int_{-\infty}^{\infty}\int_{-\infty}^{\infty}F(x,y)\phi(x,y;\rho)\mathrm{d}x\mathrm{d}y\approx\frac{1}{\pi}\sum_{i=1}^{N}\sum_{j=1}^{N}w_{i}w_{j}F(\sqrt{2}[ax_{i}+bx_{j}],\sqrt{2}[bx_{i}+ax_{j}])$$

in which

$$a = \frac{\sqrt{1+\rho} + \sqrt{1-\rho}}{2}$$

and

$$b = \frac{\sqrt{1+\rho} - \sqrt{1-\rho}}{2}$$

as given in Jäckel (2005).

Jäckel (2005) discusses other Gauss-Hermite approximations to the two-dimensional integral, but found that the approximation given above generally gives the most accurate results. For the general bivariate normal distribution with means (μ_x, μ_y) and variances (σ_x^2, σ_y^2) the integral can be approximated by means of

$$\frac{1}{\pi}\sum_{i=1}^N\sum_{j=1}^N w_iw_jF(\mu_x+\sigma_x\sqrt{2}[ax_i+bx_j],\mu_y+\sigma_y\sqrt{2}[bx_i+ax_j])$$

The product $w_i w_j$ can be very small, especially when many quadrature points are used, thus wasting possibly precious calculation time. This can be remedied by pruning, i.e. by dropping combinations of (i, j) with very small values of the product $w_i w_j$.

8.3.3 Maximum likelihood for the LNN model with two-dimensional Gauss-Hermite integration

Denote non-consumption on day j for individual i as $Y_{ij} = 0$. The conditional likelihood, i.e. given random effects b_i and v_i , of a non-consumption on day j equals, with H() the inverse of the logit function

$$P(Y_{ij} = 0 | b_i, v_j) = 1 - H(\lambda + v_j).$$

The conditional likelihood of a positive intake $Y_{ij} > 0$ equals, with ϕ the density of the normal distribution

$$f(Y_{ij} = y_{ij} | y_{ij} > 0, b_i, v_i) = H(\lambda + v_i)\phi(y_{ij} - \mu - b_i; 0, \sigma_w^2)$$

The conditional likelihood contribution for individual i is the product of the individual contributions for each day. The marginal likelihood contribution for individual i is obtained by integrating over the possible values of b_i and v_i . Since the pair (b_i, v_i) follows a bivariate normal distribution, the likelihood contribution for individual i can be approximated by means of two-dimensional Gauss-Hermite integration. Individually based covariables, such as sex or age, imply that μ_i and λ_i must be used instead of μ and λ . The likelihood must be optimized by means of some general optimization routine.

8.4 Distributions

8.4.1 Uniform distribution

The uniform distribution is defined on the interval [a, b] where a and b are the minimum and maximum value and a > 0, b > 0.

The conversion factor is the nominal value of the uniform distribution, and

$$factor = \frac{a+b}{2}$$
For the variability upper value, b = upper and a is calculated as:

$$a = 2 \cdot factor - b$$

See also module exposure biomarker conversions, kinetic conversion factors.

8.4.2 Lognormal distribution

The lognormal distribution is characterised by parameters μ and σ , which are the mean and standard deviation on log-scale.

The conversion factor is the nominal value of the lognormal distribution, $\mu = ln(factor)$. The variability upper value specifies the p95 of the standard lognormal. Parameter σ is calculated as:

$$\sigma = \frac{ln(upper) - \mu}{1.645}$$

See also module exposure biomarker conversions, kinetic conversion factors.

8.4.3 Beta distribution

The standard beta distribution is defined on the interval (0, 1) and is usually characterised by two parameters a and b, with a > 0, b > 0 (see e.g. Mood et al. (1974)). Alternatively, it can be parameterised by the mean (conversion factor)

$$\mu = a/(a+b) \in (0,1)$$

and the variance

$$\sigma^2 = ab/(a+b+1)^{-1}(a+b)^{-2}$$

Note that:

$$\frac{ab}{(a+b)^2(a+b+1)} = \frac{\mu(1-\mu)}{(a+b)^2(a+b+1)} < \frac{\mu(1-\mu)}{1} = \mu(1-\mu) \in (0.0.5^2)$$

The conversion factor is the nominal value of the beta distribution, here the mean. The variability upper value is the variance of the beta distribution. The equations solved for a and b show that:

$$a = \left(\frac{1-\mu}{\sigma^2} - \frac{1}{\mu}\right) \cdot \mu^2$$

and

$$b = a \cdot \left(\frac{1}{\mu} - 1\right)$$

See also module exposure biomarker conversions, kinetic conversion factors.

8.4.4 InverseUniform distribution

The inverse uniform distribution is defined on the interval [a, b] where a and b are the minimum and maximum value and a > 0, b > 0.

The conversion factor is the median (nominal) value of the inverse uniform distribution, and the median is equal to

$$factor = \frac{2}{a+b}$$

The variability upper value is equal to $\frac{1}{a}$, so

$$a = \frac{1}{upper}$$

and

$$b = \frac{2}{factor} - a$$

Uniform distributed values are drawn from the interval [a, b] and inverted. See e.g. Inverse distribution and Inverse transform.

See also module exposure biomarker conversions, kinetic conversion factors.

