Package 'httk'

March 25, 2022

```
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Title High-Throughput Toxicokinetics
Description Generic models and chemical-specific data for simulation and
      statistical analysis of chemical toxicokinetics (`TK") as
      described by Pearce et al. (2017) <doi:10.18637/jss.v079.i04>.
      Chemical-specific in vitro data have been obtained from relatively
      high-throughput experiments. Both physiologically-based (``PBTK")
      and empirical (for example, one compartment) "TK" models can be
      parameterized with the data provided for thousands of chemicals,
      multiple exposure routes, and various species. The models consist
      of systems of ordinary differential equations which are solved
      using compiled (C-based) code for speed. A Monte Carlo sampler is
      included, which allows for simulating human biological variability
      (Ring et al., 2017 <doi:10.1016/j.envint.2017.06.004>)
      and propagating parameter uncertainty. Calibrated methods are
      included for predicting tissue:plasma partition coefficients and
      volume of distribution
      (Pearce et al., 2017 < doi:10.1007/s10928-017-9548-7>).
      These functions and data provide a set of tools for
      in vitro-in vivo extrapolation (``IVIVE") of high-throughput
      screening data (for example, Tox21, ToxCast) to real-world
      exposures via reverse dosimetry (also known as ``RTK")
      (Wetmore et al., 2015 <doi:10.1093/toxsci/kfv171>).
Depends R (>= 2.10)
Imports deSolve, msm, data.table, survey, mvtnorm, truncnorm, stats,
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RdMacros Rdpack
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      rmarkdown,
      R.rsp,
      GGally,
      gplots,
      scales,
      EnvStats,
      MASS,
      RColorBrewer,
```

Version 2.1.0

```
TeachingDemos,
       classInt.
       ks,
       stringr,
       reshape,
       reshape2,
       viridis,
       CensRegMod,
       gmodels,
       colorspace,
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       ggrepel,
       dplyr,
       forcats,
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       gridExtra,
       testthat
License GPL-3
LazyData true
LazyDataCompression xz
Encoding UTF-8
VignetteBuilder knitr, R.rsp
RoxygenNote 7.1.1
URL https:
       //www.epa.gov/chemical-research/rapid-chemical-exposure-and-dose-research
BugReports https://github.com/USEPA/CompTox-ExpoCast-httk
NeedsCompilation yes
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```

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# Description

Generic models and chemical-specific data for simulation and statistical analysis of chemical toxicokinetics ("TK") as described by Pearce et al. (2017) <doi:10.18637/jss.v079.i04>. Chemical-specific in vitro data have been obtained from relatively high-throughput experiments. Both physiologically-based ("PBTK") and empirical (for example, one compartment) "TK" models can be parameterized with the data provided for thousands of chemicals, multiple exposure routes, and various species. The models consist of systems of ordinary differential equations which are solved using compiled (C-based) code for speed. A Monte Carlo sampler is included, which allows for simulating human biological variability (Ring et al., 2017 <doi:10.1016/j.envint.2017.06.004>) and propagating parameter uncertainty. Calibrated methods are included for predicting tissue:plasma partition coefficients and volume of distribution (Pearce et al., 2017 <doi:10.1007/s10928-017-9548-7>). These functions and data provide a set of tools for in vitro-in vivo extrapolation ("IVIVE") of high-throughput screening data (for example, Tox21, ToxCast) to real-world exposures via reverse dosimetry (also known as "RTK") (Wetmore et al., 2015 <doi:10.1093/toxsci/kfv171>).

# Author(s)

John Wambaugh, Robert Pearce, Caroline Ring, Gregory Honda, Nisha Sipes, Jimena Davis, Barbara Wetmore, Woodrow Setzer, Mark Sfeir

# See Also

PowerPoint Presentation: High-Throughput Toxicokinetics (HTTK) R package

doi: 10.1080/17425255.2021.1935867Breen et al. (2021): High-throughput PBTK models for in vitro to in vivo extrapolation

doi: 10.18637/jss.v079.i04Pearce et al. (2017): httk: R Package for High-Throughput Toxicokinetics

doi: 10.1021/es501955gArmitage et al. (2014): Application of mass balance models and the chemical activity concept to facilitate the use of in vitro toxicity data for risk assessment

doi: 10.1093/toxsci/kfv171Incorporating High-Throughput Exposure Predictions with Dosimetry-Adjusted In Vitro Bioactivity to Inform Chemical Toxicity Testing

doi: 10.1093/toxsci/kfv118Wambaugh et al. (2015): Toxicokinetic Triage for Environmental Chemicals

doi: 10.1007/s1092801795487Pearce et al. (2017): Evaluation and calibration of high-throughput predictions of chemical distribution to tissues

doi: 10.1016/j.envint.2017.06.004Ring et al. (2017): Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability

doi: 10.1021/acs.est.7b00650Sipes et al. (2017): An Intuitive Approach for Predicting Potential Human Health Risk with the Tox21 10k Library

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doi: 10.1093/toxsci/kfy020Wambaugh et al. (2018): Evaluating In Vitro-In Vivo Extrapolation of Toxicokinetics

doi: 10.1371/journal.pone.0217564Honda et al. (2019): Using the concordance of in vitro and in vivo data to evaluate extrapolation assumptions

doi: 10.1093/toxsci/kfz205Wambaugh et al. (2019): Assessing Toxicokinetic Uncertainty and Variability in Risk Prioritization

doi: 10.1038/s413700200238yLinakis et al. (2020): Development and evaluation of a high-throughput inhalation model for organic chemicals

The U.S. EPA ExpoCast (Exposure Forecasting) Project

add_chemtable

Add a table of chemical information for use in making httk predictions.

## **Description**

This function adds chemical-specific information to the table chem.physical_and_invitro.data. This table is queried by the model parameterization functions when attempting to parameterize a model, so adding sufficient data to this table allows additional chemicals to be modeled.

# Usage

```
add_chemtable(
  new.table,
  data.list,
  current.table = NULL,
  reference = NULL,
  species = NULL,
  overwrite = FALSE,
  sig.fig = 4,
  clint.pvalue.overwrite = TRUE,
  allow.na = FALSE
)
```

### **Arguments**

new. table Object of class data.frame containing one row per chemical, with each chemical

minimally described by a CAS number.

data.list This list identifies which properties are to be read from the table. Each item

in the list should point to a column in the table new.table. Valid names in the list are: 'Compound', 'CAS', 'DSSTox.GSID' 'SMILES.desalt', 'Reference', 'Species', 'MW', 'logP', 'pKa_Donor', 'pKa_Accept', 'logMA', 'Clint',

'Clint.pValue', 'Funbound.plasma', 'Fgutabs', 'Rblood2plasma'.

current. table This is the table to which data are being added.

reference This is the reference for the data in the new table. This may be omitted if a

column in data.list gives the reference value for each chemical.

species This is the species for the data in the new table. This may be omitted if a column

in data. list gives the species value for each chemical or if the data are not species-

specific (e.g., MW).

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overwrite

If overwrite=TRUE then data in current.table will be replaced by any data in new.table that is for the same chemical and property. If overwrite=FALSE (DE-FAULT) then new data for the same chemical and property are ignored. Funbound.plasma values of 0 (below limit of detection) are overwritten either way.

sig.fig Sets the number of significant figures stored (defaults to 4)

clint.pvalue.overwrite

If TRUE then the Cl_int p-value is set to NA when the Cl_int value is changed unless a new p-value is provided. (defaults to TRUE)

allow.na

If TRUE (default is FALSE) then NA values are written to the table, otherwise they are ignored.

#### Value

data.frame

A new data.frame containing the data in current.table augmented by new.table

#### Author(s)

John Wambaugh

# **Examples**

```
library(httk)
my.new.data <- as.data.frame(c("A","B","C"),stringsAsFactors=FALSE)</pre>
my.new.data <- cbind(my.new.data,as.data.frame(c(</pre>
                      "111-11-2", "222-22-0", "333-33-5"),
                      stringsAsFactors=FALSE))
my.new.data <- cbind(my.new.data,as.data.frame(c("DTX1","DTX2","DTX3"),</pre>
                     stringsAsFactors=FALSE))
my.new.data <- cbind(my.new.data,as.data.frame(c(200,200,200)))</pre>
my.new.data <- cbind(my.new.data,as.data.frame(c(2,3,4)))</pre>
my.new.data <- cbind(my.new.data,as.data.frame(c(0.01,0.02,0.3)))
my.new.data \leftarrow cbind(my.new.data,as.data.frame(c(0,10,100)))
colnames(my.new.data) <- c("Name","CASRN","DTXSID","MW","LogP","Fup","CLint")</pre>
chem.physical_and_invitro.data <- add_chemtable(my.new.data,</pre>
                                    current.table=
                                      chem.physical_and_invitro.data,
                                    data.list=list(
                                    Compound="Name",
                                    CAS="CASRN",
                                    DTXSID="DTXSID",
                                    MW="MW",
                                    logP="LogP",
                                    Funbound.plasma="Fup",
                                    Clint="CLint"),
                                    species="Human",
                                    reference="MyPaper 2015")
parameterize\_steady state (\texttt{chem.name="C"})
calc_css(chem.name="B")
# Initialize a column describing proton donors ("acids")
my.new.data$pka.a <- NA
# set chemical C to an acid (pKa_donor = 5):
my.new.data[my.new.data$Name=="C","pka.a"] <- "5"</pre>
```

age_dist_smooth 9

```
chem.physical_and_invitro.data <- add_chemtable(my.new.data,</pre>
                                   current.table=
                                     chem.physical_and_invitro.data,
                                  data.list=list(
                                  Compound="Name",
                                  CAS="CASRN",
                                  DTXSID="DTXSID".
                                  pKa_Donor="pka.a"),
                                  species="Human",
                                  reference="MyPaper 2015")
# Note Rblood2plasma and hepatic bioavailability change (relative to above):
parameterize_steadystate(chem.name="C")
# Initialize a column describing proton acceptors ("bases")
my.new.data$pka.b <- NA
# set chemical B to a base with multiple pka's (pKa_accept = 7 and 8):
my.new.data[my.new.data$Name=="B","pka.b"] <- "7;8"</pre>
chem.physical_and_invitro.data <- add_chemtable(my.new.data,</pre>
                                   current.table=
                                     chem.physical_and_invitro.data,
                                  data.list=list(
                                  Compound="Name"
                                  CAS="CASRN",
                                  DTXSID="DTXSID"
                                  pKa_Accept="pka.b"),
                                  species="Human",
                                  reference="MyPaper 2015")
# Note that average and max change (relative to above):
calc_css(chem.name="B")
```

age_dist_smooth

Smoothed age distributions by race and gender.

# **Description**

Distributions of ages in months, computed from NHANES data smoothed using survey::svysmooth(), for each combination of race/ethnicity and gender.

#### **Usage**

```
age_dist_smooth
```

# **Format**

A data.table object with three variables:

```
gender Gender: Male or Female
reth Race/ethnicity
smth A list of svysmooth objects, each encoding a weighted smoothed distribution of ages.
```

# Author(s)

Caroline Ring

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## References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

age_draw_smooth	Draws ages from a smoothed distribution for a given gender/race combination

# Description

Draws ages from a smoothed distribution for a given gender/race combination

# Usage

```
age_draw_smooth(g, r, nsamp, agelim_months)
```

# **Arguments**

σ	Gender	Fither	'Male'	or	'Female'.
8	Gender.	Eluiei	wate	OI	remaie.

r Race/ethnicity. One of 'Mexican American', 'Other Hispanic', 'Non-Hispanic

Black', 'Non-Hispanic White', 'Other'.

nsamp Number of ages to draw.

agelim_months Two-element numeric vector giving the minimum and maximum ages in months

to include.

# Value

A named list with members 'ages_months' and 'ages_years', each numeric of length nsamp, giving the sampled ages in months and years.

# Author(s)

Caroline Ring

## References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

```
armitage_estimate_sarea
```

Estimate well surface area

# **Description**

Estimate geometry surface area of plastic in well plate based on well plate format suggested values from Corning. option.plastic == TRUE (default) give nonzero surface area (sarea,  $m^2$ ) option.bottom == TRUE (default) includes surface area of the bottom of the well in determining sarea. Optionally include user values for working volume (v_working,  $m^3$ ) and surface area.

## Usage

```
armitage_estimate_sarea(
  tcdata = NA,
  this.well_number = 384,
  this.cell_yield = NA,
  this.v_working = NA
)
```

# **Arguments**

A data table with well_number corresponding to plate format, optionally include v_working, sarea, option.bottom, and option.plastic

this.well_number
For single value, plate format default is 384, used if is.na(tcdata)==TRUE

this.cell_yield
For single value, optionally supply cell_yield, otherwise estimated based on well number

this.v_working
For single value, optionally supply working volume, otherwise estimated based

# Value

A data table composed of any input data.table *tcdata* with only the following columns either created or altered by this function:

on well number (m^3)

Column Name	Description	Units
well_number	number of wells on plate	
sarea	surface area	m^2
cell_yield	number of cells	cells
v_working	working (filled) volume of each well	uL
v_total	total volume of each well	uL

# Author(s)

Greg Honda

#### References

Armitage, J. M., Arnot, J. A., Wania, F., & Mackay, D. (2013). Development and evaluation of a mechanistic bioconcentration model for ionogenic organic chemicals in fish. Environmental toxicology and chemistry, 32(1), 115-128.

armitage_eval

Evaluate the updated Armitage model

# **Description**

Evaluate the Armitage model for chemical distribution in vitro. Takes input as data table or vectors of values. Outputs a data table. Updates over the model published in Armitage et al. 2014 include binding to plastic walls and lipid and protein compartments in cells.

# Usage

```
armitage_eval(
  casrn.vector = NA_character_,
 nomconc.vector = 1,
  this.well_number = 384,
  this.FBSf = NA_real_,
  tcdata = NA,
  this.sarea = NA_real_,
  this.v_total = NA_real_,
  this.v_working = NA_real_,
  this.cell_yield = NA_real_,
  this. Tsys = 37,
  this. Tref = 298.15,
  this.option.kbsa2 = FALSE,
  this.option.swat2 = FALSE,
  this.pseudooct = 0.01,
  this.memblip = 0.04,
  this.nlom = 0.2,
  this.P_nlom = 0.035,
  this.P_{dom} = 0.05,
  this.P_cells = 1,
  this.csalt = 0.15,
  this.celldensity = 1,
  this.cellmass = 3,
  this.f_oc = 1,
  this.conc_ser_alb = 24,
  this.conc_ser_lip = 1.9,
  this.Vdom = 0
)
```

# Arguments

```
casrn.vector For vector or single value, CAS number nomconc.vector For vector or single value, micromolar nominal concentration (e.g. AC50 value)
```

this.well_numbe	er
	For single value, plate format default is 384, used if is.na(tcdata)==TRUE
this.FBSf	Fraction fetal bovine serum, must be entered by user.
tcdata	A data.table with casrn, nomconc, MP, gkow, gkaw, gswat, sarea, v_total, v_working. Otherwise supply single values to this.params.
this.sarea	Surface area per well (m^2)
this.v_total	Total volume per well (m^3)
this.v_working	Working volume per well (m^3)
this.cell_yield	
	Number of cells per well
this.Tsys	System temperature (degrees C)
this.Tref	Reference temperature (degrees K)
this.option.kbs	
*6:-	Use alternative bovine-serum-albumin partitioning model
this.option.swa	Use alternative water solubility correction
this pseudooct	Pseudo-octanol cell storage lipid content
this.memblip	Membrane lipid content of cells
this.membiip	•
	Structural protein conent of cells
this.P_nlom	Proportionality constant to octanol structural protein
this.P_dom	Proportionality constant to dissolve organic material
this.P_cells	Proportionality constant to octanol storage lipid
this.csalt	Ionic strength of buffer, mol/L
this.celldensi	
464 11	Cell density kg/L, g/mL
this.cellmass	Mass per cell, ng/cell
this.f_oc	1, everything assumed to be like proteins
this.conc_ser_a	24 g/L, mass concentration of albumin in serum.
this.conc_ser_]	
	1.9 g/L, mass concentration of lipids in serum.
this.Vdom	0 ml, the volume of dissolved organic matter (DOM)
	, , ,

# Value

Column casrn	<b>Description</b> Chemical Abstracts Service Registry Number	units
nomconc	Nominal Concentration	mol/L
well_number	Number of wells in plate	unitless
sarea	Surface area of well	m^2
v_total	Total volume of well	m^3
v_working	Filled volume of well	m^3
cell_yield	Number of cells	cells
gkow	log10 octanol to water partition coefficient (PC)	log10
logHenry	log10 Henry's law constant '	log10 atm-m3/mol
gswat	log10 Water solubility	log10 mol/L

MP	Malting Point	degrees Celsius
	Melting Point	_
MW	Molecular Weight	g/mol
gkaw	air-water partition coefficient	(mol/m3)/(mol/m3)
dsm		
duow		
duaw		
dumw		
gkmw		
gkcw		
gkbsa		
gkpl		
ksalt		
Tsys		
Tref		
option.kbsa2		
option.swat2		
FBSf		
pseudooct		
memblip		
nlom		
P_nlom		
P_dom	dissolved organic matter to water PC	Dimensionless
	dissolved organic matter to water FC	Difficusioniess
P_cells		
csalt		
celldensity		
cellmass		
f_oc		
cellwat		
Tcor		
Vm	Volume of media	L
Vwell	volume of medium (aqueous phase only)	L
Vair	volume of head space	L
Vcells	volume of cells/tissue	
Valb	volume of serum albumin	
Vslip	volume of serum lipids	
Vdom	volume of dissolved organic matter	
F_ratio		
gs1.GSE		
s1.GSE		
gss.GSE		
ss.GSE		
kmw		
kow	octanol to water PC	
kaw	the air towater PC	dimensionless
swat		
kpl		
kew	cell/tissue to water PC	dimensionless
kbsa	The second to make I o	
swat_L		
oct_L		
scell_L		
SCEII_L	Total concentration	1

mol

cinit

Initial concentration

mtot	Total moles	mol
cwat	Total concentration in water	mol/L
cwat_s	Dissolved concentration in water	mol/L
csat	Is the solution saturated (1/0)	Boolean
activity		
cair		mol/L
calb		mol/L
cslip		mol/L
cdom	concentration of/in dissolved organic matter	mol/L
ccells		mol/L
cplastic		mol/L
mwat_s	Mass dissolved in water	mols
mair	Mass in air	mols
mbsa	Mass bound to bovine serum albumin	mols
mslip	Mass bound to serum lipids	mols
mdom	Mass bound to dissolved organic matter	mols
mcells	Mass in cells	mols
mplastic	Mass bond to plastic	mols
mprecip	Mass precipitated out of solution	
xwat_s	Fraction dissolved in water	fraction
xair	Fraction in the air	fraction
xbsa	Fraction bound to bovine serum albumin	fraction
xslip	Fraction bound to serum lipids	fraction
xdom	Fraction bound to dissolved organic matter	fraction
xcells	Fraction within cells	fraction
xplastic	Fraction bound to plastic	fraction
xprecip	Fraction precipitated out of solution	fraction
eta_free	effective availability ratio	fraction
cfree.invitro	Free concentration in the in vitro media (use for Honda1 and Honda2)	micromolar

# Author(s)

Greg Honda

# References

Armitage, J. M.; Wania, F.; Arnot, J. A. Environ. Sci. Technol. 2014, 48, 9770-9779. https://doi.org/10.1021/es501955g Honda et al. PloS one 14.5 (2019): e0217564. https://doi.org/10.1371/journal.pone.0217564

# **Examples**

```
library(httk)
# Check to see if we have info on the chemical:
"80-05-7" %in% get_cheminfo()

#We do:
temp <- armitage_eval(casrn.vector = c("80-05-7", "81-81-2"), this.FBSf = 0.1,
this.well_number = 384, nomconc = 10)</pre>
```

16 armitage_input

```
print(temp$cfree.invitro)
# Check to see if we have info on the chemical:
"793-24-8" %in% get_cheminfo()
# Since we don't have any info, let's look up phys-chem from dashboard:
cheminfo <- data.frame(</pre>
  Compound="6-PPD",
  CASRN="793-24-8",
  DTXSID="DTXSID9025114",
  logP=4.27,
  logHenry=log10(7.69e-8),
  logWSol=log10(1.58e-4),
  MP = 99.4,
  MW=268.404
  )
# Add the information to HTTK's database:
chem.physical_and_invitro.data <- add_chemtable(</pre>
 cheminfo,
 current.table=chem.physical_and_invitro.data,
 data.list=list(
 Compound="Compound",
 CAS="CASRN",
 DTXSID="DTXSID",
  MW="MW",
  logP="logP",
  logHenry="logHenry",
  logWSol="logWSol",
  MP="MP"),
  species="Human",
  reference="CompTox Dashboard 31921")
# Run the Armitage et al. (2014) model:
out <- armitage_eval(</pre>
  casrn.vector = "793-24-8",
  this.FBSf = 0.1,
  this.well_number = 384,
  nomconc = 10)
print(out)
```

armitage_input

Armitage et al. (2014) Model Inputs from Honda et al. (2019)

# **Description**

Armitage et al. (2014) Model Inputs from Honda et al. (2019)

```
armitage_input
```

augment.table 17

## **Format**

A data frame with 53940 rows and 10 variables:

MP MW

casrn

compound_name

gkaw gkow

gswat

# Author(s)

Greg Honda

#### Source

https://www.diamondse.info/

# References

Armitage, J. M.; Wania, F.; Arnot, J. A. Environ. Sci. Technol. 2014, 48, 9770-9779. dx.doi.org/10.1021/es501955g Honda, Gregory S., et al. "Using the Concordance of In Vitro and In Vivo Data to Evaluate Extrapolation Assumptions", PloS ONE 14.5 (2019): e0217564.

augment.table

Add a parameter value to the chem.physical_and_invitro.data table

# **Description**

This internal function is used by add_chemtable to add a single new parameter to the table of chemical parameters. It should not be typically used from the command line.

```
augment.table(
  this.table,
  this.CAS,
  compound.name = NULL,
  this.property,
  value,
  species = NULL,
  reference,
  overwrite = FALSE,
  sig.fig = 4,
  clint.pvalue.overwrite = TRUE,
  allow.na = FALSE
)
```

#### **Arguments**

this.table Object of class data.frame containing one row per chemical.

this.CAS The Chemical Abstracts Service registry number (CAS-RN) correponding to the

parameter value

compound.name A name associated with the chemical (defaults to NULL)

this.property The property being added/modified.

value The value being assigned to this.property.

species This is the species for the data in the new table. This may be omitted if a column

in data.list gives the species value for each chemical or if the data are not species-

specific (e.g., MW).

reference This is the reference for the data in the new table. This may be omitted if a

column in data.list gives the reference value for each chemical.

overwrite If overwrite=TRUE then data in current.table will be replaced by any data in

new.table that is for the same chemical and property. If overwrite=FALSE (DE-FAULT) then new data for the same chemical and property are ignored. Funbound.plasma values of 0 (below limit of detection) are overwritten either way.

sig.fig Sets the number of significant figures stored (defaults to 4)

clint.pvalue.overwrite

If TRUE then the Cl_int p-value is set to NA when the Cl_int value is changed

unless a new p-value is provided. (defaults to TRUE)

allow.na If TRUE (default is FALSE) then NA values are written to the table, otherwise

they are ignored.

# Value

data.frame A new data.frame containing the data in current.table augmented by new.table

## Author(s)

John Wambaugh

available_rblood2plasma

Find the best available ratio of the blood to plasma concentration constant.

# Description

This function finds the best available constant ratio of the blood concentration to the plasma concentration, using get_rblood2plasma and calc_rblood2plasma.