CHAPTER NINE

GLOSSARY

ADI

Acceptable daily intake. The ADI is an estimate of the amount of a substance in food or drinking water that can be consumed daily over a lifetime without presenting an appreciable risk to health. It is usually expressed as milligrams of the substance per kilogram of body weight and day and applies to chemical substances such as food additives, pesticide residues and veterinary drugs.

ADME

An abbreviation for "absorption, distribution, metabolism and excretion", the four key processes which describe how drugs and chemicals get into the body, what happens to them while they are there, and how they are eliminated

AF

Assessment factor

AIC

Akaike Information Criterion.

AOP

Adverse Outcome Pathways. An AOP is a structured representation of biological events leading to adverse effects and is considered relevant to risk assessment.

ARfD

Acute reference dose. Estimate of the amount of a substance in food and/or drinking water, normally expressed on a body weight basis, that can be ingested in a period of 24 h or less without appreciable health risk to the consumer on the basis of all known facts at the time of the evaluation.

AU

Agricultural Use.

BBN

Beta binomial normal model.

BMD

Benchmark dose. A dose or concentration that produces a predetermined change in response rate of an adverse effect (called the benchmark response or *BMR*) compared to background.

BMDL

Benchmark dose lower confidence limit.

BMDU

Benchmark dose upper confidence limit.

BMI

The body mass index is a measurement that expresses the relationship between an individual's weight and height. BMI is calculated by dividing weight in kilograms by height in metres squared (i.e. height x height). Used to assess whether someone's weight is appropriate.

BMR

Benchmark response.

BoD

Burden of disease.

BREAM

Bystander and resident exposure assessment model for plant protection products.

BROWSE

Bystanders, residents, operators and workers exposure models for plant protection products.

BSA

Body surface area.

BW

Body weight.

CA

Concentration Addition. This model is based on a dilution principle, and was designed for chemicals with a similar mechanism of action.

CAG

Cumulative assessment group. A group of chemicals that could plausibly act by a common mode of action, not all of which will necessarily do so. Membership of a CAG can usually be refined (reduced) by application of successively higher tiers of assessment.

ConsExpo

Consumer exposure model.

CRA

Cumulative risk assessment. Risk assessment for combined exposure to two or more chemicals by all relevant pathways and routes.

DA

Dose addition. A process to establish the response of organisms to a mixture of chemicals with similar toxicity. This involves adding up their individual effects to predict the likely impact of the overall mixture.

DALY

Disability-adjusted life year. One DALY represents the loss of the equivalent of one year of full health. DALYs for a disease or health condition are the sum of the years of life lost to due to premature mortality YLLs (*YLL*) and the years lived with a disability YLDs (term:*YLD*) due to prevalent cases of the disease or health condition in a population.

DR

Dose response. The relationship between the amount of a substance to which an individual organism, population or ecosystem is exposed and the way in which it responds (e.g. in terms of toxicity).

E/H

Risk characterisation ratio (Exposure/Hazard).

EBD

Environmental burden of disease.

EEA

European Environmant Agency.

ERF

Exposure response function.

FA

Food additive. A substance deliberately added to foods or beverages for beneficial technological reasons (e.g. to preserve, flavour, colour or ensure a particular texture). Food additives are not normally consumed by themselves nor used as typical ingredients in food.

FC

Focal commodity. A commodity for which an *MRL* is to be set or for which a high residue event has been monitored, and which is therefore the focus of an exposure assessment.

GAP

Good agricultural practice. GAP is a certification system for agriculture, specifying procedures (and attendant documentation) that must be implemented to create food for consumers or further processing that is safe and wholesome, using sustainable methods.

GM

Geometric mean

GSD

Geometric standard deviation

HBM

Human biomonitoring. A direct measurement of the level of toxic chemical compounds present in the body. Often, these measurements are made using blood and urine.

HBGV

Health based guidance value. HBGV is a science-based recommendation for the maximum (oral) exposure to a substance that is not expected to result in an appreciable health risk, taking into account current safety data, uncertainties in these data, and the likely duration of consumption.

HC

Hazard characterisations is a generic term for any reference value for a substances at a chosen biological target level (external or internal) beyond which exposure is associated with potential adverse health effects. Hazard characterisations can be specified as external values (e.g., a human based guidance value, such as an *ADI* or *ARfD*) or are based on a point of departure (*POD*), such as BMDs from dose-response models or externally specified points of departure (*NOAEL*, *LOAEL*, MDS). The computation may involve assessment factors, e.g., for inter-species conversion, intra-species variation or additional sources of uncertainty. The calculation may also use kinetic models or absorption factors to convert external doses to internal doses or vice versa.

HCPs

House Cleaning Products.

HI

The hazard index is the sum of all hazard quotiens (HQ) of the substances that are associated with the same potential adverse health effect.

HQ

The hazard quotient is the ratio of the exposure to a substance and the reference level at which no adverse effects are expected (i.e., exposure divided by the reference level). A HQ smaller than 1 is associated with no expected adverse health effect and a HQ larger than 1 is associated with possible adverse health effects. The HQ is closely related to the MOE, which can be seen as the inverse metric.

HR

Highest residue. The HR is the highest residue level (expressed as mg/kg) in a composite sample of the edible portion of a food commodity when a pesticide has been used according to maximum GAP conditions. The HR is estimated as the highest of the residue values (one from each trial) from supervised trials conducted according to maximum GAP conditions, and includes residue components defined by the JMPR for estimation of dietary intake.

H/E

Risk characterisation ratio (Hazard/Exposure).

ICED

Individual critical effect dose.

In-silico

Research theoretical method, particularly involving computer models, to predict the likely toxicological, or other, effects of substances.

In-vitro

Research method which involves testing cells or tissues extracted from living organisms.

In-vivo

Research method which involves testing individual live animals or populations of live animals.

IVIVE

In vitro to in vivo extrapolation. Refers to the qualitative or quantitative transposition of experimental results or observations made in vitro to predict phenomena in vivo, biological organisms.

JRC

Joint Research Centre

Lipid

Fat and fat-like substance.

LNN

Logistic normal normal model.

LOAEL

Lowest observed adverse effect level.

LOD

Limit of detection. Lowest concentration of a pesticide residue in a defined matrix where positive identification can be achieved using a specified method (IUPAC, 2006).

LOQ

Limit of quantification. Lowest concentration of a pesticide residue in a defined matrix where positive identification and quantification measurement can be achieved using a specified analytical method (IUPAC, 2006).

LOR

Limit of reporting. Practical limit of residue quantification at or above the *LOQ*. The conservative limit of quantification for a defined matrix and method which may vary between laboratories or within the one laboratory from time to time because of different equipment, techniques, and reagents. Commonly either the lower limit of the calibrated range of the method or the lowest level at which quantitative recovery of the analyse has been demonstrated (IUPAC, 2006).

MCR

Maximum cumulative ratio.

MCRA

Monte Carlo Risk Assessment.

MIE

Molecular initiating event.

MoA

Mode of Action.

MOE

The margin of exposure (MOE) is the ratio between the reference level at which no adverse effects are expected to the exposure to a substance (i.e., reference level divided by exposure). Commonly, the reference level is assumed to be a point of departure (*POD*) based on animal studies that does not incorporate all factors to translate to a human reference value. A MOE is therefore typically compared to a uncertainty/safety factor (UF) composed of the product of the uncertainty factors. An MOE is smaller than the UF is associated with risk. A MOE larger than the UF is associated with no expected adverse health effects.

MOET

The harmonic sum of all individual MOEs.

MRA

Mixture risk assessment.

MRL

Maximum residue level. Maximum concentration of a residue that is legally permitted or recognized as acceptable in, or on, a food, agricultural commodity, or animal feedstuff as set by Codex or a national regulatory authority(IUPAC, 2006).

MV

Missing value

NAMs

New approach methodologies. The term NAM has been emerged as a descriptive reference to any non-animalbased approaches that can be used to provide information in the context of chemical hazard and risk assessment

NMF

Non-negative Matrix Factorization

NOAEL

No observed adverse effect level is the greatest concentration or amount of a substance at which no detectable adverse effects occur in an exposed population.

NOEC

No observed effect concentration.

NOEL

No Observed Effect Level

OIM

Observed Individual Means approach. An approach for estimating longer term exposures by taking each individual's observed mean consumption over the duration of a dietary survey.

OP

Occurence pattern

PACEM

Probabilistic Aggregate Consumer Exposure Model. PACEM evaluates aggregate exposure in a population considering individual use and co-use patterns as well as variation in product composition. Product usage data is included on personal care products (*PCPs*) and household cleaning products (*HCPs*).

PAF

Population attributable fraction

PARC

Partnership for the Assessment of the Risk of Chemicals

PBK

Physiologically based kinetic models

PBPK

Physiologically based pharmacokinetic/toxicokinetic models

PCPs

Personal care products

POCE

The probability of critical exposure (PoCE) is the proportion of the HI distribution above the threshold (or of the generalised *MOE* below the threshold) is the probability of critical exposure in the particular (sub)population. The threshold value can be 1 if all assessment factors have already been accounted for in the calculation of HI or MOE.

POD

A point of departure is defined as a point on a toxicological dose-response curve obtained from a dose dose-response experiment in the region at which the curve transitions from no effects to effects. It is used as the base value for deriving toxicological reference values, or hazard characterisations. Common PODs are the no-observed adverse effect level (*NOAEL*) and benchmark dose (*BMD*).

PPP

Plant protection products. Products used to protect, preserve or influence the growth of desirable plants or to destroy or control the growth of unwanted plants or parts of plants.

PRIMo

EFSA pesticide residue intake model.

QSAR

Quantitative structure activity relationship. The quantitative/qualitative structure activity relationships are a set of methods by which the effects of different compounds are related to their molecular structures. It allows the likely adverse or beneficial effects of a particular chemical to be predicted by comparing it with others which have similar structures.

RA

Response Addition. An approach to the risk assessment of mixtures of substances in which responses to each of the individual components are determined and added together in order to predict the response to the mixture as a whole. This approach is only valid if the individual components do not interact with each other, i.e. their effects are completely independent.

RAC

Raw agricultural commodity. Part of a crop used as a food or feed commodity directly from the harvested crop without processing.

RIVM

Rijksinstituut voor Volksgesondheid en Milieu (Dutch National Institute for Public Health and the Environment). (Dutch) National Institute for Public Health and the Environment.