```
available_rblood2plasma(
  chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
  species = "Human",
  adjusted.Funbound.plasma = TRUE,
  suppress.messages = FALSE
)
```

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# **Arguments**

chem. cas Either the CAS number or the chemical name must be specified. chem. name Either the chemical name or the CAS number must be specified.

dtxsid EPA's 'DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chemical

must be identified by either CAS, name, or DTXSIDs

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

adjusted.Funbound.plasma

Whether or not to use Funbound.plasma adjustment if calculating Rblood2plasma. suppress.messages

Whether or not to display relevant warning messages to user.

#### **Details**

Either retrieves a measured blood:plasma concentration ratio from the chem.physical_and_invitro.data table or calculates it using the red blood cell partition coefficient predicted with Schmitt's method

If available, in vivo data (from chem.physical_and_invitro.data) for the given species is returned, substituting the human in vivo value when missing for other species. In the absence of in vivo data, the value is calculated with calc_rblood2plasma for the given species. If Funbound.plasma is unvailable for the given species, the human Funbound.plasma is substituted. If none of these are available, the mean human Rblood2plasma from chem.physical_and_invitro.data is returned. details than the description above ~~

#### Value

The blood to plasma chemical concentration ratio - measured if available, calculated if not.

### Author(s)

Robert Pearce

# **Examples**

```
available_rblood2plasma(chem.name="Bisphenol A",adjusted.Funbound.plasma=FALSE)
available_rblood2plasma(chem.name="Bisphenol A",species="Rat")
```

aylward2014

Aylward et al. 2014

# Description

Aylward et al. (2014) compiled measurements of the ratio of maternal to fetal cord blood chemical concentrations at birth for a range of chemicals with environmental routes of exposure, including bromodiphenyl ethers, fluorinated compounds, organochlorine pesticides, polyaromatic hydrocarbons, tobacco smoke components, and vitamins.

# Usage

aylward2014

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#### **Format**

data.frame

#### Source

Kapraun et al. 2021 (submitted)

#### References

Aylward LL, Hays SM, Kirman CR, Marchitti SA, Kenneke JF, English C, Mattison DR, Becker RA (2014). "Relationships of chemical concentrations in maternal and cord blood: a review of available data." *Journal of Toxicology and Environmental Health, Part B*, **17**(3), 175–203. doi: 10.1080/10937404.2014.884956.

blood_mass_correct

Find average blood masses by age.

# **Description**

If blood mass from blood_weight is negative or very small, then just default to the mean blood mass by age. (Geigy Scientific Tables, 7th ed.)

# Usage

blood_mass_correct(blood_mass, age_months, age_years, gender, weight)

# **Arguments**

blood_mass A vector of blood masses in kg to be replaced with averages.

age_months A vector of ages in months.
age_years A vector of ages in years.

gender A vector of genders (either 'Male' or 'Female').

weight A vector of body weights in kg.

# Value

A vector of blood masses in kg.

# Author(s)

Caroline Ring

# References

Geigy Pharmaceuticals, "Scientific Tables", 7th Edition, John Wiley and Sons (1970)

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

blood_weight 21

# **Description**

Predict blood mass based on body surface area and gender, using equations from Bosgra et al. 2012

## Usage

```
blood_weight(BSA, gender)
```

# **Arguments**

BSA Body surface area in m^2. May be a vector. gender Either 'Male' or 'Female'. May be a vector.

#### Value

A vector of blood masses in kg the same length as BSA and gender.

## Author(s)

Caroline Ring

## References

Bosgra, Sieto, et al. "An improved model to predict physiologically based model parameters and their inter-individual variability from anthropometry." Critical reviews in toxicology 42.9 (2012): 751-767.

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

bmiage

CDC BMI-for-age charts

# Description

Charts giving the BMI-for-age percentiles for boys and girls ages 2-18

# Usage

bmiage

# **Format**

A data.table object with variables

Sex 'Male' or 'Female' Agemos Age in months

 $\textbf{L,M,S} \ \, \textbf{LMS} \ \, \textbf{parameters; see https://www.cdc.gov/growthcharts/percentile_data_files.}$ 

P3, P5, P10, P25, P50, P75, P85, P90, P95, and P97 BMI percentiles

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## **Details**

For children ages 2 to 18, weight class depends on the BMI-for-age percentile.

**Underweight** <5th percentile

Normal weight 5th-85th percentile

Overweight 85th-95th percentile

**Obese** >=95th percentile

## Author(s)

Caroline Ring

## Source

https://www.cdc.gov/growthcharts/percentile_data_files.htm

## References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

body_surface_area

Predict body surface area.

# Description

Predict body surface area from weight, height, and age, using Mosteller's formula for age>18 and Haycock's formula for age<18

# Usage

```
body_surface_area(BW, H, age_years)
```

# Arguments

BW A vector of body weights in kg.

H A vector of heights in cm.

age_years A vector of ages in years.

# Value

A vector of body surface areas in cm².

# Author(s)

Caroline Ring

bone_mass_age 23

#### References

Mosteller, R. D. "Simplified calculation of body surface area." N Engl J Med 317 (1987): 1098...

Haycock, George B., George J. Schwartz, and David H. Wisotsky. "Geometric method for measuring body surface area: a height-weight formula validated in infants, children, and adults." The Journal of pediatrics 93.1 (1978): 62-66.

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

bone_mass_age

Predict bone mass

# **Description**

Predict bone mass from age_years, height, weight, gender, using logistic equations fit to data from Baxter-Jones et al. 2011, or for infants < 1 year, using equation from Koo et al. 2000 (See Price et al. 2003)

#### Usage

bone_mass_age(age_years, age_months, height, weight, gender)

# **Arguments**

age_years Vector of ages in years.
age_months Vector of ages in months.
height Vector of heights in cm.
weight Vector of body weights in kg.

gender Vector of genders, either 'Male' or 'Female'.

### Value

Vector of bone masses.

#### Author(s)

Caroline Ring

# References

Baxter-Jones, Adam DG, et al. "Bone mineral accrual from 8 to 30 years of age: an estimation of peak bone mass." Journal of Bone and Mineral Research 26.8 (2011): 1729-1739.

Koo, Winston WK, and Elaine M. Hockman. "Physiologic predictors of lumbar spine bone mass in neonates." Pediatric research 48.4 (2000): 485-489.

Price, Paul S., et al. "Modeling interindividual variation in physiological factors used in PBPK models of humans." Critical reviews in toxicology 33.5 (2003): 469-503.

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

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brain_mass

Predict brain mass.

# **Description**

Predict brain mass from gender and age.

## Usage

```
brain_mass(gender, age_years)
```

# **Arguments**

gender Vector of genders, either 'Male' or 'Female' age_years Vector of ages in years.

# Value

A vector of brain masses in kg.

#### Author(s)

Caroline Ring

## References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

calc_analytic_css

Calculate the analytic steady state plasma concentration.

# **Description**

This function calculates the analytic steady state plasma or venous blood concentrations as a result of infusion dosing for the three compartment and multiple compartment PBTK models.

```
calc_analytic_css(
  chem.name = NULL,
  chem.cas = NULL,
  dtxsid = NULL,
  parameters = NULL,
  species = "human",
  daily.dose = 1,
  route = "oral",
  exp.conc = 1,
  period = 24,
  exp.duration = 24,
```

calc_analytic_css 25

```
output.units = "uM",
 model = "pbtk",
  concentration = "plasma".
  suppress.messages = FALSE,
  tissue = NULL,
  restrictive.clearance = TRUE,
 bioactive.free.invivo = FALSE,
  IVIVE = NULL,
 parameterize.args = list(default.to.human = FALSE, adjusted.Funbound.plasma = TRUE,
    regression = TRUE, minimum.Funbound.plasma = 1e-04),
)
```

#### **Arguments**

Either the chemical name, CAS number, or the parameters must be specified. chem.name Either the chemical name, CAS number, or the parameters must be specified. chem.cas dtxsid EPA's DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chemical must be identified by either CAS, name, or DTXSIDs Chemical parameters from parameterize_pbtk (for model = 'pbtk'), parameparameters terize_3comp (for model = '3compartment), parameterize_1comp(for model = '1compartment') or parameterize_steadystate (for model = '3compartmentss'), overrides chem.name and chem.cas. Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human"). species Total daily dose, mg/kg BW. daily.dose Route of exposure (either "oral", "iv", or "inhalation" default "oral"). route exp.conc Specified inhalation exposure concentration for use in assembling 'forcings' data series argument for integrator. Defaults to uM/L For use in assembling forcing function data series 'forcings' argument, specified period in hours For use in assembling forcing function data series 'forcings' argument, specified exp.duration in hours output.units Units for returned concentrations, defaults to uM (specify units = "uM") but can

also be mg/L. model

Model used in calculation, 'gas_pbtk' for the gas pbtk model, 'pbtk' for the multiple compartment model, '3compartment' for the three compartment model, '3compartmentss' for the three compartment steady state model, and '1compartment' for one compartment model.

Desired concentration type, 'blood', 'tissue', or default 'plasma'. concentration suppress.messages

Whether or not the output message is suppressed.

Desired tissue concentration (defaults to whole body concentration.) tissue restrictive.clearance

If TRUE (default), then only the fraction of chemical not bound to protein is available for metabolism in the liver. If FALSE, then all chemical in the liver is metabolized (faster metabolism due to rapid off-binding).

bioactive.free.invivo

If FALSE (default), then the total concentration is treated as bioactive in vivo. If TRUE, the the unbound (free) plasma concentration is treated as bioactive in vivo. Only works with tissue = NULL in current implementation.

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**IVIVE** 

Honda et al. (2019) identified four plausible sets of assumptions for *in vitro-in vivo* extrapolation (IVIVE) assumptions. Argument may be set to "Honda1" through "Honda4". If used, this function overwrites the tissue, restrictive.clearance, and bioactive.free.invivo arguments. See Details below for more information.

#### parameterize.args

List of arguments passed to model's associated parameterization function, including default.to.human, adjusted.Funbound.plasma, regression, and minimum.Funbound.plasma. The default.to.human argument substitutes missing animal values with human values if true, adjusted.Funbound.plasma returns adjusted Funbound.plasma when set to TRUE along with parition coefficients calculated with this value, regression indicates whether or not to use the regressions in calculating partition coefficients, and minimum.Funbound.plasma is the value to which Monte Carlo draws less than this value are set (default is 0.0001 – half the lowest measured Fup in our dataset).

Additional parameters passed to parameterize function if parameters is NULL.

#### **Details**

Concentrations are calculated for the specifed model with constant oral infusion dosing. All tissues other than gut, liver, and lung are the product of the steady state plasma concentration and the tissue to plasma partition coefficient.

	in vivo Conc.	Metabolic Clearance	Bioactive Chemical Conc.	TK Statistic Used*
Honda1	Veinous (Plasma)	Restrictive	Free	Mean Conc.
Honda2	Veinous	Restrictive	Free	Max Conc.
Honda3	Veinous	Non-restrictive	Total	Mean Conc.
Honda4	Veinous	Non-restrictive	Total	Max Conc.
Honda5	Target Tissue	Non-restrictive	Total	Mean Conc.
Honda6	Target Tissue	Non-restrictive	Total	Max Conc.

^{*}Assumption is currently ignored because analytical steady-state solutions are currently used by this function.

#### Value

Steady state plasma concentration in specified units

# Author(s)

Robert Pearce, John Wambaugh, Greg Honda, Miyuki Breen

# References

Honda, Gregory S., et al. "Using the Concordance of In Vitro and In Vivo Data to Evaluate Extrapolation Assumptions." 2019. PLoS ONE 14(5): e0217564.

# Examples

calc_analytic_css_1comp

Calculate the analytic steady state concentration for the one compartment model.

# **Description**

This function calculates the analytic steady state plasma or venous blood concentrations as a result of infusion dosing.

# Usage

```
calc_analytic_css_1comp(
  chem.name = NULL,
  chem.cas = NULL,
  dtxsid = NULL,
  parameters = NULL,
  hourly.dose = 1/24,
  concentration = "plasma",
  suppress.messages = FALSE,
  recalc.blood2plasma = FALSE,
  tissue = NULL,
  restrictive.clearance = TRUE,
  bioactive.free.invivo = FALSE,
  ...
)
```

# Arguments

chem. name

Either the chemical name, CAS number, or the parameters must be specified.

Either the chemical name, CAS number, or the parameters must be specified.

Either the chemical name, CAS number, or the parameters must be specified.

EPA's 'DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chemical must be identified by either CAS, name, or DTXSIDs

Chemical parameters from parameterize_pbtk (for model = 'pbtk'), parameterize_3comp (for model = '3compartment), parameterize_1comp(for model = '1compartment') or parameterize_steadystate (for model = '3compartmentss'), overrides chem.name and chem.cas.

```
hourly.dose Hourly dose rate mg/kg BW/h.

concentration Desired concentration type, 'blood' or default 'plasma'.

suppress.messages
```

Whether or not the output message is suppressed.

recalc.blood2plasma

Recalculates the ratio of the amount of chemical in the blood to plasma using the input parameters. Use this if you have altered hematocrit, Funbound.plasma, or Krbc2pu.

tissue Desired tissue conentration (defaults to whole body concentration.) restrictive.clearance

If TRUE (default), then only the fraction of chemical not bound to protein is available for metabolism in the liver. If FALSE, then all chemical in the liver is metabolized (faster metabolism due to rapid off-binding).

bioactive.free.invivo

If FALSE (default), then the total concentration is treated as bioactive in vivo. If TRUE, the the unbound (free) plasma concentration is treated as bioactive in vivo. Only works with tissue = NULL in current implementation.

Additional parameters passed to parameterize function if parameters is NULL.

## Value

Steady state plasma concentration in mg/L units

# Author(s)

Robert Pearce and John Wambaugh

```
calc_analytic_css_3comp
```

Calculate the analytic steady state concentration for model 3comp

# Description

This function calculates the analytic steady state plasma or venous blood concentrations as a result of infusion dosing.

```
calc_analytic_css_3comp(
  chem.name = NULL,
  chem.cas = NULL,
  dtxsid = NULL,
  parameters = NULL,
  hourly.dose = 1/24,
  concentration = "plasma",
  suppress.messages = FALSE,
  recalc.blood2plasma = FALSE,
  tissue = NULL,
  restrictive.clearance = TRUE,
  bioactive.free.invivo = FALSE,
  ...
)
```

# **Arguments**

chem. name Either the chemical name, CAS number, or the parameters must be specified.

chem. cas Either the chemical name, CAS number, or the parameters must be specified.

dtxsid EPA's 'DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chem-

ical must be identified by either CAS, name, or DTXSIDs

parameters Chemical parameters from parameterize_pbtk (for model = 'pbtk'), parame-

terize_3comp (for model = '3compartment), parameterize_1comp(for model = '1compartment') or parameterize_steadystate (for model = '3compartmentss'),

overrides chem.name and chem.cas.

hourly.dose Hourly dose rate mg/kg BW/h.

concentration Desired concentration type, 'blood' or default 'plasma'.

suppress.messages

Whether or not the output message is suppressed.

recalc.blood2plasma

Recalculates the ratio of the amount of chemical in the blood to plasma using the input parameters. Use this if you have 'altered hematocrit, Funbound.plasma, or

Krbc2pu.

tissue Desired tissue conentration (defaults to whole body concentration.)

restrictive.clearance

If TRUE (default), then only the fraction of chemical not bound to protein is available for metabolism in the liver. If FALSE, then all chemical in the liver is

metabolized (faster metabolism due to rapid off-binding).

bioactive.free.invivo

If FALSE (default), then the total concentration is treated as bioactive in vivo. If TRUE, the the unbound (free) plasma concentration is treated as bioactive in vivo. Only weeks with this case. All this correct implementation.

vivo. Only works with tissue = NULL in current implementation.

... Additional parameters passed to parameterize function if parameters is NULL.

### Value

Steady state plasma concentration in mg/L units

#### Author(s)

Robert Pearce and John Wambaugh

calc_analytic_css_3compss

Calculate the analytic steady state concentration for the three compartment steady-state model

# **Description**

This function calculates the analytic steady state plasma or venous blood concentrations as a result of infusion dosing.

#### Usage

```
calc_analytic_css_3compss(
  chem.name = NULL,
  chem.cas = NULL,
  dtxsid = NULL,
  parameters = NULL,
  hourly.dose = 1/24,
  concentration = "plasma",
  suppress.messages = FALSE,
  recalc.blood2plasma = FALSE,
  tissue = NULL,
  restrictive.clearance = TRUE,
  bioactive.free.invivo = FALSE,
  ...
)
```

# **Arguments**

chem.name

Either the chemical name, CAS number, or the parameters must be specified.

chem. cas Either the chemical name, CAS number, or the parameters must be specified.

dtxsid EPA's 'DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chem-

ical must be identified by either CAS, name, or DTXSIDs

parameters Chemical parameters from parameterize_pbtk (for model = 'pbtk'), parame-

terize_3comp (for model = '3compartment), parameterize_1comp(for model = '1compartment') or parameterize_steadystate (for model = '3compartmentss'),

overrides chem.name and chem.cas.

hourly.dose Hourly dose rate mg/kg BW/h.

concentration Desired concentration type, 'blood' or default 'plasma'.

suppress.messages

Whether or not the output message is suppressed.

recalc.blood2plasma

Recalculates the ratio of the amount of chemical in the blood to plasma using the input parameters. Use this if you have 'altered hematocrit, Funbound.plasma, or

Krbc2pu.

tissue Desired tissue concentration (defaults to whole body concentration.)

restrictive.clearance

If TRUE (default), then only the fraction of chemical not bound to protein is available for metabolism in the liver. If FALSE, then all chemical in the liver is metabolized (faster metabolism due to rapid off-binding).

bioactive.free.invivo

If FALSE (default), then the total concentration is treated as bioactive in vivo. If TRUE, the the unbound (free) plasma concentration is treated as bioactive in vivo. Only works with tissue = NULL in current implementation.

. . Additional parameters passed to parameterize function if parameters is NULL.

# Value

Steady state plasma concentration in mg/L units

calc_analytic_css_pbtk

## Author(s)

Robert Pearce and John Wambaugh

```
calc_analytic_css_pbtk
```

Calculate the analytic steady state plasma concentration for model pbtk.

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# Description

This function calculates the analytic steady state plasma or venous blood concentrations as a result of infusion dosing.

# Usage

```
calc_analytic_css_pbtk(
  chem.name = NULL,
  chem.cas = NULL,
  dtxsid = NULL,
  parameters = NULL,
  hourly.dose = 1/24,
  concentration = "plasma",
  suppress.messages = FALSE,
  recalc.blood2plasma = FALSE,
  tissue = NULL,
  restrictive.clearance = TRUE,
  bioactive.free.invivo = FALSE,
  ...
)
```

# **Arguments**

chem.name	Either the chemical name, CAS number, or the parameters must be specified.	
chem.cas	Either the chemical name, CAS number, or the parameters must be specified.	
dtxsid	EPA's 'DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chemical must be identified by either CAS, name, or DTXSIDs	
parameters	Chemical parameters from parameterize_pbtk (for model = 'pbtk'), parameterize_3comp (for model = '3compartment), parameterize_1comp(for model = '1compartment') or parameterize_steadystate (for model = '3compartmentss'), overrides chem.name and chem.cas.	
hourly.dose	Hourly dose rate mg/kg BW/h.	
concentration suppress.messag	Desired concentration type, 'blood', 'tissue', or default 'plasma'.	
	Whether or not the output message is suppressed.	
recalc.blood2plasma		

Recalculates the ratio of the amount of chemical in the blood to plasma using the input parameters. Use this if you have 'altered hematocrit, Funbound.plasma, or Krbc2pu.

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tissue Desired tissue conentration (defaults to whole body concentration.) restrictive.clearance

If TRUE (default), then only the fraction of chemical not bound to protein is available for metabolism in the liver. If FALSE, then all chemical in the liver is metabolized (faster metabolism due to rapid off-binding).

bioactive.free.invivo

If FALSE (default), then the total concentration is treated as bioactive in vivo. If TRUE, the the unbound (free) plasma concentration is treated as bioactive in vivo. Only works with tissue = NULL in current implementation.

Additional parameters passed to parameterize function if parameters is NULL.

#### Value

Steady state plasma concentration in mg/L units

# Author(s)

Robert Pearce and John Wambaugh

calc_css

Find the steady state concentration and the day it is reached.

# Description

This function finds the day a chemical comes within the specified range of the analytical steady state venous blood or plasma concentration(from calc_analytic_css) for the multiple compartment, three compartment, and one compartment models, the fraction of the true steady state value reached on that day, the maximum concentration, and the average concentration at the end of the simulation.

```
calc_css(
  chem.name = NULL,
  chem.cas = NULL,
 dtxsid = NULL,
 parameters = NULL,
  species = "Human",
  f = 0.01,
  daily.dose = 1,
  doses.per.day = 3,
  days = 21,
 output.units = "uM",
  suppress.messages = FALSE,
  tissue = NULL,
 model = "pbtk",
 default.to.human = FALSE,
  f.change = 1e-05,
  adjusted.Funbound.plasma = TRUE,
  regression = TRUE,
 well.stirred.correction = TRUE,
 restrictive.clearance = TRUE,
```

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```
dosing = NULL,
    ...
)
```

### **Arguments**

chem. name Either the chemical name, CAS number, or parameters must be specified.

chem. cas Either the chemical name, CAS number, or parameters must be specified.

dtxsid EPA's DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chem-

ical must be identified by either CAS, name, or DTXSIDs

parameters Chemical parameters from parameterize_pbtk function, overrides chem.name

and chem.cas.

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

f Fractional distance from the final steady state concentration that the average

concentration must come within to be considered at steady state.

daily.dose Total daily dose, mg/kg BW.

doses.per.day Number of doses per day.

days Initial number of days to run simulation that is multiplied on each iteration.

output.units Units for returned concentrations, defaults to uM (specify units = "uM") but can

also be mg/L.

suppress.messages

Whether or not to suppress messages.

tissue Desired tissue concentration (default value is NULL, will depend on model –

see steady.state.compartment in model.info file for further details.)

model Model used in calculation, 'pbtk' for the multiple compartment model, '3compartment'

for the three compartment model, and '1compartment' for the one compartment

model.

default.to.human

Substitutes missing animal values with human values if true (hepatic intrinsic

clearance or fraction of unbound plasma).

f. change Fractional change of daily steady state concentration reached to stop calculating.

adjusted.Funbound.plasma

Uses adjusted Funbound.plasma when set to TRUE along with partition coeffi-

cients calculated with this value.

regression Whether or not to use the regressions in calculating partition coefficients.

well.stirred.correction

Uses correction in calculation of hepatic clearance for well-stirred model if TRUE for model 1compartment elimination rate. This assumes clearance relative to amount unbound in whole blood instead of plasma, but converted to use

with plasma concentration.

restrictive.clearance

Protein binding not taken into account (set to 1) in liver clearance if FALSE.

dosing The dosing object for more complicated scenarios. Defaults to repeated daily.dose

spread out over doses.per.day

.. Additional arguments passed to model solver (default of solve_pbtk).

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## Value

frac	Ratio of the mean concentration on the day steady state is reached (baed on doses.per.day) to the analytical Css (based on infusion dosing).
max	The maximum concentration of the simulation.
avg	The average concentration on the final day of the simulation.
the.day	The day the average concentration comes within $100 * p$ percent of the true steady state concentration.

# Author(s)

Robert Pearce, John Wambaugh

# **Examples**

```
calc_css(chem.name='Bisphenol-A',doses.per.day=5,f=.001,output.units='mg/L')
parms <- parameterize_3comp(chem.name='Bisphenol-A')</pre>
parms$Funbound.plasma <- .07</pre>
calc_css(parameters=parms,model='3compartment')
out <- solve_pbtk(chem.name = "Bisphenol A",</pre>
  days = 50,
  daily.dose=1,
  doses.per.day = 3)
plot.data <- as.data.frame(out)</pre>
css <- calc_analytic_css(chem.name = "Bisphenol A")</pre>
library("ggplot2")
c.vs.t <- ggplot(plot.data,aes(time, Cplasma)) + geom_line() +</pre>
geom_hline(yintercept = css) + ylab("Plasma Concentration (uM)") +
xlab("Day") + theme(axis.text = element_text(size = 16), axis.title =
element_text(size = 16), plot.title = element_text(size = 17)) +
ggtitle("Bisphenol A")
print(c.vs.t)
```

calc_elimination_rate Calculate the elimination rate for a one compartment model.