RPC

Raw Primary Commodity.

RPC derivatives

Single-component foods altered by processing.

RPF

Relative potency factor. The ratio of the toxic potency of a given chemical to that of an index chemical in the Cumulative Assessment Group (*CAG*). Relative potency factors are used to convert exposures of all chemicals in the CAG into their exposure equivalents of the index chemical.

RR

Relative risk.

RSExpo

Modelling tool developed by ANSES, to estimate aggregate internal systemic exposures from multiple sources and routes using absorption factors

SA

Standard action

SBML

Systems Biology Markup Language

SD

Standard deviation

SG

Specific gravity of urine

SNMU

Sparse Nonnegative Matrix Underapproximation

SRA

Standard Regulatory Action

SSC

Source/Substance Combination

SSD

EFSA Standard sample description.

TDI

Tolerable daily intake. Is an estimate of the amount of a substance in food or drinking water which is not added deliberately (e.g. contaminants) and which can be consumed over a lifetime without presenting an appreciable risk to health.

TDS

Total diet study. A study designed to estimate the likely consumption of harmful or beneficial substances in the diet. When undertaking such a study, commonly-consumed foods are purchased from shops in a particular country before being analysed.

TEF

Toxic equivalency factor.

ТК

Toxicokinetics. The study of the processes by which potentially toxic substances are handled in the body. This involves an understanding of the absorption, distribution, metabolism and excretion of such substances *ADME*.

ТР

Thermal paper.

TTC

Threshold of toxicological concern. A screening tool that provides conservative exposure limits in the absence of sufficient chemical-specific toxicological data. It is a science-based approach for prioritising chemicals with low-level exposures that require more data over those that can be presumed to present no appreciable human health risk.

TWI

The tolerable weekly intake is the maximum intake of substances in food, such as nutrients or contaminants, that can be consumed weekly over a lifetime without risking adverse health effects.

YLD

Years of healthy life lost due to disability. One YLD represents the equivalent of one full year of healthy life lost due to disability or ill-health.

YLL

Years of life lost from mortality. One YLL represents the loss of one year of life. YLLs are calculated from the number of deaths multiplied by a global standard life expectancy at the age at which death occurs.

Part III

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CHAPTER ELEVEN

COLOPHON



WUR/Biometris, Wageningen University & Research FERA, Food and Environmental Research Agency RIVM, National Institute for Public Health and the Environment

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CHAPTER TWELVE

CHANGE LOG

12.1 Version 10.2.0 (2025-07-07)

12.1.1 Added

- Implementation ConsumerProductExposures module in TargetExposures action (#2148)
- Implementation of individual subset selection in ConsumerProductUseFrequencies (#2217)
- Add Download full HTML report (zip) to output details top right menu
- Add option for lifetime PBK modelling in target exposures module

12.1.2 Fixed

· Create settings HTML instead of PDF in output zip files

12.2 Version 10.1.10 (2025-05-08)

12.2.1 Added

• New setting EbdApproach in ebd module (#2120)

12.2.2 Changed

- Update downloadable excel data template file prospective CRA 2024 (#2196)
- Update prospective scenario options in SRA prospective CRA (2024) (#2198)
- Rename sensitivity analysis option SRA prospective CRA (2024) (#2199)

12.2.3 Fixed

- Multiple selection dialog immediately closes when ticking an option (#2187)
- Cannot select focal concentrations and deterministic conversions data sources simultaneously in prospective SRA 2024 (#2192)

12.3 Version 10.1.9 (2025-04-28)

12.3.1 Added

- New columns for uncertainty in exposure response functions data (#2149)
- New table population characteristics (#2176)
- New column 'population size' in populations table (#2185)
- New module 'Individuals' (#2188)

12.3.2 Changed

• Remove standard action per data source template download button (#2190)

12.3.3 Fixed

• SamplePropertyValues table allows duplicate entries for SampleId and PropertyName (#2114)

12.4 Version 10.1.8 (2025-04-08)

12.4.1 Changed

• Rename exposure effect functions to exposure response functions (#2174)

12.4.2 Fixed

• Hide exposure routes selection in risk action wizard when conditions don't apply (#2181)

12.5 Version 10.1.7 (2025-03-24)

12.5.1 Added

- Add data template download for the user-specified data of the prospective dietary CRA (2024) (#2103)
- Add setting BoD indicator selection for EBD and allow for multiple exposure effect functions (#2122)
- Add option targetLevel in ebd module (#2123)
- Enable dietary exposures as input in ebd module (#2124)
- Add option for selecting custom bins in ebd module (#2125)
- Allow for selecting multiple indicators and eefs in ebd module (#2153)
- Update prospective SRA 2024 with option to run sensitivity analysis (#2164)
- Implementation RS Expo Air Exposures module (#2167)

12.5.2 Changed

- Move PBK model simulation settings to the modules actually running the PBK models (#2118)
- Rename uncertainty analysis options display text in all SRAs (#2159)
- Rename contribution section title in short output HTML in EFSA SRAs (#2160)
- Remove background only option in SRA prospective CRA (2024) from scenario list and make separate advanced checkbox (#2163)