# Description

This function calculates an elimination rate from the three compartment steady state model where elimination is entirely due to metablism by the liver and glomerular filtration in the kidneys.

```
calc_elimination_rate(
  chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
```

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```
parameters = NULL,
  species = "Human",
  suppress.messages = FALSE,
  default.to.human = FALSE,
  restrictive.clearance = TRUE,
  adjusted.Funbound.plasma = TRUE,
  regression = TRUE,
  well.stirred.correction = TRUE,
  clint.pvalue.threshold = 0.05,
  minimum.Funbound.plasma = 1e-04
)
```

# **Arguments**

chem. cas Either the cas number or the chemical name must be specified.

chem. name Either the chemical name or the cas number must be specified.

dtxsid EPA's 'DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chem-

ical must be identified by either CAS, name, or DTXSIDs

parameters Chemical parameters from parameterize_steadystate or 1compartment function,

overrides chem.name and chem.cas.

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

suppress.messages

Whether or not the output message is suppressed.

default.to.human

Substitutes missing animal values with human values if true.

restrictive.clearance

In calculating elimination rate, protein binding is not taken into account (set to 1) in liver clearance if FALSE.

adjusted.Funbound.plasma

Uses adjusted Funbound.plasma when set to TRUE along with partition coefficients calculated with this value.

regression Whether or not to use the regressions in calculating partition coefficients.

well.stirred.correction

Uses correction in calculation of hepatic clearance for -stirred model if TRUE. This assumes clearance relative to amount unbound in whole blood instead of plasma, but converted to use with plasma concentration.

clint.pvalue.threshold

Hepatic clearance for chemicals where the in vitro clearance assay result has a p-values greater than the threshold are set to zero.

minimum.Funbound.plasma

Monte Carlo draws less than this value are set equal to this value (default is 0.0001 – half the lowest measured Fup in our dataset).

### **Details**

Elimination rate calculated by dividing the total clearance (using the default -stirred hepatic model) by the volume of distribution. When species is specified as rabbit, dog, or mouse, the function uses the appropriate physiological data(volumes and flows) but substitues human fraction unbound, partition coefficients, and intrinsic hepatic clearance.

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#### Value

Elimination rate

Units of 1/h.

### Author(s)

John Wambaugh

# **Examples**

```
calc_elimination_rate(chem.name="Bisphenol A")
calc_elimination_rate(chem.name="Bisphenol A", species="Rat")
calc_elimination_rate(chem.cas="80-05-7")
```

calc_fetal_phys

Calculate maternal-fetal physiological parameters

## **Description**

This function uses the equations from Kapraun (2019) to calculate chemical- independent physiological paramreters as a function of gestational age in weeks.

# Usage

```
calc_fetal_phys(week = 12, ...)
```

## **Arguments**

week

Gestational week

... Additional arguments to parameterize_fetal_pbtk

# **Details**

```
BW = pre_{p}regnant_{B}W + BW_{c}ubic_{t}heta1*tw + BW_{c}ubic_{t}heta2*tw^{2} + BW_{c}ubic_{t}heta3*tw^{3}
```

 $Wadipose = Wadipose_linear_theta0 + Wadipose_linear_theta1 * tw;$ 

 $Wfthyroid = 0.001*Wfthyroid_qompertz_theta0*exp(Wfthyroid_qompertz_theta1/Wfthyroid_qompertz_theta2*exp(Wfthyroid_qompertz_theta3/Wfthyroid_qomper$ 

 $Wfliver = 0.001*Wfliver_qompertz_theta0*exp(Wfliver_qompertz_theta1/Wfliver_qompertz_theta2*(1-exp(-1)))$ 

 $Wfbrain_gompertz_theta0*exp(Wfbrain_gompertz_theta1/Wfbrain_gompertz_theta2*(1-exp(Wfbrain_gompertz_theta2))))) is a simple of the property of the property$ 

 $Wfgut = 0.001*Wfgut_qompertz_theta0*exp(Wfgut_qompertz_theta1/Wfgut_qompertz_theta2*(1-exp(-Wfgut_qompertz_theta2))))$ 

```
Wflung = 0.001*Wflung_q ompertz_t heta0*exp(Wflung_q ompertz_t heta1/Wflung_q ompertz_t heta2*(1-exp(-Wflung_q ompertz_t heta2)))))
hematocrit_quadratic_theta0 + hematocrit_quadratic_theta1 * tw + hematocrit_quadratic_theta2 * powledge for the property of 
              Rblood2plasma = 1 - hematocrit + hematocrit * Krbc2pu * Fraction_unbound_plasma;
fhematocrit = (fhematocrit_cubic_theta1*tw + fhematocrit_cubic_theta2*pow(tw, 2) + fhematocrit_cubic_theta3*pow(tw, 2) + fhematocrit_cubic_theta
Rfblood2plasma = 1 - fhematocrit + fhematocrit * Kfrbc2pu * Fraction_unbound_plasma_fetus;
fBW = 0.001*fBW_q ompertz_t heta0*exp(fBW_q ompertz_t heta1/fBW_q ompertz_t heta2*(1-exp(-fBW_q ompertz_t heta2)))
Vplacenta = 0.001*(Vplacenta_cubic_theta1*tw + Vplacenta_cubic_theta2*pow(tw, 2) + Vplacenta_cubic_theta3*pow(tw, 2) + Vplacenta_cubic_t
Vamnf = 0.001*Vamnf_logistic_theta0/(1+exp(-Vamnf_logistic_theta1*(tw-Vamnf_logistic_theta2)));
Vplasma = Vplasma_mod_logistic_theta0/(1 + exp(-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_logistic_theta1*(tw-Vplasma_mod_
                                                                                                                                        Vrbcs = hematocrit/(1 - hematocrit) * Vplasma;
                                                                                                                                    Vven = venous_blood_fraction * (Vrbcs + Vplasma);
                                                                                                                                  Vart = arterial_blood_fraction * (Vrbcs + Vplasma);
                                                                                                                                                                   Vadipose = 1/adipose_density * Wadipose;
Vffmx = 1/ffmx_density*(BW-Wadipose-(fBW+placenta_density*Vplacenta+amnf_density*Vamnf));
                                                          Vallx = Vart + Vven + Vthyroid + Vkidney + Vgut + Vliver + Vlung; \\
```

Vrest = Vffmx - Vallx;

 $V fart = 0.001 * arterial_blood_fraction * fblood_weight_ratio * fBW;$  $V f ven = 0.001 * venous_blood_f raction * fblood_w eight_ratio * fBW;$  $Vfkidney = 1/kidney_density * Wfkidney;$  $Vfthyroid = 1/thyroid_density * Wfthyroid;$  $Vfliver = 1/liver_density * Wfliver;$  $V fbrain = 1/brain_density * W fbrain;$  $Vfgut = 1/gut_density * Wfgut;$  $V flung = 1/lung_d ensity * W flung;$ Vfrest = fBW - (Vfart + Vfven + Vfbrain + Vfkidney + Vfthyroid + Vfliver + Vfgut + Vflung); $Q cardiac = 24*(Q cardiac_c ubic_t heta 0 + Q cardiac_c ubic_t heta 1*tw + Q cardiac_c ubic_t heta 2*pow(tw,2) + Q cardiac_c ubic_t heta 1*tw + Q cardiac_c ubic_t heta 2*pow(tw,2) + Q cardiac_c ubic_t heta 1*tw + Q$  $Qgut = 0.01*(Qgut_percent_initial + (Qgut_percent_terminal - Qgut_percent_initial)/term*tw)*Qcardiac;$  $Qkidney = 24*(Qkidney_cubic_theta0 + Qkidney_cubic_theta1 *tw + Qkidney_cubic_theta2 *pow(tw, 2) + Qkidney_cubic_theta2$  $Qliver = 0.01*(Qliver_percent_initial + (Qliver_percent_terminal - Qliver_percent_initial) / term*tw)*Qcardiac;$  $Qthyroid = 0.01*(Qthyroid_percent_initial + (Qthyroid_percent_terminal - Qthyroid_percent_terminal)/term*tw)$  $Qplacenta = 24 * Qplacenta_linear_theta1 * 1000 * Vplacenta;$  $Qadipose = 0.01*(Qadipose_percent_initial + (Qadipose_percent_terminal - Qadipose_percent_initial)/term*tw)*(Qadipose_percent_initial)/t$ 

Qrest = Qcardiac - (Qgut + Qkidney + Qliver + Qthyroid + Qplacenta + Qadipose);

 $Qgfr = 60*24*0.001*(Qgfr_quadratic_theta0 + Qgfr_quadratic_theta1*tw + Qgfr_quadratic_theta2*pow(tw, 2));$  $Qfrvtl = 60*24*0.001*Qfrvtl_logistic_theta0/(1+exp(-Qfrvtl_logistic_theta1*(tw-Qfrvtl_logistic_theta2)));$  $Qflvtl = 60*24*0.001*Qflvtl_logistic_theta0/(1+exp(-Qflvtl_logistic_theta1*(tw-Qflvtl_logistic_theta2)));$  $Qfda = 60*24*0.001*Qfda_logistic_theta0/(1+exp(-Qfda_logistic_theta1*(tw-Qfda_logistic_theta2)));$ Qfartb = Qflvtl + Qfda;Qfcardiac = Qfartb;Qflung = Qfrvtl - Qfda; $Qfplacenta = 60*24*0.001*Qfplacenta_logistic_theta0/(1+exp(-Qfplacenta_logistic_theta1*(tw-Qfplacenta_logistic_theta1)))) + (1+exp(-Qfplacenta_logistic_theta1))) +$  $Qfdv = 60*24*0.001*Qfdv_qompertz_theta0*exp(Qfdv_qompertz_theta1/Qfdv_qompertz_theta2*(1-exp(-Qfdv_qompertz_theta1/Qfdv_qompertz_theta2*(1-exp(-Qfdv_qompertz_theta1/Qfdv_qompertz_theta2*(1-exp(-Qfdv_qompertz_theta1/Qfdv_qompertz_theta1/Qfdv_qompertz_theta2*(1-exp(-Qfdv_qompertz_theta1/Qfdv_qompertz_t$  $Qfgut = Qfgut_percent/Qfnonplacental_percent*(1 - Qfplacenta/Qfartb)*Qfartb;$  $Qfkidney = Qfkidney_percent/Qfnonplacental_percent*(1-Qfplacenta/Qfartb)*Qfartb;$  $Qfbrain = Qfbrain_percent/Qfnonplacental_percent*(1 - Qfplacenta/Qfartb)*Qfartb;$  $Qfliver = Qfliver_percent/(100 - (Qbrain_percent + Qkidney_percent + Qgut_percent)) * (1 - (Qfbrain_percent + Qkidney_percent)) * (1 - (Qfbrain_percent)) * (1 - (Qfbrain_$  $Qfthyroid = Qfthyroid_percent/(100 - (Qbrain_percent + Qkidney_percent + Qgut_percent)) * (1 - (Qfbrain_percent + Qkidney_percent) + (100 - Qkrain_percent + Qkidney_percent)) * (1 - Qkrain_percent + Qkidney_percent) * (1 - Qkrain_percent) * (1 - Qkra$ 

Qfbypass = Qfcardiac - Qflung;

Qfrest = Qfcardiac - (Qfplacenta + Qfgut + Qfliver + Qfthyroid + Qfkidney + Qfbrain);

#### Value

list containing:

BW Maternal body weight, kg

Wadipose Maternal adipose fraction of total weight Wfkidney Fetal kidney fraction of total weight Wfthyroid Fetal thyroid fraction of total weight Fetal liver fraction of total weight Wfliver Wfbrain Fetal brain fraction of total weight Wfgut Fetal gut fraction of total weight Fetal lung fraction of total weight Wflung hematocrit Maternal hematocrit fraction of blood

Rblood2plasma Maternal Rblood2plasma

fhematocrit Fetal hematocrit fraction of blood

Rfblood2plasma

Fetal Rfblood2plasma

fBW

Fetal body weight, kg

Vplacenta

Volume of Vplacenta, L

Vamnf

Volume of amniotic fluid, L

Vplasma

Maternal volume of plasma, L

Vrbcs Maternal volume of red blood cells, L Vven Maternal volume of venous blood, L Vart Maternal volume of arterial blood, L

Vadipose Maternal volume of adipose, L
Vffmx Fetal volume ofVffmx, L

Vallx, L

Vrest Maternal volume of rest of body, L<br/>
Vfart Fetal volume of arterial blood, L<br/>
Vfven Fetal volume of venous blood, L

Vfkidney Fetal volume of kidney, L
Vfthyroid Fetal volume of thyroid, L
Vfliver Fetal volume of liver, L
Vfbrain Fetal volume of brain, L
Vfgut Fetal volume of gut, L
Vflung Fetal volume of lung, L

Vfrest Fetal volume of rest of body, L

Qcardiac Maternal cardiac output blood flow, L/day

Qgut Maternal blood flow to gut, L/day
Qkidney Maternal blood flow to kidney, L/day
Qliver Maternal blood flow to liver, L/day
Qthyroid Maternal blood flow to thyroid, L/day
Qplacenta Maternal blood flow to placenta, L/day

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Qadipose Maternal blood flow to adipose, L/day
Qrest Maternal blood flow to rest, L/day

Qgfr Maternal glomerular filtration rate in kidney, L/day

Qfrvtl Fetal blood flow to right ventricle, L/day
Qflvtl Fetal blood flow to left ventircle, L/day

Qfda Fetal blood flow to Qfda, L/day
Qfartb Fetal blood flow to Qfartb, L/day
Qfcardiac Fetal cardiac output blood flow, L/day

Qflung Fetal blood flow to lung, L/day Qfplacenta Fetal blood flow to placenta, L/day Qfdv Fetal blood flow to Qfdv, L/day Qfgut Fetal blood flow to gut, L/day Qfkidney Fetal blood flow to kidney, L/day Ofbrain Fetal blood flow to brain, L/day Ofliver Fetal blood flow to liver, L/day Qfthyroid Fetal blood flow to thyroid, L/day Qfrest Fetal blood flow to rest, L/day **Qfbypass** Fetal blood flow to Qfbypass, L/day

### Author(s)

John Wambaugh

### References

Kapraun, Dustin F., et al. "Empirical models for anatomical and physiological changes in a human mother and fetus during pregnancy and gestation." PloS one 14.5 (2019): e0215906.

calc_half_life

Calculates the half-life for a one compartment model.

### Description

This function calculates the half life from the three compartment steady state model where elimination is entirely due to metabolism by the liver and glomerular filtration in the kidneys.

# Usage

```
calc_half_life(
  chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
  parameters = NULL,
  species = "Human",
  suppress.messages = FALSE,
  default.to.human = FALSE,
```

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```
restrictive.clearance = TRUE,
adjusted.Funbound.plasma = TRUE,
regression = TRUE,
well.stirred.correction = TRUE,
clint.pvalue.threshold = 0.05,
minimum.Funbound.plasma = 1e-04
)
```

### **Arguments**

chem. cas Either the cas number or the chemical name must be specified.

chem. name Either the chemical name or the cas number must be specified.

dtxsid EPA's 'DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chem-

ical must be identified by either CAS, name, or DTXSIDs

parameters Chemical parameters from parameterize_steadystate or 1 compartment function,

overrides chem.name and chem.cas.

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

suppress.messages

Whether or not the output message is suppressed.

default.to.human

Substitutes missing animal values with human values if true.

restrictive.clearance

In calculating elimination rate, protein binding is not taken into account (set to 1) in liver clearance if FALSE.

adjusted.Funbound.plasma

Uses adjusted Funbound.plasma when set to TRUE along with partition coefficients calculated with this value.

regression Whether or not to use the regressions in calculating partition coefficients.

well.stirred.correction

Uses correction in calculation of hepatic clearance for -stirred model if TRUE. This assumes clearance relative to amount unbound in whole blood instead of plasma, but converted to use with plasma concentration.

clint.pvalue.threshold

Hepatic clearance for chemicals where the in vitro clearance assay result has a p-values greater than the threshold are set to zero.

 $\verb|minimum.Funbound.plasma|\\$ 

Monte Carlo draws less than this value are set equal to this value (default is 0.0001 – half the lowest measured Fup in our dataset).

### **Details**

Half life is calculated by dividing the natural-log of 2 by the elimination rate from the one compartment model.

#### Value

Half life Units of h.

### Author(s)

Sarah E. Davidson

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#### See Also

[calc_elimination_rate()] for the elimination rate calculation

## **Examples**

```
calc_half_life(chem.name="Bisphenol A")
calc_half_life(chem.name="Bisphenol A", species="Rat")
calc_half_life(chem.cas="80-05-7")
```

calc_hepatic_clearance

Calculate the hepatic clearance (deprecated).

### **Description**

This function is included for backward compatibility. It calls calc_hep_clearance which calculates the hepatic clearance in plasma for a well-stirred model or other type if specified. Based on Ito and Houston (2004)

# Usage

```
calc_hepatic_clearance(
  chem.name = NULL,
  chem.cas = NULL,
  dtxsid = NULL,
  parameters = NULL,
  species = "Human",
  default.to.human = FALSE,
  hepatic.model = "well-stirred",
  suppress.messages = FALSE,
  well.stirred.correction = TRUE,
  restrictive.clearance = TRUE,
  adjusted.Funbound.plasma = TRUE,
  ...
)
```

# Arguments

```
chem.name Either the chemical name, CAS number, or the parameters must be specified.

chem.cas Either the chemical name, CAS number, or the parameters must be specified.

dtxsid EPA's DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chemical must be identified by either CAS, name, or DTXSIDs

parameters Chemical parameters from parameterize_steadystate function, overrides chem.name and chem.cas.

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

default.to.human
```

Substitutes missing animal values with human values if true.

hepatic.model Model used in calculating hepatic clearance, unscaled, parallel tube, dispersion, or default well-stirred.

suppress.messages

Whether or not to suppress the output message.

well.stirred.correction

Uses correction in calculation of hepatic clearance for well-stirred model if TRUE for hepatic.model well-stirred. This assumes clearance relative to amount unbound in whole blood instead of plasma, but converted to use with plasma concentration.

restrictive.clearance

Protein binding not taken into account (set to 1) in liver clearance if FALSE.

adjusted.Funbound.plasma

Uses adjusted Funbound.plasma when set to TRUE.

. Additional parameters passed to parameterize_steadystate if parameters is NULL.

#### Value

Hepatic Clearance

Units of L/h/kg BW.

### Author(s)

John Wambaugh and Robert Pearce

#### References

Ito, K., & Houston, J. B. (2004). "Comparison of the use of liver models for predicting drug clearance using in vitro kinetic data from hepatic microsomes and isolated hepatocytes." Pharmaceutical Tesearch, 21(5), 785-792.

# **Examples**

```
calc_hep_clearance(chem.name="Ibuprofen",hepatic.model='unscaled')
calc_hep_clearance(chem.name="Ibuprofen",well.stirred.correction=FALSE)
```

calc_hep_bioavailability

Calculate first pass metabolism

## **Description**

For models that don't described first pass blood flow from the gut, need to cacluate a hepatic bioavailability, that is, the fraction of chemical systemically available after metabolism during the first pass through the liver (Rowland, 1973 Equaation 29, where k21 is blood flow through the liver and k23 is clearance from the liver in Figure 1).

calc_hep_clearance 45

## Usage

```
calc_hep_bioavailability(
  chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
  parameters = NULL,
  restrictive.clearance = TRUE,
  flow.34 = TRUE
)
```

## **Arguments**

chem.cas	Chemical Abstract Services Registry Number (CAS-RN) – if parameters is not specified then the chemical must be identified by either CAS, name, or DTXISD	
chem.name	Chemical name (spaces and capitalization ignored) – if parameters is not specified then the chemical must be identified by either CAS, name, or DTXISD	
dtxsid	EPA's 'DSSTox Structure ID (https://comptox.epa.gov/dashboard) – if parameters is not specified then the chemical must be identified by either CAS, name, or DTXSIDs	
parameters	Parameters from the appropriate parameterization function for the model indicated by argument model	
restrictive.clearance		
	Protein binding not taken into account (set to 1) in liver clearance if FALSE.	
flow.34	A logical constraint	

## Value

A data.table whose columns are the parameters of the HTTK model specified in model.

# Author(s)

John Wambaugh

# References

Rowland, Malcolm, Leslie Z. Benet, and Garry G. Graham. "Clearance concepts in pharmacokinetics." Journal of pharmacokinetics and biopharmaceutics 1.2 (1973): 123-136.

calc_hep_clearance Calculate the hepatic clearance.

# Description

This function calculates the hepatic clearance in plasma for using the Houston (2004) are also available. In vitro measured hepatic clearace is corrected for the free fraction in the assay using the model of Kilford et al. (2008).

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### **Usage**

```
calc_hep_clearance(
  chem.name = NULL,
  chem.cas = NULL,
  dtxsid = NULL,
  parameters = NULL,
  species = "Human",
  default.to.human = FALSE,
  hepatic.model = "well-stirred",
  suppress.messages = FALSE,
  well.stirred.correction = TRUE,
  restrictive.clearance = TRUE,
  adjusted.Funbound.plasma = TRUE,
  ...
)
```

### **Arguments**

chem. name Either the chemical name, CAS number, or the parameters must be specified.

Chem. cas Either the chemical name, CAS number, or the parameters must be specified.

dtxsid EPA's DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chem-

ical must be identified by either CAS, name, or DTXSIDs

parameters Chemical parameters from parameterize_steadystate function, overrides chem.name

and chem.cas.

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

default.to.human

Substitutes missing animal values with human values if true.

hepatic.model Model used in calculating hepatic clearance, unscaled, parallel tube, dispersion,

or default well-stirred.

suppress.messages

Whether or not to suppress the output message.

well.stirred.correction

Uses correction in calculation of hepatic clearance for well-stirred model if TRUE for hepatic.model well-stirred. This assumes clearance relative to amount unbound in whole blood instead of plasma, but converted to use with plasma concentration.

restrictive.clearance

Protein binding not taken into account (set to 1) in liver clearance if FALSE.

adjusted.Funbound.plasma

Uses adjusted Funbound.plasma when set to TRUE.

. Additional parameters passed to parameterize_steadystate if parameters is NULL.

### Value

Hepatic Clearance

Units of L/h/kg BW.

### Author(s)

John Wambaugh and Robert Pearce

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#### References

Ito, K., & Houston, J. B. (2004). "Comparison of the use of liver models for predicting drug clearance using in vitro kinetic data from hepatic microsomes and isolated hepatocytes." Pharmaceutical Tesearch, 21(5), 785-792.

Kilford, P. J., Gertz, M., Houston, J. B. and Galetin, A. (2008). Hepatocellular binding of drugs: correction for unbound fraction in hepatocyte incubations using microsomal binding or drug lipophilicity data. Drug Metabolism and Disposition 36(7), 1194-7, 10.1124/dmd.108.020834.

## **Examples**

```
calc_hep_clearance(chem.name="Ibuprofen",hepatic.model='unscaled')
calc_hep_clearance(chem.name="Ibuprofen",well.stirred.correction=FALSE)
```

calc_hep_fu

Calculate the free chemical in the hepaitic clearance assay

## **Description**

Method from Kilford et al. (2008) for fraction of unbound chemical in the hepatocyte intrinsic clearance assay

# Usage

```
calc_hep_fu(
  chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
  parameters = NULL,
  Vr = 0.005,
  pH = 7.4
)
```

# Arguments

chem.cas	Chemical Abstract Services Registry Number (CAS-RN) – if parameters is not specified then the chemical must be identified by either CAS, name, or DTXISD
chem.name	Chemical name (spaces and capitalization ignored) – if parameters is not specified then the chemical must be identified by either CAS, name, or DTXISD
dtxsid	EPA's 'DSSTox Structure ID (https://comptox.epa.gov/dashboard) – if parameters is not specified then the chemical must be identified by either CAS, name, or DTXSIDs
parameters	Parameters from the appropriate parameterization function for the model indicated by argument model
Vr	Rratio of cell volume to incubation volume. Default is taken from
рН	pH of the incupation medium.

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#### Value

A numeric fraction between zero and one

#### Author(s)

John Wambaugh and Robert Pearce

#### References

Kilford, Peter J., et al. "Hepatocellular binding of drugs: correction for unbound fraction in hepatocyte incubations using microsomal binding or drug lipophilicity data." Drug Metabolism and Disposition 36.7 (2008): 1194-1197.

Wetmore, Barbara A., et al. "Incorporating high-throughput exposure predictions with dosimetry-adjusted in vitro bioactivity to inform chemical toxicity testing." Toxicological Sciences 148.1 (2015): 121-136.

calc_ionization

Calculate the ionization.

# Description

This function calculates the ionization of a compound at a given pH. The pKa's are either entered as parameters or taken from a specific compound in the package.

## Usage

```
calc_ionization(
  chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
  parameters = NULL,
  pH = NULL,
  pKa_Donor = NULL,
  pKa_Accept = NULL)
```

chem.cas.

## **Arguments**

chem.cas Either the chemical name or the CAS number must be specified. Either the chemical name or the CAS number must be specified. chem.name dtxsid EPA's 'DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chemical must be identified by either CAS, name, or DTXSIDs parameters Chemical parameters from a parameterize MODEL function, overrides chem.name and chem.cas. pH where ionization is evaluated. Compound H dissociation equilibirum constant(s). Overwrites chem.name and pKa_Donor chem.cas. Compound H association equilibirum constant(s). Overwrites chem.name and pKa_Accept

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#### **Details**

The arguments pKa_Donor and pKa_Accept may be single numbers, characters, or vectors. We support characters because there are many instances with multiple predicted values and all those values can be included by concatenating with commas (for example, pKa_Donor = "8.1,8.6". Finally, pka_Donor and pKa_Accept may be vectors of characters representing different chemicals or instances of chemical parameters to allow for uncertainty analysis.

The fractions are calculated by determining the coefficients for each species and dividing the particular species by the sum of all three. The positive, negative and zwitterionic/neutral coefficients are given by:

```
zwitter/netural = 1 for(iin1: pkabove)negative = negative + 10^{(i*pH - pKa1 - ... - pKai)} for(iin1: pkbelow)positive = positive + 10^{(pKa1 + ... + pKai - i*pH)}
```

where i begins at 1 and ends at the number of points above(for negative) or below(for positive) the neutral/zwitterionic range. The neutral/zwitterionic range is either the pH range between 2 pKa's where the number of acceptors above is equal to the number of donors below, everything above the pKa acceptors if there are no donors, or everything below the pKa donors if there are no acceptors. Each of the terms in the sums represent a different ionization.

#### Value

```
fraction_neutral
fraction of compound neutral
fraction_charged
fraction of compound charged
fraction_negative
fraction of compound negative
fraction_positive
fraction of compound positive
fraction_zwitter
fraction of compound zwitterionic
```

# Author(s)

Robert Pearce and John Wambaugh

### References

Pearce, Robert G., et al. "Evaluation and calibration of high-throughput predictions of chemical distribution to tissues." Journal of Pharmacokinetics and Pharmacodynamics 44.6 (2017): 549-565.

Strope, Cory L., et al. "High-throughput in-silico prediction of ionization equilibria for pharmacokinetic modeling." Science of The Total Environment 615 (2018): 150-160.

### **Examples**

```
# Donor pKa's 9.78,10.39 -- Should be almost all neutral at plasma pH:
out <- calc_ionization(chem.name='bisphenola',pH=7.4)
print(out)
out[["fraction_neutral"]]==max(unlist(out))

# Donor pKa's 9.78,10.39 -- Should be almost all negative (anion) at higher pH:
out <- calc_ionization(chem.name='bisphenola',pH=11)</pre>
```

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```
print(out)
out[["fraction_negative"]]==max(unlist(out))
# Fictious compound, should be almost all all negative (anion):
out <- calc_ionization(pKa_Donor=8,pKa_Accept="1,4",pH=9)</pre>
print(out)
out[["fraction_negative"]]>0.9
# Donor pKa 6.54 -- Should be mostly negative (anion):
out <- calc_ionization(chem.name='Acephate',pH=7)</pre>
print(out)
out[["fraction_negative"]]==max(unlist(out))
#Acceptor pKa's "9.04,6.04" -- Should be almost all positive (cation) at plasma pH:
out <- calc_ionization(chem.cas="145742-28-5",pH=7.4)</pre>
print(out)
out[["fraction_positive"]]==max(unlist(out))
#Fictious Zwitteron:
out <- calc_ionization(pKa_Donor=6,pKa_Accept="8",pH=7.4)
print(out)
out[["fraction_zwitter"]]==max(unlist(out))
```

calc_krbc2pu

Back-calculates the Red Blood Cell to Unbound Plasma Partition Coefficient

# **Description**

Given an observed ratio of chemical concentration in blood to plasma, this function calculates a Red Blood Cell to unbound plasma (Krbc2pu) partition coefficient that would be consistent with that observation.

### Usage

```
calc_krbc2pu(
  Rb2p,
  Funbound.plasma,
  hematocrit = NULL,
  default.to.human = FALSE,
  species = "Human",
  suppress.messages = TRUE
)
```

# Arguments

Rb2p The chemical blood:plasma concentration ratop

Funbound.plasma

The free fraction of chemical in the presence of plasma protein Rblood2plasma.

hematocrit Overwrites default hematocrit value in calculating Rblood2plasma.

default.to.human

Substitutes missing animal values with human values if true.

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species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human"). suppress.messages

Determine whether to display certain usage feedback.

### Value

The red blood cell to unbound chemical in plasma partition coefficient.

#### Author(s)

John Wambaugh and Robert Pearce

## References

Pearce, Robert G., et al. "Evaluation and calibration of high-throughput predictions of chemical distribution to tissues." Journal of pharmacokinetics and pharmacodynamics 44.6 (2017): 549-565.

Ruark, Christopher D., et al. "Predicting passive and active tissue: plasma partition coefficients: interindividual and interspecies variability." Journal of pharmaceutical sciences 103.7 (2014): 2189-2198.

calc_maternal_bw

Calculate maternal body weight

### **Description**

This function initializes the parameters needed in the functions solve_fetal_pbtk by calling solve_pbtk and adding additional parameters.

### Usage

```
calc_maternal_bw(week = 12)
```

## **Arguments**

week

Gestational week

### **Details**

```
BW <- params$pre_pregnant_BW + params$BW_cubic_theta1 * tw + params$BW_cubic_theta2 * tw^2 + params$BW_cubic_theta3 * tw^3
```

## Value

BW

Maternal Body Weight, kg.

## Author(s)

John Wambaugh

## References

Kapraun, Dustin F., et al. "Empirical models for anatomical and physiological changes in a human mother and fetus during pregnancy and gestation." PloS one 14.5 (2019): e0215906.

calc_mc_css

Find the monte carlo steady state concentration.

### **Description**

This function finds the analytical steady state plasma concentration(from calc_analytic_css) using a monte carlo simulation (monte_carlo).

# Usage

```
calc_mc_css(
  chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
  parameters = NULL,
  samples = 1000,
  which.quantile = 0.95,
  species = "Human",
  suppress.messages = FALSE,
  model = "3compartmentss",
  httkpop = TRUE,
  invitrouv = TRUE.
  calcrb2p = TRUE,
  censored.params = list(),
  vary.params = list(),
  return.samples = FALSE,
  tissue = NULL,
  output.units = "mg/L",
  invitro.mc.arg.list = list(adjusted.Funbound.plasma = TRUE, poormetab = TRUE,
  fup.censored.dist = FALSE, fup.lod = 0.01, fup.meas.cv = 0.4, clint.meas.cv = 0.3,
    fup.pop.cv = 0.3, clint.pop.cv = 0.3),
 httkpop.generate.arg.list = list(method = "direct resampling", gendernum = NULL,
   agelim_years = NULL, agelim_months = NULL, weight_category = c("Underweight"
   "Normal", "Overweight", "Obese"), gfr_category = c("Normal", "Kidney Disease",
    "Kidney Failure"), reths = c("Mexican American", "Other Hispanic",
    "Non-Hispanic White", "Non-Hispanic Black", "Other")),
  convert.httkpop.arg.list = list(),
 parameterize.arg.list = list(default.to.human = FALSE, clint.pvalue.threshold = 0.05,
    restrictive.clearance = TRUE, regression = TRUE),
  calc.analytic.css.arg.list = list(well.stirred.correction = TRUE,
  adjusted.Funbound.plasma = TRUE, regression = TRUE, IVIVE = NULL, tissue = tissue,
    restrictive.clearance = TRUE, bioactive.free.invivo = FALSE),
  parameterize.args = list(default.to.human = F, adjusted.Funbound.plasma = T,
    regression = T, minimum.Funbound.plasma = 1e-04)
)
```

## **Arguments**

chem.cas

Chemical Abstract Services Registry Number (CAS-RN) – if parameters is not specified then the chemical must be identified by either CAS, name, or DTXISD

chem. name Chemical name (spaces and capitalization ignored) – if parameters is not speci-

fied then the chemical must be identified by either CAS, name, or DTXISD

dtxsid EPA's DSSTox Structure ID (https://comptox.epa.gov/dashboard) – if pa-

rameters is not specified then the chemical must be identified by either CAS,

name, or DTXSIDs

parameters Parameters from the appropriate parameterization function for the model indi-

cated by argument model

samples Number of samples generated in calculating quantiles.

which quantile Which quantile from Monte Carlo simulation is requested. Can be a vector.

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

Species must be set to "Human" to run httkpop model.

suppress.messages

Whether or not to suppress output message.

model Model used in calculation, 'gas pbtk' for the gas pbtk model, 'pbtk' for the mul-

tiple compartment model, '3compartment' for the three compartment model, '3compartmentss' for the three compartment steady state model, and '1compartment' for one compartment model. This only applies when httkpop=TRUE

and species="Human", otherwise '3compartmentss' is used.

httkpop Whether or not to use population generator and sampler from httkpop. This is

overwrites censored.params and vary.params and is only for human physiology.

Species must also be set to 'Human'.

invitrouv Logical to indicate whether to include in vitro parameters in uncertainty and

variability analysis

calcrb2p Logical determining whether or not to recalculate the chemical ratio of blood to

plasma

censored.params

The parameters listed in censored.params are sampled from a normal distribution that is censored for values less than the limit of detection (specified separately for each parameter). This argument should be a list of sublists. Each sublist is named for a parameter in "parameters" and contains two elements: "CV" (coefficient of variation) and "LOD" (limit of detection, below which parameter values are censored. New values are sampled with mean equal to the value in "parameters" and standard deviation equal to the mean times the CV. Censored values are sampled on a uniform distribution between 0 and the limit

of detection. Not used with httkpop model.

vary.params The parameters listed in vary.params are sampled from a normal distribution that

is truncated at zero. This argument should be a list of coefficients of variation (CV) for the normal distribution. Each entry in the list is named for a parameter in "parameters". New values are sampled with mean equal to the value in "parameters" and standard deviation equal to the mean times the CV. Not used with

httkpop model.

return.samples Whether or not to return the vector containing the samples from the simulation

instead of the selected quantile.

tissue Desired steady state tissue concentration.

output.units Plasma concentration units, either uM or default mg/L.

invitro.mc.arg.list

List of additional parameters passed to invitro_mc

httkpop.generate.arg.list

Additional parameters passed to httkpop_generate.

convert.httkpop.arg.list

Additional parameters passed to the convert_httkpop_* function for the model.

parameterize.arg.list

Additional parameters passed to the parameterize_* function for the model.

calc.analytic.css.arg.list

Additional parameters passed to calc_analytic_css.

parameterize.args

A list of arguments to be passed to the model parameterization function (that is, parameterize_MODEL) corresponding to argument "model". (Defaults to NULL.)

#### **Details**

All arguments after httkpop only apply if httkpop is set to TRUE and species to "Human".

When species is specified as rabbit, dog, or mouse, the function uses the appropriate physiological data (volumes and flows) but substitutes human fraction unbound, partition coefficients, and intrinsic hepatic clearance.

Tissue concentrations are calculated for the pbtk model with a default oral infusion dosing. All tissues other than gut, liver, and lung are the product of the steady state plasma concentration and the tissue to plasma partition coefficient.

The six sets of plausible *in vitro-in vivo* extrapolation (IVIVE) assumptions identified by Honda et al. (2019) are:

	in vivo Conc.	Metabolic Clearance	Bioactive Chemical Conc.	TK Statistic Used*
Honda1	Veinous (Plasma)	Restrictive	Free	Mean Conc.
Honda2	Veinous	Restrictive	Free	Max Conc.
Honda3	Veinous	Non-restrictive	Total	Mean Conc.
Honda4	Veinous	Non-restrictive	Total	Max Conc.
Honda5	Target Tissue	Non-restrictive	Total	Mean Conc.
Honda6	Target Tissue	Non-restrictive	Total	Max Conc.

^{*}Assumption is currently ignored because analytical steady-state solutions are currently used by this function.

## Value

Quantiles (specified by which quantile) of the distribution of plasma steady-stae concentration (Css) from the Monte Carlo simulation

## Author(s)

Caroline Ring, Robert Pearce, John Wambaugh, Miyuki Breen

## References

Wambaugh, John F., et al. "Toxicokinetic triage for environmental chemicals." Toxicological Sciences 147.1 (2015): 55-67.

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment international 106 (2017): 105-118.

Honda, Gregory S., et al. "Using the Concordance of In Vitro and In Vivo Data to Evaluate Extrapolation Assumptions." 2019. PLoS ONE 14(5): e0217564.

Rowland, Malcolm, Leslie Z. Benet, and Garry G. Graham. "Clearance concepts in pharmacokinetics." Journal of pharmacokinetics and biopharmaceutics 1.2 (1973): 123-136.

## **Examples**

```
set.seed(1234)
calc_mc_css(chem.name='Bisphenol A',output.units='uM',
            samples=100,return.samples=TRUE)
set.seed(1234)
calc_mc_css(chem.name='Bisphenol A',output.units='uM',httkpop.generate.arg.list=list(method='vi'))
set.seed(1234)
calc_mc_css(chem.name='2,4-d',which.quantile=.9,httkpop=FALSE,tissue='heart')
set.seed(1234)
calc_mc_css(chem.cas = "80-05-7", which.quantile = 0.5,
            output.units = "uM", samples = 2000,
            httkpop.generate.arg.list=list(method='vi', gendernum=NULL,
            agelim_years=NULL, agelim_months=NULL, weight_category =
            c("Underweight", "Normal", "Overweight", "Obese")))
params <- parameterize_pbtk(chem.cas="80-05-7")</pre>
set.seed(1234)
calc_mc_css(parameters=params,model="pbtk")
set.seed(1234)
# Standard HTTK Monte Carlo:
NSAMP = 500
calc_mc_css(chem.cas="90-43-7",model="pbtk",samples=NSAMP)
set.seed(1234)
calc_mc_css(chem.cas="90-43-7",
model="pbtk",
samples=NSAMP,
invitro.mc.arg.list = list(
  adjusted.Funbound.plasma = TRUE,
  poormetab = TRUE,
  fup.censored.dist = FALSE,
  fup.lod = 0.01,
  fup.meas.cv = 0.0,
  clint.meas.cv = 0.0,
  fup.pop.cv = 0.3,
  clint.pop.cv = 0.3))
set.seed(1234)
# HTTK Monte Carlo with no HTTK-Pop physiological variability):
calc_mc_css(chem.cas="90-43-7",model="pbtk",samples=NSAMP,httkpop=FALSE)
set.seed(1234)
# HTTK Monte Carlo with no in vitro uncertainty and variability):
```

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```
calc_mc_css(chem.cas="90-43-7",model="pbtk",samples=NSAMP,invitrouv=FALSE)
set.seed(1234)
# HTTK Monte Carlo with no HTTK-Pop and no in vitro uncertainty and variability):
calc_mc_css(chem.cas="90-43-7",model="pbtk",samples=NSAMP,httkpop=FALSE,invitrouv=FALSE)
# Should be the same as the mean result:
calc_analytic_css(chem.cas="90-43-7",model="pbtk",output.units="mg/L")
set.seed(1234)
# HTTK Monte Carlo using basic Monte Carlo sampler:
calc_mc_css(chem.cas="90-43-7",
model="pbtk",
samples=NSAMP,
httkpop=FALSE,
invitrouv=FALSE,
invitrouv=FALSE,
vary.params=list(Pow=0.3))
```

calc_mc_oral_equiv

Calculate Monte Carlo Oral Equivalent Dose

## **Description**

This functions converts a chemical plasma concetration to an oral equivalent dose using a concentration obtained from calc_mc_css.

## Usage

```
calc_mc_oral_equiv(
 conc,
 chem.name = NULL,
 chem.cas = NULL,
 dtxsid = NULL,
 which.quantile = 0.95,
  species = "Human",
  input.units = "uM",
 output.units = "mgpkgpday",
  suppress.messages = FALSE,
 return.samples = FALSE,
  concentration = "plasma",
  restrictive.clearance = TRUE,
 bioactive.free.invivo = FALSE,
  tissue = NULL,
 IVIVE = NULL,
)
```

## **Arguments**

conc Bioactive in vitro concentration in units of uM.

chem. name Either the chemical name or the CAS number must be specified.

chem. cas Either the CAS number or the chemical name must be specified.

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dtxsid EPA's 'DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chem-

ical must be identified by either CAS, name, or DTXSIDs

which quantile Which quantile from Monte Carlo steady-state simulation (calc_mc_css) is re-

quested. Can be a vector. Note that 95th concentration quantile is the same

population as the 5th dose quantile.

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

input.units Units of given concentration, default of uM but can also be mg/L.

output.units Units of dose, default of 'mgpkgpday' for mg/kg BW/ day or 'umolpkgpday'

for umol/ kg BW/ day.

suppress.messages

Suppress text messages.

return.samples Whether or not to return the vector containing the samples from the simulation

instead of the selected quantile.

concentration Desired concentration type, 'blood', 'tissue', or default 'plasma'.

restrictive.clearance

Protein binding not taken into account (set to 1) in liver clearance if FALSE.

bioactive.free.invivo

If FALSE (default), then the total concentration is treated as bioactive in vivo. If TRUE, the the unbound (free) plasma concentration is treated as bioactive in

vivo. Only works with tissue = NULL in current implementation.

tissue Desired steady state tissue conentration.

IVIVE Honda et al. (2019) identified six plausible sets of assumptions for in vitro-

*in vivo* extrapolation (IVIVE) assumptions. Argument may be set to "Honda1" through "Honda6". If used, this function overwrites the tissue, restrictive.clearance,

and plasma.binding arguments. See Details below for more information.

... Additional parameters passed to calc_mc_css for httkpop and variance of pa-

rameters.

## **Details**

All arguments after httkpop only apply if httkpop is set to TRUE and species to "Human".

When species is specified as rabbit, dog, or mouse, the function uses the appropriate physiological data(volumes and flows) but substitutes human fraction unbound, partition coefficients, and intrinsic hepatic clearance.

Tissue concentrations are calculated for the pbtk model with oral infusion dosing. All tissues other than gut, liver, and lung are the product of the steady state plasma concentration and the tissue to plasma partition coefficient.

The six sets of plausible *in vitro-in vivo* extrapolation (IVIVE) assumptions identified by Honda et al. (2019) are:

	in vivo Conc.	Metabolic Clearance	Bioactive Chemical Conc.	TK Statistic Used*
Honda1	Veinous (Plasma)	Restrictive	Free	Mean Conc.
Honda2	Veinous	Restrictive	Free	Max Conc.
Honda3	Veinous	Non-restrictive	Total	Mean Conc.
Honda4	Veinous	Non-restrictive	Total	Max Conc.
Honda5	Target Tissue	Non-restrictive	Total	Mean Conc.
Honda6	Target Tissue	Non-restrictive	Total	Max Conc.

*Assumption is currently ignored because analytical steady-state solutions are currently used by this function.

#### Value

Equivalent dose in specified units, default of mg/kg BW/day.

## Author(s)

John Wambaugh

#### References

Wetmore, Barbara A., et al. "Incorporating high-throughput exposure predictions with dosimetry-adjusted in vitro bioactivity to inform chemical toxicity testing." Toxicological Sciences 148.1 (2015): 121-136.

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment international 106 (2017): 105-118.

Honda, Gregory S., et al. "Using the Concordance of In Vitro and In Vivo Data to Evaluate Extrapolation Assumptions." 2019. PLoS ONE 14(5): e0217564.

Rowland, Malcolm, Leslie Z. Benet, and Garry G. Graham. "Clearance concepts in pharmacokinetics." Journal of pharmacokinetics and biopharmaceutics 1.2 (1973): 123-136.

# **Examples**

```
 calc\_mc\_oral\_equiv(\emptyset.1, chem. cas="34256-82-1", which.quantile=c(\emptyset.05, \emptyset.5, \emptyset.95), \\ tissue='brain')
```

calc_mc_tk

Conduct multiple TK simulations using Monte Carlo

## **Description**

This function finds the analytical steady state plasma concentration(from calc_analytic_css) using a monte carlo simulation (monte_carlo).

# Usage

```
calc_mc_tk(
  chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
  parameters = NULL,
  samples = 1000,
  which.quantile = 0.95,
  species = "Human",
```

```
suppress.messages = FALSE,
 model = "pbtk",
 httkpop = TRUE,
  invitrouv = TRUE,
 calcrb2p = TRUE,
 censored.params = list(),
 vary.params = list(),
 return.samples = FALSE,
  tissue = NULL,
 output.units = "mg/L",
 solvemodel.arg.list = list(times = c(0, 0.25, 0.5, 0.75, 1, 1.5, 2, 2.5, 3, 4, 5)),
 invitro.mc.arg.list = list(adjusted.Funbound.plasma = TRUE, poormetab = TRUE,
  fup.censored.dist = FALSE, fup.lod = 0.01, fup.meas.cv = 0.4, clint.meas.cv = 0.3,
    fup.pop.cv = 0.3, clint.pop.cv = 0.3),
 httkpop.generate.arg.list = list(method = "direct resampling", gendernum = NULL,
   agelim_years = NULL, agelim_months = NULL, weight_category = c("Underweight",
   "Normal", "Overweight", "Obese"), gfr_category = c("Normal", "Kidney Disease",
    "Kidney Failure"), reths = c("Mexican American", "Other Hispanic",
    "Non-Hispanic White", "Non-Hispanic Black", "Other")),
 convert.httkpop.arg.list = list(),
 parameterize.arg.list = list(default.to.human = FALSE, clint.pvalue.threshold = 0.05,
    restrictive.clearance = TRUE, regression = TRUE),
 return.all.sims = FALSE
)
```

### **Arguments**

chem. cas Either the CAS number, parameters, or the chemical name must be specified.

chem. name Either the chemical parameters, name, or the CAS number must be specified.

dtxsid EPA's DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chem-

ical must be identified by either CAS, name, or DTXSIDs

parameters Parameters from parameterize_steadystate. Not used with httkpop model.

samples Number of samples generated in calculating quantiles.

which quantile Which quantile from Monte Carlo simulation is requested. Can be a vector.

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

Species must be set to "Human" to run httkpop model.

suppress.messages

Whether or not to suppress output message.

model Model used in calculation: 'pbtk' for the multiple compartment model, '3compartment'

for the three compartment model, '3compartmentss' for the three compartment steady state model, and '1compartment' for one compartment model. This only applies when httkpop=TRUE and species="Human", otherwise '3compart-

mentss' is used.

httkpop Whether or not to use population generator and sampler from httkpop. This is

overwrites censored.params and vary.params and is only for human physiology.

Species must also be set to 'Human'.

invitrouv Logical to indicate whether to include in vitro parameters in uncertainty and

variability analysis

calcrb2p Logical determining whether or not to recalculate the chemical ratio of blood to

plasma

censored.params

The parameters listed in censored.params are sampled from a normal distribution that is censored for values less than the limit of detection (specified separately for each parameter). This argument should be a list of sub-lists. Each sublist is named for a parameter in "parameters" and contains two elements: "CV" (coefficient of variation) and "LOD" (limit of detection, below which parameter values are censored. New values are sampled with mean equal to the value in "parameters" and standard deviation equal to the mean times the CV. Censored values are sampled on a uniform distribution between 0 and the limit of detection. Not used with httkpop model.

vary.params

The parameters listed in vary.params are sampled from a normal distribution that is truncated at zero. This argument should be a list of coefficients of variation (CV) for the normal distribution. Each entry in the list is named for a parameter in "parameters". New values are sampled with mean equal to the value in "parameters" and standard deviation equal to the mean times the CV. Not used with httkpop model.

return.samples Whether or not to return the vector containing the samples from the simulation instead of the selected quantile.

tissue Desired steady state tissue conentration.

output.units Plasma concentration units, either uM or default mg/L.

solvemodel.arg.list

Additional arguments ultimately passed to solve_model

invitro.mc.arg.list

List of additional parameters passed to invitro_mc

httkpop.generate.arg.list

Additional parameters passed to httkpop_generate.

convert.httkpop.arg.list

Additional parameters passed to the convert_httkpop_* function for the model.

parameterize.arg.list

Additional parameters passed to the parameterize_* function for the model.

return.all.sims

Logical indicating whether to return the results of all simulations, in addition to the default toxicokinetic statistics

## Details

All arguments after httkpop only apply if httkpop is set to TRUE and species to "Human".

When species is specified as rabbit, dog, or mouse, the function uses the appropriate physiological data(volumes and flows) but substitues human fraction unbound, partition coefficients, and intrinsic hepatic clearance.

Tissue concentrations are calculated for the pbtk model with oral infusion dosing. All tissues other than gut, liver, and lung are the product of the steady state plasma concentration and the tissue to plasma partition coefficient.

The six sets of plausible *in vitro-in vivo* extrpolation (IVIVE) assumptions identified by Honda et al. (2019) are:

	in vivo Conc.	Metabolic Clearance	Bioactive Chemical Conc.	TK Statistic Used*
Honda1	Veinous (Plasma)	Restrictive	Free	Mean Conc.
Honda2	Veinous	Restrictive	Free	Max Conc.
Honda3	Veinous	Non-restrictive	Total	Mean Conc.

Honda4	Veinous	Non-restrictive	Total	Max Conc.
Honda5	Target Tissue	Non-restrictive	Total	Mean Conc.
Honda6	Target Tissue	Non-restrictive	Total	Max Conc.

^{*}Assumption is currently ignored because analytical steady-state solutions are currently used by this function.

### Value

If return.all.sims == FALSE (default) a list with:

means The mean concentration for each model compartment as a function of time

across the Monte Carlo simulation

sds The standard deviation for each model compartment as a function of time across

the Monte Carlo simulation

If return.all.sums == TRUE then a list is returned with:

stats The list of means and sds from return.all.sums=FALSE

sims The concentration vs. time results for each compartment for every (samples) set

of parameters in the Monte Carlo simulation

### Author(s)

John Wambaugh

## **Examples**

```
NSAMP <- 50
chemname="Abamectin"
times<- c(0,0.25,0.5,0.75,1,1.5,2,2.5,3,4,5)
age.ranges <- seq(6,80,by=10)
forward <- NULL
for (age.lower in age.ranges)
  label <- paste("Ages ",age.lower,"-",age.lower+4,sep="")</pre>
  set.seed(1234)
  forward[[label]] <- calc_mc_tk(</pre>
                         chem.name=chemname,
                         samples=NSAMP,
                         httkpop.generate.arg.list=list(
                           method="d",
                           agelim_years = c(age.lower, age.lower+9)),
                         solvemodel.arg.list = list(
                           times=times))
}
```

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calc_rblood2plasma	Calculate the constant ratio of the blood concentration to the plasma concentration.
	concentration.

# Description

This function calculates the constant ratio of the blood concentration to the plasma concentration.

# Usage

```
calc_rblood2plasma(
  chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
  parameters = NULL,
  hematocrit = NULL,
  Krbc2pu = NULL,
  Funbound.plasma = NULL,
  default.to.human = FALSE,
  species = "Human",
  adjusted.Funbound.plasma = TRUE,
  suppress.messages = TRUE
)
```

# Arguments

chem.cas	Either the CAS number or the chemical name must be specified.	
chem.name	Either the chemical name or the CAS number must be specified.	
dtxsid	EPA's DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chemical must be identified by either CAS, name, or DTXSIDs	
parameters	Parameters from parameterize_schmitt	
hematocrit	Overwrites default hematocrit value in calculating Rblood2plasma.	
Krbc2pu	The red blood cell to unbound plasma chemical partition coefficient, typically from predict_partitioning_schmitt	
Funbound.plasma		
	The fraction of chemical unbound (free) in the presence of plasma protein	
default.to.human		
	Substitutes missing animal values with human values if true.	
species	Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").	
adjusted.Funbound.plasma		
	Whether or not to use Funbound.plasma adjustment.	
suppress.messages		
	Determine whether to display certain usage feedback.	

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#### **Details**

The red blood cell (RBC) parition coefficient as predicted by the Schmitt (2008) method is used in the calculation. The value is calculated with the equation: 1 - hematocrit + hematocrit * Krbc2pu * Funbound.plasma, summing the red blood cell to plasma and plasma:plasma (equal to 1) partition coefficients multiplied by their respective fractional volumes. When species is specified as rabbit, dog, or mouse, the function uses the appropriate physiological data (hematocrit and temperature), but substitutes human fraction unbound and tissue volumes.

#### Value

The blood to plasma chemical concentration ratio

## Author(s)

John Wambaugh and Robert Pearce

### References

Schmitt W. "General approach for the calculation of tissue to plasma partition coefficients." Toxicology In Vitro, 22, 457-467 (2008).

Pearce, Robert G., et al. "Evaluation and calibration of high-throughput predictions of chemical distribution to tissues." Journal of pharmacokinetics and pharmacodynamics 44.6 (2017): 549-565.

Ruark, Christopher D., et al. "Predicting passive and active tissue: plasma partition coefficients: interindividual and interspecies variability." Journal of pharmaceutical sciences 103.7 (2014): 2189-2198.