12.5.3 Fixed

- Multiple substances should not be visible for a single substance (#2115)
- Data source selection does not appear after saving a prospective EFSA 2024 SRA (#2155)
- Retrospective SRAs 2023/24 don't work for effect NCN-Chronic (#2158)
- Cloning EFSA SRA prospective 2024 from workspace doesn't copy some settings (#2169)

12.6 Version 10.1.6 (2025-03-04)

12.6.1 Added

- New data module burden of disease infos (#2129)
- Add StandardAction ID column in admin Standard Actions table, optionally visible using a check box
- Remove set frontend limit number of simulated days for PBK model simulations to 25,000 (#2119)

12.6.2 Changed

- Remove deprecated functionality to hard-link additional SBML PBK models
- Change exposure route of dust and soil ingestion to oral

12.6.3 Fixed

• Add documentation for soil exposure module (#2113)

12.7 Version 10.1.5 (2025-02-10)

12.7.1 Added

- Update SRA EU prospective dietary CRA (2024) to use the option to apply conversion of processed derivatives (#2105)
- Implement method to run chronic PBK model simulations using single repeated exposures (#2106)
- New module Soil exposures

12.7.2 Changed

• Refactor 'IsCompute' functionality, move to ModuleConfiguration in project settings XML (#2101)

12.8 Version 10.1.4 (2025-01-31)

12.8.1 Added

- Update concentrations module with functionality to exclude occurrence data at the level of the processed RPC the focal commodity (#2066)
- Focal food/substance functionality for MRL replacement using a settings input field for the proposed MRL (#2085)
- Filter focal food concentrations/samples based on fieldTrialType (#2094)
- Implement default processing factor for missing processing factors (#2095)
- Add option to replace measurements of processed derivatives of the focal food, including correction for processing (#2102)

12.8.2 Changed

• Update SRA EU prospective dietary CRA (2024) with option to choose MRL/GAP/USE scenario (#2092)

12.8.3 Fixed

• Focal food-substance combination not saved in EFSA prospective CRA 2024 (#2087)

12.9 Version 10.1.3 (2024-12-13)

12.9.1 Added

- Implement first draft version of a working templates example in MCRA Web (#1894)
- Documentation on 'combining dietary and non-dietary exposures' (#2040)
- Initial implementation of uploadable PBK models via data module (#2061)
- Initial version of standard action for prospective dietary CRA 2024 (#2068)
- Initial version of standard action for retrospective dietary CRA 2024 (#2079)
- Implement uploading of SBML models (#2072)
- Add ActionType column to MCRA Tasklogs database table (#2077)

12.9.2 Fixed

- Can't send mail from within MCRA to other user (#1846)
- Model-then-add modeltypes and transformations are not saved (#2080)

12.10 Version 10.1.2 (2024-11-08)

12.10.1 Added

- Show last successful and failed login (#1810)
- Implement internal concentration and internal dose (systemic) (#1891)
- Download modular design based on action mappings (#2012)
- Implement expression type for kinetic conversion factors (#2019)
- Implement population alignment methods (#2040)

12.10.2 Changed

- Rename modules: "Human monitoring data" to "HBM occurences" and "Human monitoring analysis" to "HBM analysis" (#2001)
- Hide not-working proast link (#2015)
- Remove enum value AbsorptionFactorModel from InternalModelType, add TargetDoseLevelType.Systemic (#2026)
- Rename HBM Occurences module (previously named human monitoring data) to HBM concentrations (#2039)

12.10.3 Fixed

- Sharing with use rights allows users with administrator role still to download the data (#1934)
- After removing a share, the sharee can still run the action (#1935)
- Terms of use window keeps appearing on repeated logins and doesn't get saved (#2032)
- Create internal exposures action: remove option "Undefined", remove option "External dose" issue 2059, 2060 (#2059, #2060)

12.11 Version 10.1.1 (2024-09-20)

12.11.1 Added

- Implement PbkModels interface
- · Allow plug-in of additional SBML PBK models in frontend and simulation service
- Update frontend with panels and forms for new dust exposure assessment modules

12.11.2 Changed

- Move compartment to target exposures module
- Split kinetic models into AbsorptionFactors (KineticModels) and KineticConversionFactors module
- Remove deprecated and unused setting CodeKineticModel (#2007)

12.11.3 Fixed

- Imported zip of converted full action from standard action opens as standard action (#1984)
- Correct visibility update behaviour of form controls for dynamic setting groups (#2002)
- Notification of new user registration email is not sent (#2016)

12.12 Version 10.1.0 (2024-08-15)

12.12.1 Added

- Automated job to disable inactive user accounts (#1715)
- Add rate limiting for failed login attempts (#1809)

12.12.2 Changed

- Restructure MCRA settings XML format to group settings per module (#581)
- Migrate code base from .NET 6 to .NET 8 (#1833)
- Upgrade R to version 4.4.1 and rtools to 4.4 (#1966)
- EFSA 2023 SRAs pro- and retrospective (#1961)

12.12.3 Fixed

• Cannot convert to target matrix blood when blood is not present as sampling method in HBM data (#1960)

12.13 Version 10.0.15 (2024-06-21)

12.13.1 Changed

• Update EFSA 2023 Standard actions pro- and retrospective (#1961)