## **Examples**

```
calc_rblood2plasma(chem.name="Bisphenol A")
calc_rblood2plasma(chem.name="Bisphenol A",species="Rat")
```

calc_stats

Calculate toxicokinetic summary statistics (deprecated).

### **Description**

#' This function is included for backward compatibility. It calls calc_tkstats which calculates the area under the curve, the mean, and the peak values for the venous blood or plasma concentration of a specified chemical or all chemicals if none is specified for the multiple compartment model with a given number of days, dose, and number of doses per day.

## Usage

```
calc_stats(
  chem.name = NULL,
  chem.cas = NULL,
  dtxsid = NULL,
  parameters = NULL,
  route = "oral",
```

64 calc_stats

```
stats = c("AUC", "peak", "mean"),
  species = "Human",
  days = 28,
  daily.dose = 1,
  dose = NULL,
  doses.per.day = 1,
  output.units = "uM"
  concentration = "plasma",
  tissue = "plasma",
 model = "pbtk",
  default.to.human = FALSE,
  adjusted.Funbound.plasma = TRUE,
  regression = TRUE,
  restrictive.clearance = TRUE,
  suppress.messages = FALSE,
)
```

## **Arguments**

Name of desired chemical. chem.name

chem.cas CAS number of desired chemical.

dtxsid EPA's DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chem-

ical must be identified by either CAS, name, or DTXSIDs

Chemical parameters from parameterize_pbtk function, overrides chem.name parameters

and chem.cas.

route String specification of route of exposure for simulation: "oral", "iv", "inhala-

tion", ...

stats Desired values (either 'AUC', 'mean', 'peak', or a vector containing any com-

bination).

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

Length of the simulation. days Total daily dose, mg/kg BW. daily.dose

Amount of a single dose at time zero, mg/kg BW. dose

doses.per.day Number of doses per day.

Desired units (either "mg/L", "mg", "umol", or default "uM"). output.units Desired concentration type, 'blood' or default 'plasma'.

tissue Desired steady state tissue conentration.

mode1 Model used in calculation, 'pbtk' for the multiple compartment model, '3compartment'

for the three compartment model, '3compartmentss' for the three compartment

steady state model, and '1compartment' for one compartment model.

default.to.human

concentration

Substitutes missing animal values with human values if true (hepatic intrinsic clearance or fraction of unbound plasma).

adjusted.Funbound.plasma

Uses adjusted Funbound.plasma when set to TRUE along with partition coeffi-

cients calculated with this value.

Whether or not to use the regressions in calculating partition coefficients. regression

calc_tkstats 65

```
restrictive.clearance
```

Protein binding not taken into account (set to 1) in liver clearance if FALSE.

suppress.messages

Whether to suppress output message.

... Arguments passed to solve function.

### Details

Default value of 0 for doses.per.day solves for a single dose.

When species is specified as rabbit, dog, or mouse, the function uses the appropriate physiological data(volumes and flows) but substitues human fraction unbound, partition coefficients, and intrinsic hepatic clearance.

### Value

AUC Area under the plasma concentration curve.

mean.conc The area under the curve divided by the number of days.

peak.conc The highest concentration.

## Author(s)

Robert Pearce and John Wambaugh

calc_tkstats

Calculate toxicokinetic summary statistics.

### **Description**

This function calculates the area under the curve, the mean, and the peak values for the venous blood or plasma concentration of a specified chemical or all chemicals if none is specified for the multiple compartment model with a given number of days, dose, and number of doses per day.

### Usage

```
calc_tkstats(
  chem.name = NULL,
  chem.cas = NULL,
 dtxsid = NULL,
 parameters = NULL,
  route = "oral",
  stats = c("AUC", "peak", "mean"),
  species = "Human",
  days = 28,
 daily.dose = 1,
 dose = NULL,
 doses.per.day = 1,
 output.units = "uM",
  concentration = "plasma",
  tissue = "plasma",
 model = "pbtk",
```

66 calc_tkstats

```
default.to.human = FALSE,
adjusted.Funbound.plasma = TRUE,
regression = TRUE,
restrictive.clearance = TRUE,
suppress.messages = FALSE,
...
)
```

### **Arguments**

chem.name Name of desired chemical.

chem. cas CAS number of desired chemical.

dtxsid EPA's DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chem-

ical must be identified by either CAS, name, or DTXSIDs

parameters Chemical parameters from parameterize_pbtk function, overrides chem.name

and chem.cas.

route String specification of route of exposure for simulation: "oral", "iv", "inhala-

tion", ...

stats Desired values (either 'AUC', 'mean', 'peak', or a vector containing any com-

bination).

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

days Length of the simulation.

daily.dose Total daily dose, mg/kg BW.

dose Amount of a single dose at time zero, mg/kg BW.

doses.per.day Number of doses per day.

 $output.units \qquad Desired\ units\ (either\ "mg/L",\ "mg",\ "umol",\ or\ default\ "uM").$ 

concentration Desired concentration type, 'blood' or default 'plasma'.

tissue Desired steady state tissue conentration.

model Model used in calculation, 'pbtk' for the multiple compartment model, '3compartment'

for the three compartment model, '3compartmentss' for the three compartment

steady state model, and '1compartment' for one compartment model.

default.to.human

Substitutes missing animal values with human values if true (hepatic intrinsic

clearance or fraction of unbound plasma).

adjusted.Funbound.plasma

Uses adjusted Funbound.plasma when set to TRUE along with partition coeffi-

cients calculated with this value.

regression Whether or not to use the regressions in calculating partition coefficients.

restrictive.clearance

Protein binding not taken into account (set to 1) in liver clearance if FALSE.

 ${\tt suppress.messages}$ 

Whether to suppress output message.

... Arguments passed to solve function.

### **Details**

Default value of 0 for doses.per.day solves for a single dose.

When species is specified as rabbit, dog, or mouse, the function uses the appropriate physiological data(volumes and flows) but substitues human fraction unbound, partition coefficients, and intrinsic hepatic clearance.

calc_total_clearance 67

### Value

AUC Area under the plasma concentration curve.

mean.conc The area under the curve divided by the number of days.

peak.conc The highest concentration.

## Author(s)

Robert Pearce and John Wambaugh

## **Examples**

```
calc_tkstats(chem.name='Bisphenol-A',days=100,stats='mean',model='3compartment')
calc_tkstats(chem.name='Bisphenol-A',days=100,stats=c('peak','mean'),species='Rat')
triclosan.stats <- calc_tkstats(days=10, chem.name = "triclosan")</pre>
```

calc_total_clearance Calculate the total plasma clearance.

# **Description**

This function calculates the total clearance rate for a one compartment model for plasma where clearance is entirely due to metablism by the liver and glomerular filtration in the kidneys, identical to clearance of three compartment steady state model.

## Usage

```
calc_total_clearance(
  chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
  parameters = NULL,
  species = "Human",
  suppress.messages = FALSE,
  default.to.human = FALSE,
  well.stirred.correction = TRUE,
  restrictive.clearance = TRUE,
  adjusted.Funbound.plasma = TRUE,
  ...
)
```

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#### **Arguments**

chem. cas Either the chemical name, CAS number, or the parameters must be specified.

chem. name Either the chemical name, CAS number, or the parameters must be specified.

dtxsid EPA's 'DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chemical

must be identified by either CAS, name, or DTXSIDs

parameters Chemical parameters from parameterize_steadystate function, overrides chem.name

and chem.cas.

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human"). suppress.messages

Whether or not the output message is suppressed.

default.to.human

Substitutes missing animal values with human values if true.

well.stirred.correction

Uses correction in calculation of hepatic clearance for well-stirred model if TRUE. This assumes clearance relative to amount unbound in whole blood instead of plasma, but converted to use with plasma concentration.

restrictive.clearance

Protein binding is not taken into account (set to 1) in liver clearance if FALSE.

adjusted.Funbound.plasma

Uses adjusted Funbound.plasma when set to TRUE.

... Additional parameters passed to parameterize_steadystate if parameters is NULL.

#### Value

Total Clearance

Units of L/h/kg BW.

## Author(s)

John Wambaugh

## **Examples**

calc_total_clearance(chem.name="Ibuprofen")

calc_vdist

Calculate the volume of distribution for a one compartment model.

## **Description**

This function predicts partition coefficients for all tissues, then lumps them into a single compartment.

calc_vdist 69

### Usage

```
calc_vdist(
  chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
  parameters = NULL,
  default.to.human = FALSE,
  species = "Human",
  suppress.messages = FALSE,
  adjusted.Funbound.plasma = TRUE,
  regression = TRUE,
  minimum.Funbound.plasma = 1e-04
)
```

### **Arguments**

chem.cas Either the CAS number or the chemical name must be specified when Fun-

bound.plasma is not given in parameter list.

chem.name Either the chemical name or the CAS number must be specified when Fun-

bound.plasma is not given in parameter list.

dtxsid EPA's DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chem-

ical must be identified by either CAS, name, or DTXSIDs

parameters Parameters from parameterize_3comp, parameterize_pbtk or predict_partitioning_schmitt.

default.to.human

Substitutes missing animal values with human values if true.

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

suppress.messages

Whether or not the output message is suppressed.

adjusted.Funbound.plasma

Uses adjusted Funbound.plasma when set to TRUE along with parition coeffi-

cients calculated with this value.

regression Whether or not to use the regressions in calculating partition coefficients.

minimum.Funbound.plasma

Monte Carlo draws less than this value are set equal to this value (default is 0.0001 – half the lowest measured Fup in our dataset).

# Details

The effective volume of distribution is calculated by summing each tissues volume times it's partition coefficient relative to plasma. Plasma, and the paritioning into RBCs are also added to get the total volume of distribution in L/KG BW. Partition coefficients are calculated using Schmitt's (2008) method. When species is specified as rabbit, dog, or mouse, the function uses the appropriate physiological data(volumes and flows) but substitues human fraction unbound, partition coefficients, and intrinsic hepatic clearance.

## Value

```
Volume of distribution \label{eq:Units} Units \ of \ L/ \ kg \ BW.
```

70 CAS.checksum

#### Author(s)

John Wambaugh and Robert Pearce

#### References

Schmitt W. "General approach for the calculation of tissue to plasma partition coefficients." Toxicology In Vitro, 22, 457-467 (2008). Peyret, T., Poulin, P., Krishnan, K., "A unified algorithm for predicting partition coefficients for PBPK modeling of drugs and environmental chemicals." Toxicology and Applied Pharmacology, 249, 197-207 (2010).

## **Examples**

```
calc_vdist(chem.cas="80-05-7")
calc_vdist(chem.name="Bisphenol A")
calc_vdist(chem.name="Bisphenol A",species="Rat")
```

CAS.checksum

Test the check digit of a CAS number to confirm validity

## **Description**

Chemical abstracts services registry numbers (CAS-RN) include a final digit as a "checksum" to test for validity (that is, that the number has not been corrupted).

# Usage

```
CAS.checksum(CAS.string)
```

### **Arguments**

CAS.string

A character string of three numbers separated by two dashes

## **Details**

The check digit (final number) is calculated by working from right to left, starting with the second to last digit of the CAS-RN. We multiply each digit by an increasing digit (1, 2, 3...) and sum as we work from right to left. The check digit should equal the final digit of the sum.

## Value

logical (TRUE if final digit of CAS is consistent with other digits)

# Author(s)

John Wambaugh

chem.invivo.PK.aggregate.data

Parameter Estimates from Wambaugh et al. (2018)

# Description

This table includes 1 and 2 compartment fits of plasma concentration vs time data aggregated from chem.invivo.PK.data, performed in Wambaugh et al. 2018. Data includes volume of distribution (Vdist, L/kg), elimination rate (kelim, 1/h), gut absorption rate (kgutabs, 1/h), fraction absorbed (Fgutabs), and steady state concentration (Css, mg/L).

# Usage

chem.invivo.PK.aggregate.data

#### **Format**

data.frame

### Author(s)

John Wambaugh

#### **Source**

Wambaugh et al. 2018 Toxicological Sciences, in press

chem.invivo.PK.data

Published toxicokinetic time course measurements

### **Description**

This data set includes time and dose specific measurements of chemical concentration in tissues taken from animals administered control doses of the chemicals either orally or intravenously. This plasma concentration-time data is from rat experiments reported in public sources. Toxicokinetic data were retrieved from those studies by the Netherlands Organisation for Applied Scientific Research (TNO) using curve stripping (TechDig v2). This data is provided for statistical analysis as in Wambaugh et al. 2018.

## Usage

chem.invivo.PK.data

#### **Format**

A data.frame containing 597 rows and 13 columns.

### Author(s)

Sieto Bosgra

72 chem.invivo.PK.data

#### Source

Wambaugh et al. 2018 Toxicological Sciences, in press

#### References

Aanderud L, Bakke OM (1983). Pharmacokinetics of antipyrine, paracetamol, and morphine in rat at 71 ATA. Undersea Biomed Res. 10(3):193-201. PMID: 6636344

Aasmoe L, Mathiesen M, Sager G (1999). Elimination of methoxyacetic acid and ethoxyacetic acid in rat. Xenobiotica. 29(4):417-24. PMID: 10375010

Ako RA. Pharmacokinetics/pharmacodynamics (PK/PD) of oral diethylstilbestrol (DES) in recurrent prostate cancer patients and of oral dissolving film (ODF)-DES in rats. PhD dissertation, College of Pharmacy, University of Houston, USA, 2011.

Anadon A, Martinez-Larranaga MR, Fernandez-Cruz ML, Diaz MJ, Fernandez MC, Martinez MA (1996). Toxicokinetics of deltamethrin and its 4'-HO-metabolite in the rat. Toxicol Appl Pharmacol. 141(1):8-16. PMID: 8917670

Binkerd PE, Rowland JM, Nau H, Hendrickx AG (1988). Evaluation of valproic acid (VPA) developmental toxicity and pharmacokinetics in Sprague-Dawley rats. Fundam Appl Toxicol. 11(3):485-93. PMID: 3146521

Boralli VB, Coelho EB, Cerqueira PM, Lanchote VL (2005). Stereoselective analysis of metoprolol and its metabolites in rat plasma with application to oxidative metabolism. J Chromatogr B Analyt Technol Biomed Life Sci. 823(2):195-202. PMID: 16029965

Chan MP, Morisawa S, Nakayama A, Kawamoto Y, Sugimoto M, Yoneda M (2005). Toxicokinetics of 14C-endosulfan in male Sprague-Dawley rats following oral administration of single or repeated doses. Environ Toxicol. 20(5):533-41. PMID: 16161119

Cruz L, Castaneda-Hernandez G, Flores-Murrieta FJ, Garcia-Lopez P, Guizar-Sahagun G (2002). Alteration of phenacetin pharmacokinetics after experimental spinal cord injury. Proc West Pharmacol Soc. 45:4-5. PMID: 12434508

Della Paschoa OE, Mandema JW, Voskuyl RA, Danhof M (1998). Pharmacokinetic-pharmacodynamic modeling of the anticonvulsant and electroencephalogram effects of phenytoin in rats. J Pharmacol Exp Ther. 284(2):460-6. PMID: 9454785

Du B, Li X, Yu Q, A Y, Chen C (2010). Pharmacokinetic comparison of orally disintegrating, beta-cyclodextrin inclusion complex and conventional tablets of nicardipine in rats. Life Sci J. 7(2):80-4.

Farris FF, Dedrick RL, Allen PV, Smith JC (1993). Physiological model for the pharmacokinetics of methyl mercury in the growing rat. Toxicol Appl Pharmacol. 119(1):74-90. PMID: 8470126

Hays SM, Elswick BA, Blumenthal GM, Welsch F, Conolly RB, Gargas ML (2000). Development of a physiologically based pharmacokinetic model of 2-methoxyethanol and 2-methoxyacetic acid disposition in pregnant rats. Toxicol Appl Pharmacol. 163(1):67-74. PMID: 10662606

Igari Y, Sugiyama Y, Awazu S, Hanano M (1982). Comparative physiologically based pharmacokinetics of hexobarbital, phenobarbital and thiopental in the rat. J Pharmacokinet Biopharm. 10(1):53-75. PMID: 7069578

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Jia L, Wong H, Wang Y, Garza M, Weitman SD (2003). Carbendazim: disposition, cellular permeability, metabolite identification, and pharmacokinetic comparison with its nanoparticle. J Pharm Sci. 92(1):161-72. PMID: 12486692

Kawai R, Mathew D, Tanaka C, Rowland M (1998). Physiologically based pharmacokinetics of cyclosporine A: extension to tissue distribution kinetics in rats and scale-up to human. J Pharmacol Exp Ther. 287(2):457-68. PMID: 9808668

Kim YC, Kang HE, Lee MG (2008). Pharmacokinetics of phenytoin and its metabolite, 4'-HPPH, after intravenous and oral administration of phenytoin to diabetic rats induced by alloxan or streptozotocin. Biopharm Drug Dispos. 29(1):51-61. PMID: 18022993

Kobayashi S, Takai K, Iga T, Hanano M (1991). Pharmacokinetic analysis of the disposition of valproate in pregnant rats. Drug Metab Dispos. 19(5):972-6. PMID: 1686245

Kotegawa T, Laurijssens BE, Von Moltke LL, Cotreau MM, Perloff MD, Venkatakrishnan K, Warrington JS, Granda BW, Harmatz JS, Greenblatt DJ (2002). In vitro, pharmacokinetic, and pharmacodynamic interactions of ketoconazole and midazolam in the rat. J Pharmacol Exp Ther. 302(3):1228-37. PMID: 12183684

Krug AK, Kolde R, Gaspar JA, Rempel E, Balmer NV, Meganathan K, Vojnits K, Baquie M, Waldmann T, Ensenat-Waser R, Jagtap S, Evans RM, Julien S, Peterson H, Zagoura D, Kadereit S, Gerhard D, Sotiriadou I, Heke M, Natarajan K, Henry M, Winkler J, Marchan R, Stoppini L, Bosgra S, Westerhout J, Verwei M, Vilo J, Kortenkamp A, Hescheler J, Hothorn L, Bremer S, van Thriel C, Krause KH, Hengstler JG, Rahnenfuhrer J, Leist M, Sachinidis A (2013). Human embryonic stem cell-derived test systems for developmental neurotoxicity: a transcriptomics approach. Arch Toxicol. 87(1):123-43. PMID: 23179753

Leon-Reyes MR, Castaneda-Hernandez G, Ortiz MI (2009). Pharmacokinetic of diclofenac in the presence and absence of glibenclamide in the rat. J Pharm Pharm Sci. 12(3):280-7. PMID: 20067705

Nagata M, Hidaka M, Sekiya H, Kawano Y, Yamasaki K, Okumura M, Arimori K (2007). Effects of pomegranate juice on human cytochrome P450 2C9 and tolbutamide pharmacokinetics in rats. Drug Metab Dispos. 35(2):302-5. PMID: 17132763

Okiyama M, Ueno K, Ohmori S, Igarashi T, Kitagawa H (1988). Drug interactions between imipramine and benzodiazepines in rats. J Pharm Sci. 77(1):56-63. PMID: 2894451

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Piersma AH, Bosgra S, van Duursen MB, Hermsen SA, Jonker LR, Kroese ED, van der Linden SC, Man H, Roelofs MJ, Schulpen SH, Schwarz M, Uibel F, van Vugt-Lussenburg BM, Westerhout J, Wolterbeek AP, van der Burg B (2013). Evaluation of an alternative in vitro test battery for detecting reproductive toxicants. Reprod Toxicol. 38:53-64. PMID: 23511061

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Satterwhite JH, Boudinot FD (1991). Effects of age and dose on the pharmacokinetics of ibuprofen in the rat. Drug Metab Dispos. 19(1):61-7. PMID: 1673423

Szymura-Oleksiak J, Panas M, Chrusciel W (1983). Pharmacokinetics of imipramine after single and multiple intravenous administration in rats. Pol J Pharmacol Pharm. 35(2):151-7. PMID: 6622297

Tanaka C, Kawai R, Rowland M (2000). Dose-dependent pharmacokinetics of cyclosporin A in rats: events in tissues. Drug Metab Dispos. 28(5):582-9. PMID: 10772639

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Yeung PK, Alcos A, Tang J (2009). Pharmacokinetics and Hemodynamic Effects of Diltiazem in Rats Following Single vs Multiple Doses In Vivo. Open Drug Metab J. 3:56-62.

chem.invivo.PK.summary.data

Summary of published toxicokinetic time course experiments

## Description

This data set summarizes the time course data in the chem.invivo.PK.data table. Maximum concentration (Cmax), time integrated plasma concentration for the duration of treatment (AUC.treatment) and extrapolated to zero concentration (AUC.infinity) as well as half-life are calculated. Summary values are given for each study and dosage. These data can be used to evaluate toxicokinetic model predictions.

# Usage

chem.invivo.PK.summary.data

## **Format**

A data.frame containing 100 rows and 25 columns.

### Author(s)

John Wambaugh

#### Source

Wambaugh et al. 2018 Toxicological Sciences, in press

#### References

Aanderud L, Bakke OM (1983). Pharmacokinetics of antipyrine, paracetamol, and morphine in rat at 71 ATA. Undersea Biomed Res. 10(3):193-201. PMID: 6636344

Aasmoe L, Mathiesen M, Sager G (1999). Elimination of methoxyacetic acid and ethoxyacetic acid in rat. Xenobiotica. 29(4):417-24. PMID: 10375010

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Anadon A, Martinez-Larranaga MR, Fernandez-Cruz ML, Diaz MJ, Fernandez MC, Martinez MA (1996). Toxicokinetics of deltamethrin and its 4'-HO-metabolite in the rat. Toxicol Appl Pharmacol. 141(1):8-16. PMID: 8917670

Binkerd PE, Rowland JM, Nau H, Hendrickx AG (1988). Evaluation of valproic acid (VPA) developmental toxicity and pharmacokinetics in Sprague-Dawley rats. Fundam Appl Toxicol. 11(3):485-93. PMID: 3146521

Boralli VB, Coelho EB, Cerqueira PM, Lanchote VL (2005). Stereoselective analysis of metoprolol and its metabolites in rat plasma with application to oxidative metabolism. J Chromatogr B Analyt Technol Biomed Life Sci. 823(2):195-202. PMID: 16029965

Chan MP, Morisawa S, Nakayama A, Kawamoto Y, Sugimoto M, Yoneda M (2005). Toxicokinetics of 14C-endosulfan in male Sprague-Dawley rats following oral administration of single or repeated doses. Environ Toxicol. 20(5):533-41. PMID: 16161119

Cruz L, Castaneda-Hernandez G, Flores-Murrieta FJ, Garcia-Lopez P, Guizar-Sahagun G (2002). Alteration of phenacetin pharmacokinetics after experimental spinal cord injury. Proc West Pharmacol Soc. 45:4-5. PMID: 12434508

Della Paschoa OE, Mandema JW, Voskuyl RA, Danhof M (1998). Pharmacokinetic-pharmacodynamic modeling of the anticonvulsant and electroencephalogram effects of phenytoin in rats. J Pharmacol Exp Ther. 284(2):460-6. PMID: 9454785

Du B, Li X, Yu Q, A Y, Chen C (2010). Pharmacokinetic comparison of orally disintegrating, beta-cyclodextrin inclusion complex and conventional tablets of nicardipine in rats. Life Sci J. 7(2):80-4.

Farris FF, Dedrick RL, Allen PV, Smith JC (1993). Physiological model for the pharmacokinetics of methyl mercury in the growing rat. Toxicol Appl Pharmacol. 119(1):74-90. PMID: 8470126

Hays SM, Elswick BA, Blumenthal GM, Welsch F, Conolly RB, Gargas ML (2000). Development of a physiologically based pharmacokinetic model of 2-methoxyethanol and 2-methoxyacetic acid disposition in pregnant rats. Toxicol Appl Pharmacol. 163(1):67-74. PMID: 10662606

Igari Y, Sugiyama Y, Awazu S, Hanano M (1982). Comparative physiologically based pharmacokinetics of hexobarbital, phenobarbital and thiopental in the rat. J Pharmacokinet Biopharm. 10(1):53-75. PMID: 7069578

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chem.lists

Chemical membership in different research projects

# Description

A static list of lists identifying chemical membership in different research projects. While it is our intent to keep these lists up-to-date, the information here is only for convenience and should not be considered to be definitive.

# Usage

chem.lists

# **Format**

A list containing ten lists.

## Author(s)

John Wambaugh

Units

log10(mole/L)

#### References

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chem.physical_and_invitro.data

Physico-chemical properties and in vitro measurements for toxicokinetics

## **Description**

This data set contains the necessary information to make basic, high-throughput toxicokinetic (HTTK) predictions for compounds, including Funbound.plasma, molecular weight (g/mol), logP, logMA (membrane affinity), intrinsic clearance(uL/min/10^6 cells), and pKa. These data have been compiled from multiple sources, and can be used to parameterize a variety of toxicokinetic models. See variable EPA.ref for information on the reference EPA.

Description

## Usage

chem.physical_and_invitro.data

#### **Format**

Column Name

#' logWSol

A data frame containing 9411 rows and 54 columns.

Compound	The preferred name of the chemical compound	none
CAS	The preferred Chemical Abstracts Service Registry Number	none
CAS.Checksum	A logical indicating whether the CAS number is valid	none
DTXSID	DSSTox Structure ID (http://comptox.epa.gov/dashboard)	none
Formula	The proportions of atoms within the chemical compound	none
SMILES.desalt	The simplified molecular-input line-entry system structure	none
All.Compound.Names	All names of the chemical as they occured in the data	none
logHenry	The log10 Henry's law constant	log10(atmosphers
logHenry.Reference	Reference for Henry's law constant	
logP	The log10 octanol:water partition coefficient (PC)	log10 unitless rati
logP.Reference	Reference for logPow	
logPwa	The log10 water:air PC	log10 unitless rati
logPwa.Reference	Reference for logPwa	
logMA	The log10 phospholipid:water PC or "Membrane affinity"	unitless ratio
logMA.Reference	Reference for membrane affinity	

The log10 water solubility

uL/min/10⁶ hepa

unitless fraction

unitless fraction

unitless ratio

none

logWSol.Reference	Reference for logWsol

MP The chemical compound melting point degrees Celsius

MP.Reference Reference for melting point

MW The chemical compound molecular weight g/mol

MW.Reference Reference for molecular weight

pKa_Accept The hydrogen acceptor equilibria concentrations logarithm

pKa_Accept.Reference Reference for pKa_Accept

pKa_Donor The hydrogen acceptor equilibria concentrations logarithm

pKa_Donor.Reference Reference for pKa_Donor

All Species All species for which data were available none

DTXSID.Reference Reference for DTXSID

Formula.Reference Reference for chemical formulat

[SPECIES].Clint (Primary hepatocyte suspension) intrinsic hepatic clearance

[SPECIES].Clint.pValue Probability that there is no clearance observed.

[SPECIES].Clint.pValue.Ref Reference for Clint pValue

[SPECIES].Clint.Reference Reference for Clint

[SPECIES]. Figure Fraction of chemical absorbed from the gut

[SPECIES].Fgutabs.Reference Reference for Fgutabs

[SPECIES].Funbound.plasma Chemical fraction unbound in presence of plasma proteins

[SPECIES].Funbound.plasma.Ref Reference for Funbound.plasma

[SPECIES].Rblood2plasma Chemical concentration blood to plasma ratio

[SPECIES].Rblood2plasma.Ref Reference for Rblood2plasma SMILES.desalt.Reference" Reference for SMILES structure

Chemical.Class All classes to which the chemical has been assigned

#### **Details**

In some cases the rapid equilbrium dailysis method (Waters et al., 2008) fails to yield detectable concentrations for the free fraction of chemical. In those cases we assume the compound is highly bound (that is, Fup approaches zero). For some calculations (for example, steady-state plasma concentration) there is precendent (Rotroff et al., 2010) for using half the average limit of detection, that is 0.005. We do not recomend using other models where quantities like partition coefficients must be predicted using Fup. We also do not recomend including the value 0.005 in training sets for Fup predictive models.

**Note** that in some cases the **Funbound.plasma** and the **intrinsic clearance** are *provided as a series of numbers separated by commas*. These values are the result of Bayesian analysis and characterize a distribution: the first value is the median of the distribution, while the second and third values are the lower and upper 95th percentile (that is qunatile 2.5 and 97.5) respectively. For intrinsic clearance a fourth value indicating a p-value for a decrease is provided. Typically 4000 samples were used for the Bayesian analysis, such that a p-value of "0" is equivale to "<0.00025". See Wambaugh et al. (2019) for more details.

Any one chemical compound *may have multiple ionization equilibria* (see Strope et al., 2018) may both for donating or accepting a proton (and therefore changing charge state). If there are multiple equilibria of the same type (donor/accept])the are concatonated by commas.

All species-specific information is initially from experimental measurements. The functions load_sipes2017, load_pradeep2020, and load_dawson2021 may be used to add in silico, structure-based predictions for many thousands of additional compounds to this table.

## Author(s)

John Wambaugh

#### Source

Wambaugh, John F., et al. "Toxicokinetic triage for environmental chemicals." Toxicological Sciences (2015): 228-237.

#### References

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ckd_epi_eq

CKD-EPI equation for GFR.

# Description

Predict GFR from serum creatinine, gender, and age.

### **Usage**

```
ckd_epi_eq(scr, gender, reth, age_years, ckd_epi_race_coeff = FALSE)
```

# **Arguments**

scr Vector of serum creatinine values in mg/dL. gender Vector of genders (either 'Male' or 'Female').

reth Vector of races/ethnicities. Not used unless ckd_epi_race_coeff is TRUE.

age_years Vector of ages in years.

ckd_epi_race_coeff

Whether to use the "race coefficient" in the CKD-EPI equation. Default is FALSE.

## **Details**

From Levey AS, Stevens LA, Schmid CH, Zhang YL, Castro AF, Feldman HI, et al. A new equation to estimate glomerular filtration rate. Ann Intern Med 2009; 150(9):604-612. doi:10.7326/0003-4819-150-9-200905050-00006

# Value

Vector of GFR values in mL/min/1.73m².

## Author(s)

Caroline Ring

## References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

concentration_data_Linakis2020

Concentration data involved in Linakis 2020 vignette analysis.

# Description

Concentration data involved in Linakis 2020 vignette analysis.

# Usage

concentration_data_Linakis2020

## **Format**

A data.frame containing x rows and y columns.

# Author(s)

Matt Linakis

# Source

Matt Linakis

## References

DSStox database (https://www.epa.gov/ncct/dsstox

# **Description**

Converts HTTK-Pop physiology into parameters relevant to the one compartment model

## Usage

```
convert_httkpop_1comp(parameters.dt, httkpop.dt, ...)
```

# Arguments

```
parameters.dt Data table returned by create_mc_samples

httkpop.dt Data table returned by httkpop_generate

... Additional arguments passed to propagate_invitrouv_1comp
```

## Value

A data.table whose columns are the parameters of the HTTK model specified in model.

## Author(s)

Caroline Ring, John Wambaugh, and Greg Honda

### References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

# Description

This function is designed to convert compartment values estimated from one of the HTTK models (e.g. "1compartment) using the solve_model function. It takes the HTTK model output matrix, model name, desired output units, and compound information to perform the conversion default model units to user specified units.

84 convert_solve_x

### Usage

```
convert_solve_x(
  model.output.mat,
  model = NULL,
  output.units = NULL,
  MW = NULL,
  vol = NULL,
  chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
  parameters = NULL,
  monitor.vars = NULL,
  suppress.messages = FALSE,
  verbose = FALSE,
  ...
)
```

# **Arguments**

model.output.mat

Matrix of results from HTTK solve_model function.

model Specified model to use in simulation: "pbtk", "3compartment", "3compartmentss",

"1compartment", "schmitt", ...

output units Output units of interest for the compiled components. Defaults to NULL, and

will provide values in model units if unspecified.

MW Molecular weight of substance of interest in g/mole

vol Volume for the target tissue of interest in liters (L). NOTE: Volume should not

be in units of per BW, i.e. "kg".

chem. cas Either the chemical name, CAS number, or the parameters must be specified.

chem. name Either the chemical name, CAS number, or the parameters must be specified.

dtxsid EPA's DSSTox Structure ID . (http://comptox.epa.gov/dashboard) the chem-

ical must be identified by either CAS, name, or DTXSIDs.

parameters A set of model parameters, especially a set that includes MW (molecular weight)

for our conversions.

monitor.vars A vector of character strings indicating the model component variables to re-

tain in the conversion factor table (assuming suppress.messages == FALSE). It should also be noted this option does NOT exclude columns from the input matrix provided in the 'model.output.mat' parameter. (Default is NULL, i.e. conversion factors for all model components are included in the reporting ma-

trix.)

suppress.messages

Whether or not the output messages are suppressed. (Default is FALSE, i.e.

show messages.)

verbose Whether or not to display the full conversion factor table. (Default is FALSE,

i.e. only include rows where the conversion factor is 1.)

... Other parameters that can be passed to convert_units, e.g. temperature and

compound state. See details in convert_units.

convert_units 85

#### **Details**

The function can be used to convert all compartments to a single unit, only units for a single model compartment, or units for a set of model compartments.

More details on the unit conversion can be found in the documentation for convert_units.

## Value

'new.ouput.matrix' A matrix with a column for time (in days), each compartment, and the area under the curve (AUC) and a row for each time point. The compartment and AUC columns are converted from model specified units to user specified units.

'output.units.vector' A vector of character strings providing the model compartments and their corresponding units after convert_solve_x.

## Author(s)

Sarah E. Davidson

## See Also

convert_units

# **Examples**

convert_units

convert units

# **Description**

This function is designed to accept input units, output units, and the molecular weight (MW) of a substance of interest to then use a table lookup to return a scaling factor that can be readily applied for the intended conversion. It can also take chemical identifiers in the place of a specified molecular weight value to retrieve that value for its own use.

## Usage

```
convert_units(
  input.units = NULL,
  output.units = NULL,
  MW = NULL,
  vol = NULL,
  chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
  parameters = NULL,
```

86 convert_units

```
temp = 25,
  state = "liquid"
)
```

### **Arguments**

input.units Assigned input units of interest

MW Molecular weight of substance of interest in g/mole

vol Volume for the target tissue of interest in liters (L). NOTE: Volume should not

be in units of per BW, i.e. "kg".

chem. cas Either the chemical name, CAS number, or the parameters must be specified.

chem. name Either the chemical name, CAS number, or the parameters must be specified.

dtxsid EPA's DSSTox Structure ID (http://comptox.epa.gov/dashboard) the chem-

ical must be identified by either CAS, name, or DTXSIDs

parameters A set of model parameters, especially a set that includes MW (molecular weight)

for our conversions

temp Temperature for conversions (default = 25 degreees C)

state Chemical state (gas or default liquid)

#### **Details**

If input or output units not contained in the table are queried, it gives a corresponding error message. It gives a warning message about the handling of 'ppmv,' as the function is only set up to convert between ppmv and mass-based units (like  $mg/m^3$  or umol/L) in the context of ideal gases.

convert_units is not directly configured to accept and convert units based on BW, like mg/kg. For this purpose, see scale_dosing.

The function supports a limited set of most relevant units across toxicological models, currently including umol, uM, mg, mg/L, mg/ $m^3$  or umol/L), and in the context of gases assumed to be ideal, ppmv.

Andersen and Clewell's Rules of PBPK Modeling:

- 1Check Your Units
- 2Check Your Units
- 3Check Mass Balance

### Author(s)

Mark Sfeir, John Wambaugh, and Sarah E. Davidson

## **Examples**

```
# MW BPA is 228.29 g/mol
# 1 mg/L -> 1/228.29*1000 = 4.38 uM
convert_units("mg/L","uM",chem.cas="80-05-7")

# MW Diclofenac is 296.148 g/mol
# 1 uM -> 296.148/1000 = 0.296
convert_units("uM","mg/L",chem.name="diclofenac")
```

create_mc_samples 87

create_mc_samples

Create a data table of draws of parameter values for Monte Carlo

### **Description**

This function creates a data table of draws of parameter values for use with Monte Carlo methods

## Usage

```
create_mc_samples(
 chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
 parameters = NULL,
  samples = 1000,
  species = "Human",
  suppress.messages = FALSE,
 model = "3compartmentss",
 httkpop = TRUE,
  invitrouv = TRUE,
  calcrb2p = TRUE,
  censored.params = list(),
  vary.params = list(),
  return.samples = FALSE,
  tissue = NULL,
 httkpop.dt = NULL,
  invitro.mc.arg.list = list(adjusted.Funbound.plasma = TRUE, poormetab = TRUE,
  fup.censored.dist = FALSE, fup.lod = 0.01, fup.meas.cv = 0.4, clint.meas.cv = 0.3,
    fup.pop.cv = 0.3, clint.pop.cv = 0.3),
 httkpop.generate.arg.list = list(method = "direct resampling", gendernum = NULL,
   agelim_years = NULL, agelim_months = NULL, weight_category = c("Underweight",
   "Normal", "Overweight", "Obese"), gfr_category = c("Normal", "Kidney Disease",
    "Kidney Failure"), reths = c("Mexican American", "Other Hispanic",
    "Non-Hispanic White", "Non-Hispanic Black", "Other")),
  convert.httkpop.arg.list = list(),
 propagate.invitrouv.arg.list = list(),
 parameterize.arg.list = list(restrictive.clearance = TRUE, default.to.human = FALSE,
    clint.pvalue.threshold = 0.05, regression = TRUE)
)
```

88 create_mc_samples

## **Arguments**

chem.cas Chemical Abstract Services Registry Number (CAS-RN) – if parameters is not

specified then the chemical must be identified by either CAS, name, or DTXISD

chem.name Chemical name (spaces and capitalization ignored) – if parameters is not speci-

fied then the chemical must be identified by either CAS, name, or DTXISD

dtxsid EPA's DSSTox Structure ID (https://comptox.epa.gov/dashboard) – if pa-

rameters is not specified then the chemical must be identified by either CAS,

name, or DTXSIDs

parameters Parameters from the appropriate parameterization function for the model indi-

cated by argument model

samples Number of samples generated in calculating quantiles.

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

Species must be set to "Human" to run httkpop model.

suppress.messages

Whether or not to suppress output message.

model Model used in calculation: 'pbtk' for the multiple compartment model, '3compartment'

for the three compartment model, '3compartmentss' for the three compartment steady state model, and '1compartment' for one compartment model. This only applies when httkpop=TRUE and species="Human", otherwise '3compart-

mentss' is used.

httkpop Whether or not to use the Ring et al. (2017) "httkpop" population generator.

Species must be 'Human'.

invitrouv Logical to indicate whether to include in vitro parameters such as intrinsic hep-

atic clearance rate and fraction unbound in plasma in uncertainty and variability

analysis

calcrb2p Logical determining whether or not to recalculate the chemical ratio of blood to

plasma

censored.params

The parameters listed in censored params are sampled from a normal distribution that is censored for values less than the limit of detection (specified separately for each parameter). This argument should be a list of sub-lists. Each sublist is named for a parameter in "parameters" and contains two elements: "CV" (coefficient of variation) and "LOD" (limit of detection, below which parameter values are censored. New values are sampled with mean equal to the value in "parameters" and standard deviation equal to the mean times the CV. Censored values are sampled on a uniform distribution between 0 and the limit of detection. Not used with httkpop model.

of detection. Not used with fitting

vary.params The parameters listed in vary.params are sampled from a normal distribution that

is truncated at zero. This argument should be a list of coefficients of variation (CV) for the normal distribution. Each entry in the list is named for a parameter in "parameters". New values are sampled with mean equal to the value in "parameters" and standard deviation equal to the mean times the CV. Not used with

httkpop model.

return.samples Whether or not to return the vector containing the samples from the simulation

instead of the selected quantile.

tissue Desired steady state tissue conentration.

httkpop.dt A data table generated by httkpop_generate. This defaults to NULL, in which

case httkpop_generate is called to generate this table.

dawson2021 89

```
invitro.mc.arg.list
```

Additional parameters passed to invitro_mc.

httkpop.generate.arg.list

Additional parameters passed to httkpop_generate.

convert.httkpop.arg.list

Additional parameters passed to the convert_httkpop_* function for the model.

propagate.invitrouv.arg.list

Additional parameters passed to model's associated in vitro uncertainty and variability propagation function

parameterize.arg.list

Additional parameters passed to the parameterize_* function for the model.

## Value

A data table where each column corresponds to parameters needed for the specified model and each row represents a different Monte Carlo sample of parameter values.

### Author(s)

Caroline Ring, Robert Pearce, and John Wambaugh

#### References

Wambaugh, John F., et al. "Toxicokinetic triage for environmental chemicals." Toxicological Sciences 147.1 (2015): 55-67.

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment international 106 (2017): 105-118.

# **Examples**

```
sample_set = create_mc_samples(chem.name = 'bisphenol a')
```

dawson2021

Dawson et al. 2021 data

# **Description**

This table includes QSAR (Random Forest) model predicted values for unbound fraction plasma protein (fup) and intrinsic hepatic clearance (clint) for a subset of chemicals in the Tox21 library (see https://www.epa.gov/chemical-research/toxicology-testing-21st-century-tox21).

# Usage

dawson2021

# Format

data.frame

90 EPA.ref

## **Details**

Predictions were made with a set of Random Forest QSAR models, as reported in Dawson et al. (2021).

# Author(s)

Daniel E. Dawson

## Source

Dawson et al. 2021 Random Forest QSAR Model

## References

Dawson, Daniel E. et al. "Designing QSARs for parameters of high-throughput toxicokinetic models using open-source descriptors." Environmental Science & Technology____. (2021):_____.

EPA.ref

Reference for EPA Physico-Chemical Data

# Description

The physico-chemical data in the chem.phys_and_invitro.data table are obtained from EPA's Comptox Chemicals dashboard. This variable indicates the date the Dashboard was accessed.

# Usage

EPA.ref

# **Format**

An object of class character of length 1.

# Author(s)

John Wambaugh

## **Source**

https://comptox.epa.gov/dashboard

estimate_gfr 91

estimate_gfr

Predict GFR.

### **Description**

Predict GFR using CKD-EPI equation (for adults) or BSA-based equation (for children).

### Usage

```
estimate_gfr(gfrtmp.dt, gfr_resid_var = TRUE, ckd_epi_race_coeff = FALSE)
```

# Arguments

gfrtmp.dt A data.table with columns gender, reth, age_years, age_months, BSA_adj, serum_creat.

gfr_resid_var Logical value indicating whether or not to include residual variability when gen-

erating GFR values. (Default is TRUE.)

ckd_epi_race_coeff

Logical value indicating whether or not to use the "race coefficient" from the CKD-EPI equation when estimating GFR values. (Default is FALSE.)

#### **Details**

Add residual variability based on reported residuals for each equation.

## Value

The same data.table with a gfr_est column added, containing estimated GFR values.

# Author(s)

Caroline Ring

## References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

estimate_gfr_ped

Predict GFR in children.

# Description

BSA-based equation from Johnson et al. 2006, Clin Pharmacokinet 45(9) 931-56. Used in Wetmore et al. 2014.

## Usage

```
estimate_gfr_ped(BSA)
```

92 estimate_hematocrit

# **Arguments**

BSA

Vector of body surface areas in m².

## Value

Vector of GFRs in mL/min/1.73m².

## Author(s)

Caroline Ring

## References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

estimate_hematocrit

Predict hematocrit using smoothing spline.

# **Description**

Using precalculated smoothing splines on NHANES log hematocrit vs. age in months (and KDE residuals) by gender and race/ethnicity, generate hematocrit values for individuals specified by age, gender, and race/ethnicity.

## Usage

```
estimate_hematocrit(hcttmp_dt)
```

# **Arguments**

hcttmp_dt

A data.table with columns age_years, age_months, gender, reth.

# Value

The same data.table with a hematocrit column added.

# Author(s)

Caroline Ring

## References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

export_pbtk_jarnac 93

# **Description**

This function exports the multiple compartment PBTK model to a jarnac file.

# Usage

```
export_pbtk_jarnac(
  chem.cas = NULL,
  chem.name = NULL,
  species = "Human",
  initial.amounts = list(Agutlumen = 0),
  filename = "default.jan",
  digits = 4
)
```

## **Arguments**

chem.cas Either the chemical name or CAS number must be specified.

chem.name Either the chemical name or CAS number must be specified.

species Species desired (either "Rat", "Rabbit", "Dog", or default "Human").

initial.amounts

Must specify initial amounts in units of choice.

filename The name of the jarnac file containing the model.

digits Desired number of decimal places to round the parameters.

### **Details**

Compartments to enter into the initial.amounts list includes Agutlumen, Aart, Aven, Alung, Agut, Aliver, Akidney, and Arest.

When species is specified as rabbit, dog, or mouse, the function uses the appropriate physiological data(volumes and flows) but substitues human fraction unbound, partition coefficients, and intrinsic hepatic clearance.

## Value

Text containing a Jarnac language version of the PBTK model.

## Author(s)

Robert Pearce

# **Examples**

```
export_pbtk_jarnac(chem.name='Nicotine',initial.amounts=list(Agutlumen=1),filename='PBTKmodel.jan')
```

94 export_pbtk_sbml

export_pbtk_sbml

Export model to sbml.

# **Description**

This function exports the multiple compartment PBTK model to an sbml file.

# Usage

```
export_pbtk_sbml(
  chem.cas = NULL,
  chem.name = NULL,
  species = "Human",
  initial.amounts = list(Agutlumen = 0),
  filename = "default.xml",
  digits = 4
)
```

## **Arguments**

chem. cas Either the chemical name or CAS number must be specified.

chem. name Either the chemical name or CAS number must be specified.

species Species desired (either "Rat", "Rabbit", "Dog", or default "Human").

initial. amounts

Must specify initial amounts in units of choice.

filename The name of the jarnac file containing the model.

digits Desired number of decimal places to round the parameters.

### **Details**

Compartments to enter into the initial.amounts list includes Agutlumen, Aart, Aven, Alung, Agut, Aliver, Akidney, and Arest.

When species is specified as rabbit, dog, or mouse, the function uses the appropriate physiological data(volumes and flows) but substitues human fraction unbound, partition coefficients, and intrinsic hepatic clearance.

## Value

Text describing the PBTK model in SBML.

## Author(s)

Robert Pearce

# **Examples**

```
export_pbtk_sbml(chem.name='Nicotine',initial.amounts=list(Agutlumen=1),filename='PBTKmodel.xml')
```

fetalpcs 95

fetalpcs

Fetal Partition Coefficients

# **Description**

Partition coefficients were measured for tissues, including placenta, in vitro by Csanady et al. (2002) for Bisphenol A and Diadzen. Curley et al. (1969) measured the concentration of a variety of pesticides in the cord blood of newborns and in the tissues of infants that were stillborn.

## Usage

fetalpcs

#### **Format**

data.frame

#### **Details**

Three of the chemicals studied by Curley et al. (1969) were modeled by Weijs et al. (2013) using the same partition coefficients for mother and fetus. The values used represented "prior knowledge" summarizing the available literature.

#### Source

Kapraun et al. 2021 (submitted)

## References

Csanady G, Oberste-Frielinghaus H, Semder B, Baur C, Schneider K, Filser J (2002). "Distribution and unspecific protein binding of the xenoestrogens bisphenol A and daidzein." *Archives of toxicology*, **76**(5-6), 299–305. Curley A, Copeland MF, Kimbrough RD (1969). "Chlorinated hydrocarbon insecticides in organs of stillborn and blood of newborn babies." *Archives of Environmental Health: An International Journal*, **19**(5), 628–632. Weijs L, Yang RS, Das K, Covaci A, Blust R (2013). "Application of Bayesian population physiologically based pharmacokinetic (PBPK) modeling and Markov chain Monte Carlo simulations to pesticide kinetics studies in protected marine mammals: DDT, DDE, and DDD in harbor porpoises." *Environmental science* \& technology, **47**(9), 4365–4374.

Frank2018invivo

Literature In Vivo Data on Doses Causing Neurological Effects

# **Description**

Studies were selected from Table 1 in Mundy et al., 2015, as the studies in that publication were cited as examples of compounds with evidence for developmental neurotoxicity. There were sufficient in vitro toxicokinetic data available for this package for only 6 of the 42 chemicals.

## Usage

Frank2018invivo

#### **Format**

A data.frame containing 14 rows and 16 columns.

# Author(s)

Timothy J. Shafer

### References

Frank, Christopher L., et al. "Defining toxicological tipping points in neuronal network development." Toxicology and Applied Pharmacology 354 (2018): 81-93.

Mundy, William R., et al. "Expanding the test set: Chemicals with potential to disrupt mammalian brain development." Neurotoxicology and Teratology 52 (2015): 25-35.

gen_age_height_weight Generate ages, heights, and weights for a virtual population using the virtual-individuals method.

# **Description**

Generate ages, heights, and weights for a virtual population using the virtual-individuals method.

# Usage

```
gen_age_height_weight(
  nsamp = NULL,
  gendernum = NULL,
  reths,
  weight_category,
  agelim_years,
  agelim_months
)
```

# **Arguments**

nsamp

The desired number of individuals in the virtual population. nsamp need not be provided if gendernum is provided.

gendernum

Optional: A named list giving the numbers of male and female individuals to include in the population, e.g. list(Male=100,Female=100). Default is NULL, meaning both males and females are included, in their proportions in the NHANES data. If both nsamp and gendernum are provided, they must agree (i.e., nsamp must be the sum of gendernum).

reths

Optional: a character vector giving the races/ethnicities to include in the population. Default is c('Mexican American','Other Hispanic','Non-Hispanic White','Non-Hispanic Black','Other'), to include all races and ethnicities in their proportions in the NHANES data. User-supplied vector must contain one or more of these strings.

weight_category

Optional: The weight categories to include in the population. Default is c('Underweight', 'Normal', 'Overweight', 'Obese'). User-supplied vector must contain one or more of these strings.

gen_height_weight 97

agelim_years Optional: A two-element numeric vector giving the minimum and maximum

ages (in years) to include in the population. Default is c(0,79). If agelim_years is provided and agelim_months is not, agelim_years will override the default

value of agelim_months.

agelim_months Optional: A two-element numeric vector giving the minimum and maximum

ages (in months) to include in the population. Default is c(0, 959), equivalent to the default agelim_years. If agelim_months is provided and agelim_years is not, agelim months will override the default values of agelim_years.

## Value

A data.table containing variables

gender Gender of each virtual individual
reth Race/ethnicity of each virtual individual

 ${\tt age_months} \ \ Age \ in \ months \ of \ each \ virtual \ individual$ 

age_years Age in years of each virtual individual

weight Body weight in kg of each virtual individual

height Height in cm of each virtual individual

## Author(s)

Caroline Ring

# References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118 importFrom survey svymean

gen_height_weight

Generate heights and weights for a virtual population.