12.14 Version 10.0.14 (2024-06-18)

12.14.1 Fixed

- SA Risk steatosis from imazalil seems broken: can't download action+data as zipped CSV (#1948)
- Remove default CodeFocalEffect from Eu 2023 prospective standard action settings
- Update access level when access level was previously set without subfolders (#1928)

12.15 Version 10.0.13 (2024-05-28)

12.15.1 Added

• Add settings tiers for Prospective Dietary CRA (EFSA 2023) (#1924)

12.15.2 Fixed

- Do not show substance weighting of mixtures option in create new action dialog of dietary exposures (#1943)
- Retrospective and prospective SA 2022 defaults, remove checksums from SA definition

12.16 Version 10.0.12 (2024-05-21)

12.16.1 Added

- Add password expiration functionality (#1714)
- Kinetic conversion for specific biomarkers in hbm analysis (#1827)
- Extend registration form with fields for organisation, PARC, and purpose (#1920)
- Documentation: add more info on LOD and LOQ imputation (#1904)

12.16.2 Changed

- EU prospective SA 2023 should use gap scenario and default percentage should be 20
- Place standard actions in alphabetic order in window "Create new standard action" (#1914)
- Decouple MCR settings / analysis in different modules (e.g. MCR in Risks and HBM analysis) (#1931)

12.16.3 Fixed

• Selecting checkbox of apply biomarker conversion no longer auto selects convert to checkbox (#1923)

12.17 Version 10.0.11 (2024-04-12)

12.17.1 Added

- Filtering of hbm data on time points (#1782)
- Running two opex products for compute action single value non-dietary exposures (#1796)
- Specific gravity from creatinine urine adjustment method by Busgang et al. (#1874)
- Option in to skip privacy sensitive outputs in report (#1882)
- New expression type for specific gravity (#1896)
- Implement new option nondetects handling (#1905)

12.17.2 Changed

- Split Busgang urine specific gravity normalisation into age and non-age dependent methods (#1903)
- Draw from censored lognormal using nondetect and nonquantification information (#1904)
- Updates to EFSA CRA 2023 prospective and retrospective standard actions

12.17.3 Fixed

• Remove hbm analysis option to convert using default value 1 (#1829)

12.18 Version 10.0.10 (2024-03-07)

12.18.1 Changed

- Remove conversion factor 1 when target level external in hbm analysis (#1829)
- Update acknowledgement efsa on landing page (#1892)

12.18.2 Added

- Prevent password reuse (#1879)
- Tables for single value non-dietary exposures (#1860)

12.19 Version 10.0.9 (2024-02-27)

12.19.1 Added

- New module single value non-dietary exposures (#1733)
- New standard action prospective EFSA data 2019-2021 (#1856)
- Implement 'use kinetic conversion factor subgroup' setting (#1868)
- Add specific gravity from creatinine adjusted method Carrieri 2001 (#1874)

12.19.2 Changed

• HBM analysis: update biomarker conversion to allow for biomarker conversion subgroups (age/sex) (#1844)

12.19.3 Fixed

- Kernel calculation in R for violin plots fails when vector contains infinities (#1869)
- Human monitoring analysis settings: order of settings is incorrect (#1876)

12.20 Version 10.0.8 (2024-02-12)

12.20.1 Changed

- Update standard action retrospective CRA with EFSA 2019-2021 data (#1823)
- HBM analysis: change order of settings in input form (#1839)

12.21 Version 10.0.7 (2024-02-02)

12.21.1 Added

• First draft version of standard action retrospective SRA with EFSA 2019-2021 data (#1823)

12.21.2 Changed

• Add uncertainty columns to RawkineticConversionFactor table (#1845)

12.21.3 Fixed

• Wrong data source version is shown and used in MCRA for a datasource with multiple versions (#1848)

12.22 Version 10.0.6 (2024-01-26)

12.22.1 Fixed

- Improve loading speed frontend for large concentration datasets (#1542)
- · Add 'clickjacking' protection to MCRA website by hiding content when site is loaded in an IFrame
- PDF fails to render due to large AnalyticalMethods section (#1831)
- Can't delete an MCRA user group when repositories still reference it (#1832)
- Bug in concentration models settings form in frontend: cannot set MRL fallback model (#1834)

12.23 Version 10.0.5 (2023-12-21)

12.23.1 Changed

• Additional changes for MCR analysis in risks module (#1784)

12.24 Version 10.0.4 (2023-12-18)

12.24.1 Added

- Add general information columns (publication title e.o.) to RPF table (#1772)
- Allow for hazard-characterizations to vary as function of a covariate: example age-dependent HBM-GV for PFAS (#1778)
- Implement MCR analysis in risks module (#1784)

12.24.2 Changed

• Rename MarginOfExposure and HazardIndex of type-definition/enum RiskMetricType (#1705)

12.24.3 Fixed

• Use EN-US number input field with decimal point and not decimal comma (#1781)

12.25 Version 10.0.3 (2023-12-01)

12.25.1 Added

- Terms of use and disclaimer (#1583)
- Kinetic conversion models for metabolites (#1683)
- Extend standard action mra metals for effect dnt (#1689)
- Implement BMDL (#1788)
- HBM data exclude substance sampling method combinations (#1795)
- Show number of non-analysed samples in hbm data samples summary (#1797)