# Description

Generate heights and weights for a virtual population.

### Usage

```
gen_height_weight(hbw_dt)
```

## **Arguments**

hbw_dt

A data.table describing the virtual population by race, gender, and age (in years and months). Must have variables gender, reth, age, and age.years.

## Value

The same data.table with two new variables added: weight and height. Respectively, these give individual body weights in kg, and individual heights in cm.

98 get_cheminfo

### Author(s)

Caroline Ring

#### References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

gen_serum_creatinine Predict GFR.

# **Description**

Predict serum creatinine using smoothing splines and kernel density estimates of residual variability

## Usage

gen_serum_creatinine(serumcreat.dt)

# **Arguments**

serumcreat.dt A data.table with columns gender, reth, age_years, age_months, BSA_adj.

# Value

The same data.table with a serum_creat column added, containing spline-interpolated serum creatinine values.

### Author(s)

Caroline Ring

## References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

get_cheminfo

Retrieve chemical information from HTTK package

## **Description**

This function provides the information specified in "info=" (can be single entry or vector) for all chemicals for which a toxicokinetic model can be parameterized for a given species. Since different models have different requirements and not all chemicals have complete data, this function will return different number of chemicals depending on the model specified.

get_cheminfo 99

### **Usage**

```
get_cheminfo(
  info = "CAS",
  species = "Human",
  fup.lod.default = 0.005,
  model = "3compartmentss",
  default.to.human = FALSE,
  median.only = FALSE,
  fup.ci.cutoff = TRUE,
  clint.pvalue.threshold = 0.05,
  suppress.messages = FALSE
)
```

## **Arguments**

info A single character vector (or collection of character vectors) from "Compound",

"CAS", "DTXSID, "logP", "pKa_Donor", "pKa_Accept", "MW", "Clint", "Clint.pValue",

"Funbound.plasma", "Structure_Formula", or "Substance_Type". info="all" gives

all information for the model and species.

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

fup.lod.default

Default value used for fraction of unbound plasma for chemicals where mea-

sured value was below the limit of detection. Default value is 0.0005.

model Model used in calculation, 'pbtk' for the multiple compartment model, '1com-

partment' for the one compartment model, '3compartment' for three compartment model, '3compartmentss' for the three compartment model without partition coefficients, or 'schmitt' for chemicals with logP and fraction unbound

(used in predict_partitioning_schmitt).

default.to.human

Substitutes missing values with human values if true.

median.only Use median values only for fup and clint. Default is FALSE.

fup.ci.cutoff Cutoff for the level of uncertainty in fup estimates. This value should be between

(0,1). Default is 'NULL' specifying no filtering.

clint.pvalue.threshold

Hepatic clearance for chemicals where the in vitro clearance assay result has a

p-values greater than the threshold are set to zero.

suppress.messages

Whether or not the output messages are suppressed.

# **Details**

When default.to.human is set to TRUE, and the species-specific data, Funbound.plasma and Clint, are missing from chem.physical_and_invitro.data, human values are given instead.

In some cases the rapid equilbrium dailysis method (Waters et al., 2008) fails to yield detectable concentrations for the free fraction of chemical. In those cases we assume the compound is highly bound (that is, Fup approaches zero). For some calculations (for example, steady-state plasma concentration) there is precendent (Rotroff et al., 2010) for using half the average limit of detection, that is, 0.005 (this value is configurable via the argument fup.lod.default). We do not recomend using other models where quantities like partition coefficients must be predicted using Fup. We also do not recomend including the value 0.005 in training sets for Fup predictive models.

100 get_cheminfo

**Note** that in some cases the **Funbound.plasma** and the **intrinsic clearance** are *provided as a series of numbers separated by commas*. These values are the result of Bayesian analysis and characterize a distribution: the first value is the median of the distribution, while the second and third values are the lower and upper 95th percentile (that is qunatile 2.5 and 97.5) respectively. For intrinsic clearance a fourth value indicating a p-value for a decrease is provided. Typically 4000 samples were used for the Bayesian analysis, such that a p-value of "0" is equivale to "<0.00025". See Wambaugh et al. (2019) for more details. If argument meadian.only == TRUE then only the median is reported for parameters with Bayesian analysis distributions. If the 95 credible interval is larger than fup.ci.cutoff (defaults to NULL) then the Fup is treated as too uncertain and the value NA is given.

#### Value

vector/data.table

[SPECIES].Rblood2plasma

Table (if info has multiple entries) or vector containing a column for each valid entry specified in the argument "info" and a row for each chemical with sufficient data for the model specified by argument "model":

Column	Description	units
Compound	The preferred name of the chemical compound	none
CAS	The preferred Chemical Abstracts Service Registry Number	none
DTXSID	DSSTox Structure ID (http://comptox.epa.gov/dashboard)	none
logP	The log10 octanol:water partition coefficient	log10 unitless ratio
MW	The chemical compound molecular weight	g/mol
pKa_Accept	The hydrogen acceptor equilibria concentrations	logarithm
pKa_Donor	The hydrogen donor equilibria concentrations	logarithm
[SPECIES].Clint	(Primary hepatocyte suspension) intrinsic hepatic clearance	uL/min/10^6 hepatoc
[SPECIES].Clint.pValue	Probability that there is no clearance observed.	none
[SPECIES].Funbound.plasma	Chemical fraction unbound in presence of plasma proteins	unitless fraction

unitless ratio

Chemical concentration blood to plasma ratio

# Author(s)

John Wambaugh, Robert Pearce, and Sarah E. Davidson

## References

Rotroff, Daniel M., et al. "Incorporating human dosimetry and exposure into high-throughput in vitro toxicity screening." Toxicological Sciences 117.2 (2010): 348-358.

Waters, Nigel J., et al. "Validation of a rapid equilibrium dialysis approach for the measurement of plasma protein binding." Journal of pharmaceutical sciences 97.10 (2008): 4586-4595.

Wambaugh, John F., et al. "Assessing toxicokinetic uncertainty and variability in risk prioritization." Toxicological Sciences 172.2 (2019): 235-251.

# **Examples**

```
# List all CAS numbers for which the 3compartmentss model can be run in humans:
get_cheminfo()

get_cheminfo(info=c('compound','funbound.plasma','logP'),model='pbtk')
# See all the data for humans:
```

get_chem_id 101

```
TPO.cas <- c("741-58-2", "333-41-5", "51707-55-2", "30560-19-1", "5598-13-0",
"35575-96-3", "142459-58-3", "1634-78-2", "161326-34-7", "133-07-3", "533-74-4",
"101-05-3", "330-54-1", "6153-64-6", "15299-99-7", "87-90-1", "42509-80-8",
"10265-92-6", "122-14-5", "12427-38-2", "83-79-4", "55-38-9", "2310-17-0",
"5234-68-4", \ "330-55-2", \ "3337-71-1", \ "6923-22-4", \ "23564-05-8", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0", \ "101-02-0
"140-56-7", "120-71-8", "120-12-7", "123-31-9", "91-53-2", "131807-57-3",
"68157-60-8", "5598-15-2", "115-32-2", "298-00-0", "60-51-5", "23031-36-9"
"137-26-8", "96-45-7", "16672-87-0", "709-98-8", "149877-41-8", "145701-21-9",
"7786-34-7", "54593-83-8", "23422-53-9", "56-38-2", "41198-08-7", "50-65-7",
"28434-00-6", "56-72-4", "62-73-7", "6317-18-6", "96182-53-5", "87-86-5", "101-54-2", "121-69-7", "532-27-4", "91-59-8", "105-67-9", "90-04-0",
"134-20-3", "599-64-4", "148-24-3", "2416-94-6", "121-79-9", "527-60-6",
"99-97-8", "131-55-5", "105-87-3", "136-77-6", "1401-55-4", "1948-33-0", "121-00-6", "92-84-2", "140-66-9", "99-71-8", "150-13-0", "80-46-6", "120-95-6",
"128-39-2", "2687-25-4", "732-11-6", "5392-40-5", "80-05-7", "135158-54-2",
"29232-93-7", "6734-80-1", "98-54-4", "97-53-0", "96-76-4", "118-71-8",
"2451-62-9", "150-68-5", "732-26-3", "99-59-2", "59-30-3", "3811-73-2",
"101-61-1", "4180-23-8", "101-80-4", "86-50-0", "2687-96-9", "108-46-3",
"95-54-5", "101-77-9", "95-80-7", "420-04-2", "60-54-8", "375-95-1", "120-80-9",
"149-30-4", "135-19-3", "88-58-4", "84-16-2", "6381-77-7", "1478-61-1",
"96-70-8", "128-04-1", "25956-17-6", "92-52-4", "1987-50-4", "563-12-2",
"298-02-2", "79902-63-9", "27955-94-8")
httk.TPO.rat.table <- subset(get_cheminfo(info="all", species="rat"),</pre>
  CAS %in% TPO.cas)
httk.TPO.human.table <- subset(get_cheminfo(info="all",species="human"),</pre>
  CAS %in% TPO.cas)
```

get_chem_id

get_cheminfo(info="all")

Retrieve chemical identity from HTTK package

# Description

Given one of chem.name, chem.cas (Chemical Abstract Service Registry Number), or DTXSID (DSStox Substance Identifier https://comptox.epa.gov/dashboard) this function checks if the chemical is available and, if so, returns all three pieces of information.

### Usage

```
get_chem_id(chem.cas = NULL, chem.name = NULL, dtxsid = NULL)
```

## **Arguments**

chem.cas CAS regstry number
chem.name Chemical name

dtxsid DSSTox Substance identifier

102 get_gfr_category

## Value

A list containing the following chemical identifiers:

chem.cas CAS registry number

chem.name Name dtxsid DTXSID

# Author(s)

John Wambaugh and Robert Pearce

get_gfr_category

Categorize kidney function by GFR.

# Description

For adults: In general GFR > 60 is considered normal 15 < GFR < 60 is considered kidney disease GFR < 15 is considered kidney failure

# Usage

```
get_gfr_category(age_years, age_months, gfr_est)
```

# Arguments

age_years Vector of ages in years.

age_months Vector of ages in months.

gfr_est Vector of estimated GFR values in mL/min/1.73m^2.

### **Details**

These values can also be used for children 2 years old and greater (see PEDIATRICS IN REVIEW Vol. 29 No. 10 October 1, 2008 pp. 335-341 (doi: 10.1542/pir.29-10-335))

## Value

Vector of GFR categories: 'Normal', 'Kidney Disease', 'Kidney Failure'.

## Author(s)

Caroline Ring

# References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

get_invitroPK_param 103

get_invitroPK_param Retrieve data from chem.physical_and_invitro.data table

# **Description**

or fraction unbound in plasma) from the main HTTK data. This function looks for species-specific values.

# Usage

```
get_invitroPK_param(
  param,
  species,
  chem.name = NULL,
  chem.cas = NULL,
  dtxsid = NULL
```

# **Arguments**

The in vitro pharmacokinetic parameter needed.

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

chem. name Either the chemical name, CAS number, or the parameters must be specified.

chem. cas Either the chemical name, CAS number, or the parameters must be specified.

dtxsid EPA's DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chem-

ical must be identified by either CAS, name, or DTXSIDs

# Value

The value of the parameter, if found

## Author(s)

John Wambaugh and Robert Pearce

# Description

This function provides the information specified in "info=" for all chemicals with data from the Wetmore et al. (2012) and (2013) publications and other literature.

# Usage

```
get_lit_cheminfo(info = "CAS", species = "Human")
```

104 get_lit_css

#### **Arguments**

info A single character vector (or collection of character vectors) from "Compound",

"CAS", "MW", "Raw.Experimental.Percentage.Unbound", "Entered.Experimental.Percentage.Unbound",

"Fub", "source_PPB", "Renal_Clearance", "Met_Stab", "Met_Stab_entered", "r2",

"p.val", "Concentration..uM.", "Css_lower_5th_perc.mg.L.", "Css_median_perc.mg.L.", "Css_upper_95th_perc.mg.L.", "Css_lower_5th_perc.uM.", "Css_median_perc.uM.", "Css_upper_95th_perc.uM.", "Css_upper_95th_perc.

and "Species".

Species desired (either "Rat" or default "Human"). species

### Value

info Table/vector containing values specified in "info" for valid chemicals.

#### Author(s)

John Wambaugh

#### References

Wetmore, B.A., Wambaugh, J.F., Ferguson, S.S., Sochaski, M.A., Rotroff, D.M., Freeman, K., Clewell, H.J., Dix, D.H., Andersen, M.E., Houck, K.A., Allen, B., Judson, R.S., Sing, R., Kavlock, R.J., Richard, A.M., and Thomas, R.S., "Integration of Dosimetry, Exposure and High-Throughput Screening Data in Chemical Toxicity Assessment," Toxicological Sciences 125 157-174 (2012)

Wetmore, B.A., Wambaugh, J.F., Ferguson, S.S., Li, L., Clewell, H.J. III, Judson, R.S., Freeman, K., Bao, W, Sochaski, M.A., Chu T.-M., Black, M.B., Healy, E, Allen, B., Andersen M.E., Wolfinger, R.D., and Thomas R.S., "The Relative Impact of Incorporating Pharmacokinetics on Predicting in vivo Hazard and Mode-of-Action from High-Throughput in vitro Toxicity Assays" Toxicological Sciences, 132:327-346 (2013).

Wetmore, B. A., Wambaugh, J. F., Allen, B., Ferguson, S. S., Sochaski, M. A., Setzer, R. W., Houck, K. A., Strope, C. L., Cantwell, K., Judson, R. S., LeCluyse, E., Clewell, H.J. III, Thomas, R.S., and Andersen, M. E. (2015). "Incorporating High-Throughput Exposure Predictions with Dosimetry-Adjusted In Vitro Bioactivity to Inform Chemical Toxicity Testing" Toxicological Sciences, kfv171.

# **Examples**

```
get_lit_cheminfo()
get_lit_cheminfo(info=c('CAS','MW'))
```

get_lit_css

Get literature Css

# Description

This function retrives a steady-state plasma concentration as a result of infusion dosing from the Wetmore et al. (2012) and (2013) publications and other literature.

get_lit_css 105

# Usage

```
get_lit_css(
  chem.cas = NULL,
  chem.name = NULL,
  daily.dose = 1,
  which.quantile = 0.95,
  species = "Human",
  clearance.assay.conc = NULL,
  output.units = "mg/L",
  suppress.messages = FALSE
)
```

## **Arguments**

chem. cas Either the cas number or the chemical name must be specified.

chem. name Either the chemical name or the CAS number must be specified.

daily.dose Total daily dose infused in units of mg/kg BW/day. Defaults to 1 mg/kg/day. which.quantile Which quantile from the SimCYP Monte Carlo simulation is requested. Can be

a vector.

species Species desired (either "Rat" or default "Human").

clearance.assay.conc

Concentration of chemical used in measureing intrinsic clearance data, 1 or  $10\,$ 

uM.

output.units Returned units for function, defaults to mg/L but can also be uM (specify units

= "uM").

suppress.messages

Whether or not the output message is suppressed.

# Value

A numeric vector with the literature steady-state plasma concentration (1 mg/kg/day) for the requested quantiles

### Author(s)

John Wambaugh

### References

Wetmore, B.A., Wambaugh, J.F., Ferguson, S.S., Sochaski, M.A., Rotroff, D.M., Freeman, K., Clewell, H.J., Dix, D.H., Andersen, M.E., Houck, K.A., Allen, B., Judson, R.S., Sing, R., Kavlock, R.J., Richard, A.M., and Thomas, R.S., "Integration of Dosimetry, Exposure and High-Throughput Screening Data in Chemical Toxicity Assessment," Toxicological Sciences 125 157-174 (2012)

Wetmore, B.A., Wambaugh, J.F., Ferguson, S.S., Li, L., Clewell, H.J. III, Judson, R.S., Freeman, K., Bao, W, Sochaski, M.A., Chu T.-M., Black, M.B., Healy, E, Allen, B., Andersen M.E., Wolfinger, R.D., and Thomas R.S., "The Relative Impact of Incorporating Pharmacokinetics on Predicting in vivo Hazard and Mode-of-Action from High-Throughput in vitro Toxicity Assays" Toxicological Sciences, 132:327-346 (2013).

Wetmore, B. A., Wambaugh, J. F., Allen, B., Ferguson, S. S., Sochaski, M. A., Setzer, R. W., Houck, K. A., Strope, C. L., Cantwell, K., Judson, R. S., LeCluyse, E., Clewell, H.J. III, Thomas, R.S., and Andersen, M. E. (2015). "Incorporating High-Throughput Exposure Predictions with Dosimetry-Adjusted In Vitro Bioactivity to Inform Chemical Toxicity Testing" Toxicological Sciences, kfv171.

106 get_lit_oral_equiv

### **Examples**

```
get_lit_css(chem.cas="34256-82-1")
get_lit_css(chem.cas="34256-82-1", species="Rat", which.quantile=0.5)
get_lit_css(chem.cas="80-05-7", daily.dose = 1, which.quantile = 0.5, output.units = "uM")
```

get_lit_oral_equiv

Get Literature Oral Equivalent Dose

# **Description**

This function converts a chemical plasma concetration to an oral equivalent dose using the values from the Wetmore et al. (2012) and (2013) publications and other literature.

## Usage

```
get_lit_oral_equiv(
  conc,
  chem.name = NULL,
  chem.cas = NULL,
  suppress.messages = FALSE,
  which.quantile = 0.95,
  species = "Human",
  input.units = "uM",
  output.units = "mg",
  clearance.assay.conc = NULL,
  ...
)
```

# **Arguments**

conc Bioactive in vitro concentration in units of specified input.units, default of uM.

chem. name Either the chemical name or the CAS number must be specified. chem. cas Either the CAS number or the chemical name must be specified.

suppress.messages

Suppress output messages.

which quantile Which quantile from the SimCYP Monte Carlo simulation is requested. Can be

a vector. Papers include 0.05, 0.5, and 0.95 for humans and 0.5 for rats.

species Species desired (either "Rat" or default "Human").

input.units Units of given concentration, default of uM but can also be mg/L.

output.units Units of dose, default of 'mg' for mg/kg BW/ day or 'mol' for mol/kg BW/ day.

clearance.assay.conc

Concentration of chemical used in measureing intrinsic clearance data, 1 or 10

uM.

... Additional parameters passed to get_lit_css.

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#### Value

Equivalent dose in specified units, default of mg/kg BW/day.

### Author(s)

John Wambaugh

#### References

Wetmore, B.A., Wambaugh, J.F., Ferguson, S.S., Sochaski, M.A., Rotroff, D.M., Freeman, K., Clewell, H.J., Dix, D.H., Andersen, M.E., Houck, K.A., Allen, B., Judson, R.S., Sing, R., Kavlock, R.J., Richard, A.M., and Thomas, R.S., "Integration of Dosimetry, Exposure and High-Throughput Screening Data in Chemical Toxicity Assessment," Toxicological Sciences 125 157-174 (2012)

Wetmore, B.A., Wambaugh, J.F., Ferguson, S.S., Li, L., Clewell, H.J. III, Judson, R.S., Freeman, K., Bao, W, Sochaski, M.A., Chu T.-M., Black, M.B., Healy, E, Allen, B., Andersen M.E., Wolfinger, R.D., and Thomas R.S., "The Relative Impact of Incorporating Pharmacokinetics on Predicting in vivo Hazard and Mode-of-Action from High-Throughput in vitro Toxicity Assays" Toxicological Sciences, 132:327-346 (2013).

Wetmore, B. A., Wambaugh, J. F., Allen, B., Ferguson, S. S., Sochaski, M. A., Setzer, R. W., Houck, K. A., Strope, C. L., Cantwell, K., Judson, R. S., LeCluyse, E., Clewell, H.J. III, Thomas, R.S., and Andersen, M. E. (2015). "Incorporating High-Throughput Exposure Predictions with Dosimetry-Adjusted In Vitro Bioactivity to Inform Chemical Toxicity Testing" Toxicological Sciences, kfv171.

# **Examples**

## **Description**

This function retrieves physico-chemical properties ("param") for the chemical specified by chem.name or chem.cas from the vLiver tables.

# Usage

```
get_physchem_param(param, chem.name = NULL, chem.cas = NULL, dtxsid = NULL)
```

108 get_rblood2plasma

### **Arguments**

param The desired parameters, a vector or single value.

chem.name The chemical names that you want parameters for, a vector or single value

chem.cas The chemical CAS numbers that you want parameters for, a vector or single

value

dtxsid EPA's 'DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chemical

must be identified by either CAS, name, or DTXSIDs

## Value

The parameters, either a single value, a named list for a single chemical, or a list of lists

### Author(s)

John Wambaugh and Robert Pearce

## **Examples**

get_rblood2plasma

Get ratio of the blood concentration to the plasma concentration.

# **Description**

This function attempts to retrieve a measured species- and chemical-specific blood:plasma concentration ratio.

## Usage

```
get_rblood2plasma(
  chem.name = NULL,
  chem.cas = NULL,
  dtxsid = NULL,
  species = "Human",
  default.to.human = FALSE
)
```

## **Arguments**

chem. name Either the chemical name or the CAS number must be specified. chem. cas Either the CAS number or the chemical name must be specified.

dtxsid EPA's 'DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chemical

must be identified by either CAS, name, or DTXSIDs

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

default.to.human

Substitutes missing animal values with human values if true.

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#### **Details**

A value of NA is returned when the requested value is unavailable. Values are retrieved from chem.physical_and_invitro.data. details than the description above ~~

#### Value

A numeric value for the steady-state ratio of chemical concentration in blood to plasma

#### Author(s)

Robert Pearce

### **Examples**

```
get_rblood2plasma(chem.name="Bisphenol A")
get_rblood2plasma(chem.name="Bisphenol A",species="Rat")
```

get_weight_class

Given vectors of age, BMI, recumbent length, weight, and gender, categorizes weight classes using CDC and WHO categories.

# **Description**

Given vectors of age, BMI, recumbent length, weight, and gender, categorizes weight classes using CDC and WHO categories.

# Usage

```
get_weight_class(age_years, age_months, bmi, recumlen, weight, gender)
```

# **Arguments**

age_years A vector of ages in years.
age_months A vector of ages in months.

bmi A vector of BMIs.

recumlen A vector of heights or recumbent lengths in cm.

weight A vector of body weights in kg.

gender A vector of genders (as 'Male' or 'Female').

#### Value

A character vector of weight classes. Each element will be one of 'Underweight', 'Normal', 'Overweight', or 'Obese'.

#### Author(s)

Caroline Ring

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#### References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

hematocrit_infants

Predict hematocrit in infants under 1 year old.

# **Description**

For infants under 1 year, hematocrit was not measured in NHANES. Assume a log-normal distribution where plus/minus 1 standard deviation of the underlying normal distribution is given by the reference range. Draw hematocrit values from these distributions by age.

# Usage

```
hematocrit_infants(age_months)
```

# **Arguments**

age_months

Vector of ages in months; all must be <= 12.

#### **Details**

Age	Reference range
<1 month	31-49
1-6 months	29-42
7-12 months	33-38

#### Value

Vector of hematocrit percentages corresponding to the input vector of ages.

# Author(s)

Caroline Ring

#### References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

honda.ivive

Return the assumptions used in Honda et al. 2019

honda.ivive 111

#### **Description**

This function returns four of the better performing sets of assumptions evaluated in Honda et al. 2019 (https://doi.org/10.1371/journal.pone.0217564). These include four different combinations of hepatic clearance assumption, in vivo bioactivity assumption, and relevant tissue assumption. Generally, this function is not called directly by the user, but instead called by setting the IVIVE option in calc_mc_oral_equiv, calc_mc_css, and calc_analytic functions. Currently, these IVIVE option is not implemented the solve_1comp etc. functions.

#### Usage

```
honda.ivive(method = "Honda1", tissue = "liver")
```

#### **Arguments**

method This is set to one of "Honda1", "Honda2", "Honda3", or "Honda4".

tissue This is only relevant to "Honda4" and indicates the relevant tissue compartment.

#### **Details**

"Honda1" - tissue = NULL, restrictive.clearance = TRUE, bioactive.free.invivo = TRUE This assumption assumes restrictive hepatic clearance, and treats the free concentration in plasma as the bioactive concentration in vivo. This option must be used in combination with the concentration in vitro predicted by armitage_eval(), otherwise the result will be the same as "Honda2". This option corresponds to the result in Figure 8 panel c) restrictive, mean free plasma conc., Armitage in Honda et al. 2019. "Honda2" - tissue = NULL, restrictive.clearance = TRUE, bioactive.free.invivo = TRUE This assumption assumes restrictive hepatic clearance, and treats the free concentration in plasma as the bioactive concentration in vivo. This option corresponds to the result in Figure 8 panel b) restrictive, mean free plasma conc. in Honda et al. 2019. "Honda3" - tissue = NULL, restrictive.clearance = TRUE, bioactive.free.invivo = TRUE This assumption assumes restrictive hepatic clearance, and treats the free concentration in plasma as the bioactive concentration in vivo. This option corresponds to the result in Figure 8 panel a) restrictive, mean total plasma conc. in Honda et al. 2019. "Honda4" - tissue = tissue, restrictive.clearance = FALSE, bioactive.free.invivo = TRUE This assumption assumes restrictive hepatic clearance, and treats the free concentration in plasma as the bioactive concentration in vivo. The input tissue should be relevant to the in vitro assay endpoint used as input or that the result is being compared to. This option corresponds to the result in Figure 8 panel d) nonrestrictive, mean tissue conc. in Honda et al. 2019.