12.25.2 Changed

- MCRA security: make 2FA compulsory for all MCRA accounts (#1765)
- · Always show top right help icon in MCRA web app

12.25.3 Fixed

- User groups not visible for non-admin MCRA users
- Remove strong password validation check on Lost QR code form

12.26 Version 10.0.2 (2023-10-26)

No changes in MCRA Web

12.27 Version 10.0.1 (2023-10-23)

12.27.1 Added

- Bootstrap hbm monitoring data (#1405)
- MCRA security: two-factor authentication (2FA) (#1582)
- Implement PARC-HBM data format reader to support codebook v2.3 (#1649)
- Basic implementation kinetic conversion factors (#1658, #1660)
- Create separate matrix selection settings for kinetic models and HBM analysis (#1684)
- Exclude substances from urine or blood normalisation standardisation (#1659, #1703)
- Implement kinetic conversion factors (#1695)
- Add biological matrix and expression type to points of departure (POD) data table (#1724)
- Hazard characterisations uncertain (#1742)

12.27.2 Changed

- Update HBM demo standard action on bisphenols to include Karrer kinetic model (#1251)
- Update frontend action details form to show progress spinner on setting/change data sources (#1672)
- Remove action tiers from module definitions to tier/template files per tier (#1673)
- Using new risk metric names for section paths in standard actions (#1679)
- · Allow multiple/nested HTML tab panels in output section views

12.27.3 Fixed

- Wrong thousands separator in number of Monte Carlo iterations (#1645)
- Import HBM data should read in correct biological matrix (#1685)
- Convert to single target for hazard characterisations (#1696)
- Substance-specific intraspecies conversion of HC does not work (#1710)
- Delete data folder and show data source usage (#1713)

12.28 Version 10.0.0 (2023-06-23)

Public release of Open MCRA Core source code on GitHub.

12.28.1 Changed

- Update short report template for EFSA 2022 acute cranio
- Adjust text on MCRA home page for MCRA 10 (#1627)
- Adjust documentation, settings and URIs for MCRA 10 (#1627)

12.28.2 Fixed

- Inconsistencies in setting visibility and selected module tiers, add documentation (#1632)
- Delete action unnecessarily loads settings (#1646)
- Username is not saved when creating an external repository
- Wrong tier selection in SA EU chronic exposure assessment (2018) and Training Prospective Chronic (#1650)

12.29 Version 9.2.10 (2023-06-01)

12.29.1 Changed

• Download loop task output (html, tables, charts) creates a zip file with all sub-task outputs (#1530)

12.29.2 Fixed

- Version number of data source in RPF action input summary shows wrong version (#1604)
- Show correct tier setting for single value risk in SA cranio efsa 2022 (#1635)

12.30 Version 9.2.9 (2023-05-24)

12.30.1 Added

- Implement support for missing bodyweights in MCRA (#1585)
- Documentation imputation with zero and biological matrix conversion (#1603)
- Add 'compare sub tasks' to single value risk module (#1606)

12.30.2 Changed

- Clean up the contents of the downloadable HTML zip report: move CSV, SVG and metadata files into subfolders (#1361)
- Update EFSA 2022 SRA cranio to use the new EFSA 2022 settings tiers (#1544)
- Update create action wizard of risks module (and single value risks) to include risk metric choice (#1574)
- Update the two standard actions for training prospective CRA with option to run scenario GAP + 20% use (#1621)

12.30.3 Fixed

- Data folders in Workspace data tab are not scrollable (#1572)
- Human monitoring data allows to run without specifying survey (#1593)
- Correct update method of automatic forms so that also the item visibilities are also properly updated after form submission (#1628)
12.31 Version 9.2.8 (2023-04-25)

12.31.1 Added

- Update HTML report in zip download with toc-functionality to make it easier to browse through (#1409)
- Risks: Implement cumulative as sum of single-substance ratios (Exposure/Hazard) (#1528)
- Add documentation core and web (#1448)

12.31.2 Fixed

- Data sources view is not updated/refreshed after upload new data source version (#1512)
- User name input check for reset password is not correct (#1594)
- Setting Multiple effects gives error (#1595)

12.32 Version 9.2.7 (2023-04-03)

12.32.1 Added

- Separate substance-weighing option for MCR and mixtures analysis and add checkbox to compute or not compute MCR in HBM analysis actions (#1499)
- Show 'Created' date in admin users panel (#1523)
- Add standardisation methods for blood and urine (#1359, #1477)
- Implement initial version of SRA acute CRA of craniofacial alterations EFSA 2022 (#1464)

12.32.2 Changed

- Remove MCRA version select page, MCRA 9 is now the default
- Update exposure mixtures module with option to log-transform data before network analysis (#1531)

12.32.3 Fixed

• Hide option for reference substance equivalents when cumulative does not apply for the action (#1502)

12.33 Version 9.2.6 (2023-03-10)

12.33.1 Added

- Add compartments including cumulative amounts like urine (#1272)
- Allow hbm as input for risks (#1394)
- Add documentation about allocation of substances (#1473)
- Add new model substance approvals (#1461)

12.33.2 Changed

- Remove service worker functionality from Angular app
- Change MCRA URLs to point to static Documentation URL (#889)
- Move and update documentation section on HBM4EU/PARC HBM data format (#1427)
- Update Angular version to v15.0 (#1468)
- Unique constraint of Hazard Characterisations table is too strict (#1491)

· Keep report name tabs visible when scrolling through report comparison view

12.33.3 Fixed

- File upload failed: max upload file size was not configured correctly
- · Fixed subset range editor
- Transaction deadlocked on lock resources in Job scheduler and simulation worker (#1487)
- Job scheduler does not assign jobs correctly (#1489)
- Null reference exception when changing data source (#1501)
- Exception when trying to view task settings in admin tasks panel
- · Report comparison fails to load
- Can't compare outputs of a loop task in tabbed output view, TOC doesn't work (#1509, #1510)
- Error messages are not displayed correctly in web application
- Missing user account when e-mail verification fails (#1520)

12.34 Version 9.2.5 (2023-02-10)

12.34.1 Added

• Documentation: added section on CLI (#1450)

12.34.2 Changed

• Update third party packages (NuGet) for MCRA Web

12.34.3 Fixed

• Failed dataset upload still creates a data source record in the repository (#1483)

12.35 Version 9.2.4 (2023-02-03)

12.35.1 Added

- Add documentation of HBM4EU/PARC HBM data format (#1427)
- Add maximum percentage missing value percentage imputation hbm data (#1470)
- Add population characteristics Real Life Mixtures (#1475)

12.35.2 Changed

- Update HBM analysis module to include imputation of missing data method established within HBM4EU/PARC RLM (#1397)
- Update HBM data/analysis module to allow for analysis of concentrations of multiple sampling methods/matrices (#1398)
- Implement changed rules for sending automatic mail from WUR and RIVM (#1466)
- · Move internal concentration type to assessment settings

12.35.3 Fixed

- Download action + data (data as zipped csv) fails (#1451)
- Loop task output doesn't show the comparison between the looped actions (#1459)
- New user registration in MCRA returns 400 Bad Request (#1465)
- Load correct png mime type for safety chart
- Upload of Demo HBM bisphenols standard action data fails (#1472)

12.36 Version 9.2.3 (2023-01-16)

12.36.1 Added

• Implement chlorpyrifos kinetic model with metabolites (#1285)

12.36.2 Changed

- Removed unit test report results from admin page (#1441)
- Allow links to MCRA documentation on other servers (#1457)

12.36.3 Fixed

- Documentation shows no version of MCRA in colophon
- · Documentation links to web API documentation and web application
- Can't view info in data browser of a shared data source (#1458)

12.37 Version 9.2.2 (2022-12-20)

12.37.1 Added

• Documentation: Add PARC Real-life mixture guidance documents (pdf) and related data to the User guide - Examples section (#1366)

12.37.2 Changed

• Update HBM analysis module with non-detects imputation method according to method established within HBM4EU/PARC RLM (#1396)

12.37.3 Fixed

- Fix reference to dietary exposures section in risks output section (#1331)
- MCRA Build date-time not displayed correctly (#1410)
- Settings in overview only show headers and no settings (#1442)
- Mixtures crashes when no concentrations for HBM data are available (#1443)
- Output collection of task with subtask fails (#1444)
- MCRA settings loading fails: allow floating point literals (NaN, Inf etc) in Json serialization

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INDEX

А

ADI, 959 ADME, 959 AF, 959 AIC, 959 AOP, 959 ARFD, 959 AU, 959

В

BBN, 959 BMD, 959 BMDL, 959 BMDU, 959 BMI, 959 BMR, 959 BOD, 959 BREAM, 960 BROWSE, 960 BW, 960

С

CA, 960 CAG, 960 ConsExpo, 960 CRA, 960

D

DA, 960 DALY, 960 DR, 960

Ε

E/H, 960 EBD, 960 EEA, 960 ERF, 960

F

fa, **960** fc, **960**

G

GAP, **960** GM, **961** GSD, 961

Η

H/E, 961 HBGV, 961 HBM, 961 HC, 961 HCPs, 961 HI, 961 HQ, 961 HR, 961

ICED, 961 In-silico, 961 In-vitro, 961 In-vivo, 961 IVIVE, 961

J

JRC, 961

L

Lipid, 962 LNN, 962 LOAEL, 962 LOD, 962 LOQ, 962 LOR, 962

Μ

MCR, 962 MCRA, 962 MIE, 962 MOA, 962 MOET, 962 MRA, 962 MRL, 962 MV, 962

Ν

NAMs, 962 NMF, 962 NOAEL, 963 NOEC, 963 NOEL, 963

Ο

OIM, 963 OP, 963

Ρ

PACEM, 963 PAF, 963 PARC, 963 PBK, 963 PBPK, 963 PCPs, 963 POCE, 963 POD, 963 PPP, 963 PRIMO, 963

Q

QSAR, 963

R

RA, 963 RAC, 964 RIVM, 964 RPC, 964 RPC derivatives, 964 RPF, 964 RR, 964 RSExpo, 964

S

SA, 964 SBML, 964 SD, 964 SG, 964 SNMU, 964 SRA, 964 SSC, 964 SSD, 964

Т

TDI, 964 TDS, 964 TEF, 964 TK, 965 TP, 965 TTC, 965 TWI, 965

Y

YLD, 965 YLL, 965