#### Value

A list of tissue, bioactive.free.invivo, and restrictive.clearance assumptions.

# Author(s)

Greg Honda and John Wambaugh

# References

Honda, Gregory S., et al. "Using the Concordance of In Vitro and In Vivo Data to Evaluate Extrapolation Assumptions." 2019. PLoS ONE 14(5): e0217564.

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#### **Examples**

```
honda.ivive(method = "Honda1", tissue = NULL)
```

howgate

Howgate 2006

#### Description

This data set is only used in Vignette 5.

#### Usage

howgate

#### **Format**

A data.table containing 24 rows and 11 columns.

### Author(s)

Caroline Ring

#### References

Howgate, E. M., et al. "Prediction of in vivo drug clearance from in vitro data. I: impact of interindividual variability." Xenobiotica 36.6 (2006): 473-497.

httkpop

httkpop: Virtual population generator for HTTK.

# Description

The httkpop package generates virtual population physiologies for use in population TK.

# **Details**

To simulate inter-individual variability in the TK model, a MC approach is used: the model parameters are sampled from known or assumed distributions, and the model is evaluated for each sampled set of parameters. To simulate variability across subpopulations, the MC approach needs to capture the parameter correlation structure. For example, kidney function changes with age (Levey et al., 2009), thus the distribution of GFR is likely different in 6-year-olds than in 65-yearolds. To directly measure the parameter correlation structure, all parameters need to be measured in each individual in a representative sample population. Such direct measurements are extremely limited. However, the correlation structure of the physiological parameters can be inferred from their known individual correlations with demographic and anthropometric quantities for which direct population measurements do exist. These quantities are sex, race/ethnicity, age, height, and weight (Howgate et al., 2006; Jamei et al., 2009a; Johnson et al., 2006; McNally et al., 2014; Price et al., 2003). Direct measurements of these quantities in a large, representative sample of the U.S. population are publicly available from NHANES. NHANES also includes laboratory measurements, including

both serum creatinine, which can be used to estimate GFR (Levey et al., 2009), and hematocrit. For conciseness, sex, race/ethnicity, age, height, weight, serum creatinine, and hematocrit will be called the NHANES quantities.

HTTK-Pop's correlated MC approach begins by sampling from the joint distribution of the NHANES quantities to simulate a population. Then, for each individual in the simulated population, HTTKe-Pop predicts the physiological parameters from the NHANES quantities using regression equations from the literature (Barter et al., 2007; Baxter-Jones et al., 2011; Bosgra et al., 2012; Koo et al., 2000; Levey et al., 2009; Looker et al., 2013; McNally et al., 2014; Ogiu et al., 1997; Price et al., 2003; Schwartz and Work, 2009; Webber and Barr 2012). Correlations among the physiological parameters are induced by their mutual dependence on the correlated NHANES quantities. Finally, residual variability is added to the predicted physiological parameters using estimates of residual marginal variance (i.e., variance not explained by the regressions on the NHANES quantities) (McNally et al., 2014).

Data were combined from the three most recent publicly-available NHANES cycles: 2007-2008, 2009-2010, and 2011-2012. For each cycle, some NHANES quantities - height, weight, serum creatinine, and hematocrit - were measured only in a subset of respondents. Only these subsets were included in HTTKePop. The pooled subsets from the three cycles contained 29,353 unique respondents. Some respondents were excluded from analysis: those with age recorded as 80 years (because all NHANES respondents 80 years and older were marked as "80"); those with missing height, weight or hematocrit data; and those aged 12 years or older with missing serum creatinine data. These criteria excluded 4807 respondents, leaving 24,546 unique respondents. Each NHANES respondent was assigned a cycle-specific sample weight, which can be interpreted as the number of individuals in the total U.S. population represented by each NHANES respondent in each cycle (Johnson et al., 2013). Because data from three cycles were combined, the sample weights were rescaled (divided by the number of cycles being combined, as recommended in NHANES data analysis documentation) (Johnson et al., 2013). To handle the complex NHANES sampling structure, the R survey package was used to analyze the NHANES data (Lumley, 2004).

To allow generation of virtual populations specified by weight class, we coded a categorical variable for each NHANES respondent. The categories Underweight, Normal, Overweight, or Obese were assigned based on weight, age, and height/length (Grummer-Strawn et al., 2010; Kuczmarski et al., 2002; Ogden et al., 2014; WHO, 2006, 2010). We implemented two population simulation methods within HTTK-Pop: the direct-resampling method and the virtual-individuals method. The direct-resampling method simulated a population by sampling NHANES respondents with replacement, with probabilities proportional to the sample weights. Each individual in the resulting simulated population was an NHANES respondent, identified by a unique NHANES sequence number. By contrast, the second method generates "virtual individuals" - sets of NHANES quantities that obey the approximate joint distribution of the NHANES quantities (calculated using weighted smoothing functions and kernel density estimators), but do not necessarily correspond to any particular NHANES respondent. The direct-resampling method removed the possibility of generating unrealistic combinations of the NHANES quantities; the virtual-individuals method allowed the use of interpolation to simulate subpopulations represented by only a small number of NHANES respondents.

For either method, HTTK-Pop takes optional specifications about the population to be simulated and then samples from the appropriate conditional joint distribution of the NHANES quantities.

Once HTTK-Pop has simulated a population characterized by the NHANES quantities, the physiological parameters of the TK model are predicted from the NHANES quantities using regression equations from the literature. Liver mass was predicted for individuals over age 18 using allometric scaling with height from Reference Man (Valentin, 2002), and for individuals under 18 using regression relationships with height and weight published by Ogiu et al. (1997). Residual marginal variability was added for each individual as in PopGen (McNally et al., 2014). Similarly, hepatic portal vein blood flows (in L/h) are predicted as fixed fractions of a cardiac output allometrically

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scaled with height from Reference Man (Valentin, 2002), and residual marginal variability is added for each individual (McNally et al., 2014). Glomerular filtration rate (GFR) (in L/h/1.73 m2 body surface area) is predicted from age, race, sex, and serum creatinine using the CKD-EPI equation, for individuals over age 18 (Levey et al., 2009). For individuals under age 18, GFR is estimated from body surface area (BSA) (Johnson et al., 2006); BSA is predicted using Mosteller's formula (Verbraecken et al., 2006) for adults and Haycock's formula (Haycock et al., 1978) for children. Hepatocellularity (in millions of cells per gram of liver tissue) is predicted from age using an equation developed by Barter et al. (2007). Hematocrit is estimated from NHANES data for individuals 1 year and older. For individuals younger than 1 year, for whom NHANES did not measure hematocrit directly, hematocrit was predicted from age in months, using published reference ranges (Lubin, 1987).

In addition to the HTTK physiological parameters, the HTTK models include chemical-specific parameters representing the fraction of chemical unbound in plasma (Fup) and intrinsic clearance (CLint). Because these parameters represent interactions of the chemical with the body, their values will vary between individuals. To simulate this variability, Fub and CLint were included in MC simulations, by sampling from estimated or assumed distributions for the parameters defining them.

Variability in hematocrit was simulated either using NHANES data (for individuals ages 1 and older) or using age-based reference ranges (for individuals under age 1). Fup was treated as a random variable obeying a distribution censored below the average limit of quantification (LOQ) of the in vitro assay. Specifically, Fup was assumed to obey a normal distribution truncated below at 0 and above at 1, centered at the Fup value measured in vitro, with a 30 the average LOQ (0.01), Fup was instead drawn from a uniform distribution between 0 and 0.01. Fup was assumed to be independent of all other parameters. This censored normal distribution was chosen to match that used in Wambaugh et al. (2015).

Variability in hepatocellularity (106 cells/g liver) and Mliver (kg) were simulated. The remaining source of variability in CLint, his variability in CLint, which was simulated using a Gaussian mixture distribution to represent the population proportions of poor metabolizers (PMs) and non-PMs of each substance. The true prevalence of PMs is isozyme-specific (Ma et al., 2002; Yasuda et al., 2008); however, isozyme-specific metabolism data were not available for the majority of chemicals considered. We therefore made a simplifying assumption that 5 slower than average. With 95 a normal distribution truncated below at zero, centered at the value measured in vitro, with a 30 CLint was drawn from a PM distribution: a truncated normal distribution centered on one-tenth of the in vitro value with 30 Both CLint itself and the probability of being a PM were assumed to be independent of all other parameters. The truncated normal nonePM distribution was chosen because it has been used (with 100 in previous work (Rotroff et al., 2010; Wambaugh et al., 2015; Wetmore et al., 2014; Wetmore et al., 2015; Wetmore et al., 2012); the PM distribution was chosen to comport with the nonePM distribution.

### Main function to generate a population

If you just want to generate a table of (chemical-independent) population physiology parameters, use httkpop_generate.

### **Using HTTK-Pop with HTTK**

To generate a population and then run an HTTK model for that population, the workflow is as follows:

- 1. Generate a population using httkpop_generate.
- 2. For a given HTTK chemical and general model, convert the population data to corresponding sets of HTTK model parameters using httkpop_mc.

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#### Author(s)

Caroline Ring

#### References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

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httkpop_biotophys_default

Convert HTTK-Pop-generated parameters to HTTK physiological parameters

#### **Description**

Convert HTTK-Pop-generated parameters to HTTK physiological parameters

### Usage

```
httkpop_biotophys_default(indiv_dt)
```

### **Arguments**

indiv_dt

The data.table object returned by httkpop_generate()

#### Value

A data.table with the physiological parameters expected by any HTTK model, including body weight (BW), hematocrit, tissue volumes per kg body weight, tissue flows as fraction of CO, CO per (kg BW)^3/4, GFR per (kg BW)^3/4, portal vein flow per (kg BW)^3/4, and liver density.

#### Author(s)

Caroline Ring

### References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

```
httkpop_direct_resample
```

Generate a virtual population by directly resampling the NHANES data

# Description

Generate a virtual population by directly resampling the NHANES data.

# Usage

```
httkpop_direct_resample(
  nsamp = NULL,
  gendernum = NULL,
  agelim_years = c(0, 79),
  agelim_months = c(0, 959),
  weight_category = c("Underweight", "Normal", "Overweight", "Obese"),
  gfr_category = c("Normal", "Kidney Disease", "Kidney Failure"),
```

```
reths = c("Mexican American", "Other Hispanic", "Non-Hispanic White",
    "Non-Hispanic Black", "Other"),
    gfr_resid_var = TRUE,
    ckd_epi_race_coeff = FALSE
)
```

# **Arguments**

nsamp The desired number of individuals in the virtual population. nsamp need not be

provided if gendernum is provided.

gendernum Optional: A named list giving the numbers of male and female individuals

to include in the population, e.g. <code>list(Male=100,Female=100)</code>. Default is NULL, meaning both males and females are included, in their proportions in the NHANES data. If both nsamp and gendernum are provided, they must agree

(i.e., nsamp must be the sum of gendernum).

agelim_years Optional: A two-element numeric vector giving the minimum and maximum

ages (in years) to include in the population. Default is c(0,79). If agelim_years is provided and agelim_months is not, agelim_years will override the default

value of agelim_months.

agelim_months Optional: A two-element numeric vector giving the minimum and maximum

ages (in months) to include in the population. Default is c(0, 959), equivalent to the default agelim_years. If agelim_months is provided and agelim_years is not, agelim_months will override the default values of agelim_years.

weight_category

Optional: The weight categories to include in the population. Default is c('Underweight',

'Normal', 'Overweight', 'Obese'). User-supplied vector must contain one

or more of these strings.

gfr_category The kidney function categories to include in the population. Default is c('Normal', 'Kidney

Disease', 'Kidney Failure') to include all kidney function levels.

reths Optional: a character vector giving the races/ethnicities to include in the popula-

tion. Default is c('Mexican American','Other Hispanic','Non-Hispanic White','Non-Hispanic Black','Other'), to include all races and ethnicities in their proportions in the NHANES data. User-supplied vector must contain

one or more of these strings.

gfr_resid_var Logical value indicating whether or not to include residual variability when gen-

erating GFR values. (Default is TRUE.)

ckd_epi_race_coeff

Logical value indicating whether or not to use the "race coefficient" from the CKD-EPI equation when estimating GFR values. (Default is FALSE.)

# Value

A data.table where each row represents an individual, and each column represents a demographic, anthropometric, or physiological parameter.

# Author(s)

Caroline Ring

#### References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

```
httkpop_direct_resample_inner
```

*Inner loop function called by* httkpop_direct_resample.

### **Description**

Inner loop function called by httkpop_direct_resample.

# Usage

```
httkpop_direct_resample_inner(
   nsamp,
   gendernum,
   agelim_months,
   agelim_years,
   reths,
   weight_category,
   gfr_resid_var,
   ckd_epi_race_coeff
)
```

# Arguments

nsamp

The desired number of individuals in the virtual population. nsamp need not be provided if gendernum is provided.

gendernum

Optional: A named list giving the numbers of male and female individuals to include in the population, e.g. list(Male=100,Female=100). Default is NULL, meaning both males and females are included, in their proportions in the NHANES data. If both nsamp and gendernum are provided, they must agree (i.e., nsamp must be the sum of gendernum).

agelim_months

Optional: A two-element numeric vector giving the minimum and maximum ages (in months) to include in the population. Default is c(0, 959), equivalent to the default agelim_years. If agelim_months is provided and agelim_years is not, agelim_months will override the default values of agelim_years.

agelim_years

Optional: A two-element numeric vector giving the minimum and maximum ages (in years) to include in the population. Default is c(0,79). If agelim_years is provided and agelim_months is not, agelim_years will override the default value of agelim_months.

reths

Optional: a character vector giving the races/ethnicities to include in the population. Default is c('Mexican American','Other Hispanic','Non-Hispanic White','Non-Hispanic Black','Other'), to include all races and ethnicities in their proportions in the NHANES data. User-supplied vector must contain one or more of these strings.

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```
weight_category
```

Optional: The weight categories to include in the population. Default is c('Underweight', 'Normal', 'Overweight', 'Obese'). User-supplied vector must contain one or more of these strings.

gfr_resid_var

Logical value indicating whether or not to include residual variability when generating GFR values. (Default is TRUE, passed from 'httkpop_direct_resample'.)

ckd_epi_race_coeff

Logical value indicating whether or not to use the "race coefficient" from the CKD-EPI equation when estimating GFR values. (Default is FALSE, passed from 'httkpop_direct_resample'.)

#### Value

A data.table where each row represents an individual, and each column represents a demographic, anthropometric, or physiological parameter.

#### Author(s)

Caroline Ring

#### References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

httkpop_generate

Generate a virtual population

#### Description

Generate a virtual population

# Usage

```
httkpop_generate(
  method = "direct resampling",
  nsamp = NULL,
  gendernum = NULL,
  agelim_years = NULL,
  agelim_months = NULL,
  weight_category = c("Underweight", "Normal", "Overweight", "Obese"),
  gfr_category = c("Normal", "Kidney Disease", "Kidney Failure"),
  reths = c("Mexican American", "Other Hispanic", "Non-Hispanic White",
        "Non-Hispanic Black", "Other"),
  gfr_resid_var = TRUE,
  ckd_epi_race_coeff = FALSE
)
```

httkpop_generate 121

#### **Arguments**

method The population-generation method to use. Either "virtual individuals" or "direct

resampling." Short names may be used: "d" or "dr" for "direct resampling", and

"v" or "vi" for "virtual individuals".

nsamp The desired number of individuals in the virtual population. nsamp need not be

provided if gendernum is provided.

gendernum Optional: A named list giving the numbers of male and female individuals

to include in the population, e.g. list(Male=100, Female=100). Default is NULL, meaning both males and females are included, in their proportions in the NHANES data. If both nsamp and gendernum are provided, they must agree

(i.e., nsamp must be the sum of gendernum).

agelim_years Optional: A two-element numeric vector giving the minimum and maximum

ages (in years) to include in the population. Default is c(0,79). If only a single value is provided, both minimum and maximum ages will be set to that value; e.g. agelim_years=3 is equivalent to agelim_years=c(3,3). If agelim_years is provided and agelim_months is not, agelim_years will override the default

value of agelim_months.

agelim_months Optional: A two-element numeric vector giving the minimum and maximum

ages (in months) to include in the population. Default is c(0, 959), equivalent to the default agelim_years. If only a single value is provided, both minimum and maximum ages will be set to that value; e.g. agelim_months=36 is equivalent to agelim_months=c(36,36). If agelim_months is provided and agelim_years is not, agelim_months will override the default values of agelim_years.

weight_category

Optional: The weight categories to include in the population. Default is c('Underweight',

'Normal', 'Overweight', 'Obese'). User-supplied vector must contain one

or more of these strings.

gfr_category The kidney function categories to include in the population. Default is c('Normal', 'Kidney

Disease', 'Kidney Failure') to include all kidney function levels.

reths Optional: a character vector giving the races/ethnicities to include in the popula-

tion. Default is c('Mexican American', 'Other Hispanic', 'Non-Hispanic White', 'Non-Hispanic Black', 'Other'), to include all races and ethnicities in their proportions in the NHANES data. User-supplied vector must contain

one or more of these strings.

gfr_resid_var TRUE to add residual variability to GFR predicted from serum creatinine; FALSE

to not add residual variability

ckd_epi_race_coeff

TRUE to use the CKD-EPI equation as originally published (with a coefficient changing predicted GFR for individuals identified as "Non-Hispanic Black");

FALSE to set this coefficient to 1.

# Value

A data.table where each row represents an individual, and each column represents a demographic, anthropometric, or physiological parameter.

#### Author(s)

Caroline Ring

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#### References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

#### **Examples**

```
#Simply generate a virtual population of 100 individuals,
 #using the direct-resampling method
 set.seed(42)
httkpop_generate(method='direct resampling', nsamp=100)
#Generate a population using the virtual-individuals method,
#including 80 females and 20 males,
#including only ages 20-65,
#including only Mexican American and
 #Non-Hispanic Black individuals,
 #including only non-obese individuals
httkpop_generate(method = 'virtual individuals',
gendernum=list(Female=80,
Male=20),
agelim_years=c(20,65),
reths=c('Mexican American',
'Non-Hispanic Black'),
weight_category=c('Underweight',
'Normal',
'Overweight'))
```

httkpop_mc

Converts the HTTK-Pop population data table to a table of the parameters needed by HTTK, for a specific chemical.

# Description

Takes the data table generated by httkpop_generate, and converts it to the corresponding table of HTTK model parameters for a specified chemical and HTTK model.

# Usage

```
httkpop_mc(model, samples = 1000, httkpop.dt = NULL, ...)
```

# **Arguments**

mode1	One of the HTTK models: "1compartment", "3compartmentss", "3compartment", or "pbtk".
samples	The number of Monte Carlo samples to use (can often think of these as separate individuals)
httkpop.dt	A data table generated by httkpop_generate. This defaults to NULL, in which case httkpop_generate is called to generate this table.
	Additional arugments passed on to httkpop_generate.

httkpop_virtual_indiv 123

#### Value

A data.table with a row for each individual in the sample and a column for each parater in the model.

#### Author(s)

Caroline Ring and John Wambaugh

#### References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

Rowland, Malcolm, Leslie Z. Benet, and Garry G. Graham. "Clearance concepts in pharmacokinetics." Journal of Pharmacokinetics and Biopharmaceutics 1.2 (1973): 123-136.

# **Examples**

```
set.seed(42)
indiv_examp <- httkpop_generate(method="d", nsamp=100)
httk_param <- httkpop_mc(httkpop.dt=indiv_examp,
model="1compartment")</pre>
```

httkpop_virtual_indiv Generate a virtual population by the virtual individuals method.

# Description

Generate a virtual population by the virtual individuals method.

### Usage

```
httkpop_virtual_indiv(
   nsamp = NULL,
   gendernum = NULL,
   agelim_years = c(0, 79),
   agelim_months = c(0, 959),
   weight_category = c("Underweight", "Normal", "Overweight", "Obese"),
   gfr_category = c("Normal", "Kidney Disease", "Kidney Failure"),
   reths = c("Mexican American", "Other Hispanic", "Non-Hispanic White",
        "Non-Hispanic Black", "Other"),
   gfr_resid_var = TRUE,
   ckd_epi_race_coeff = FALSE
)
```

# **Arguments**

nsamp

The desired number of individuals in the virtual population. nsamp need not be provided if gendernum is provided.

gendernum

Optional: A named list giving the numbers of male and female individuals to include in the population, e.g. list(Male=100,Female=100). Default is NULL, meaning both males and females are included, in their proportions in the NHANES data. If both nsamp and gendernum are provided, they must agree (i.e., nsamp must be the sum of gendernum).

agelim_years

Optional: A two-element numeric vector giving the minimum and maximum ages (in years) to include in the population. Default is c(0,79). If agelim_years is provided and agelim_months is not, agelim_years will override the default value of agelim_months.

agelim_months

Optional: A two-element numeric vector giving the minimum and maximum ages (in months) to include in the population. Default is c(0, 959), equivalent to the default agelim_years. If agelim_months is provided and agelim_years is not, agelim_months will override the default values of agelim_years.

weight_category

Optional: The weight categories to include in the population. Default is c('Underweight', 'Normal', 'Overweight', 'Obese'). User-supplied vector must contain one or more of these strings.

gfr_category

The kidney function categories to include in the population. Default is c('Normal', 'Kidney Disease', 'Kidney Failure') to include all kidney function levels.

reths

Optional: a character vector giving the races/ethnicities to include in the population. Default is c('Mexican American','Other Hispanic','Non-Hispanic White','Non-Hispanic Black','Other'), to include all races and ethnicities in their proportions in the NHANES data. User-supplied vector must contain one or more of these strings.

gfr_resid_var

Logical value indicating whether or not to include residual variability when generating GFR values. (Default is TRUE.)

ckd_epi_race_coeff

Logical value indicating whether or not to use the "race coefficient" from the CKD-EPI equation when estimating GFR values. (Default is FALSE.)

# Value

A data.table where each row represents an individual, and each column represents a demographic, anthropometric, or physiological parameter.

#### Author(s)

Caroline Ring

### References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

in.list

in.list	Convenience Boolean (yes/no) functions to identify chemical membership in several key lists.

# **Description**

These functions allow easy identification of whether or not a chemical CAS is included in various research projects. While it is our intent to keep these lists up-to-date, the information here is only for convenience and should not be considered to be definitive.

# Usage

```
in.list(chem.cas = NULL, which.list = "ToxCast")
```

# **Arguments**

chem. cas The Chemical Abstracts Service Resgistry Number (CAS-RN) corresponding to

the chemical of interest.

which.list A character string that can take the following values: "ToxCast", "Tox21", "Ex-

poCast", "NHANES", ""NHANES.serum.parent", "NHANES.serum.analyte", "NHANES.blood.pare

"NHANES.urine.parent","NHANES.urine.analyte"

#### **Details**

Tox21: Toxicology in the 21st Century (Tox21) is a U.S. federal High Throughput Screening (HTS) collaboration among EPA, NIH, including National Center for Advancing Translational Sciences and the National Toxicology Program at the National Institute of Environmental Health Sciences, and the Food and Drug Administration. (Bucher et al., 2008)

ToxCast: The Toxicity Forecaster (ToxCast) is a HTS screening project led by the U.S. EPA to perform additional testing of a subset of Tox21 chemicals. (Judson et al. 2010)

ExpoCast: ExpoCast (Exposure Forecaster) is an U.S. EPA research project to generate tenetative exposure estimates (e.g., mg/kg BW/day) for thousands of chemicals that have little other information using models and informatics. (Wambaugh et al. 2014)

NHANES: The U.S. Centers for Disease Control (CDC) National Health and Nutrition Examination Survery (NHANES) is an on-going survey to characterize the health and biometrics (e.g., weight, height) of the U.S. population. One set of measurments includes the quantification of xenobiotic chemicals in various samples (blood, serum, urine) of the thousands of surveyed individuals. (CDC, 2014)

### Value

logical A Boolean (1/0) value that is TRUE if the chemical is in the list.

# Author(s)

John Wambaugh

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#### References

Bucher, J. R. (2008). Guest Editorial: NTP: New Initiatives, New Alignment. Environ Health Perspect 116(1).

Judson, R. S., Houck, K. A., Kavlock, R. J., Knudsen, T. B., Martin, M. T., Mortensen, H. M., Reif, D. M., Rotroff, D. M., Shah, I., Richard, A. M. and Dix, D. J. (2010). In Vitro Screening of Environmental Chemicals for Targeted Testing Prioritization: The ToxCast Project. Environmental Health Perspectives 118(4), 485-492.

Wambaugh, J. F., Wang, A., Dionisio, K. L., Frame, A., Egeghy, P., Judson, R. and Setzer, R. W. (2014). High Throughput Heuristics for Prioritizing Human Exposure to Environmental Chemicals. Environmental Science & Technology, 10.1021/es503583j.

CDC (2014). National Health and Nutrition Examination Survey. Available at: https://www.cdc.gov/nchs/nhanes.htm.

#### See Also

is. httk for determining inclusion in httk project

#### **Examples**

```
httk.table <- get_cheminfo(info=c("CAS","Compound"))
httk.table[,"Rat"] <- ""
httk.table[,"NHANES"] <- ""
httk.table[,"Tox21"] <- ""
httk.table[,"ToxCast"] <- ""
httk.table[,"ExpoCast"] <- ""
httk.table[,"PBTK"] <- ""
# To make this example run quickly, this loop is only over the first five
# chemicals. To build a table with all available chemicals use:
# for (this.cas in httk.table$CAS)
for (this.cas in httk.table$CAS[1:5])
{
    this.index <- httk.table$CAS==this.cas
    if (is.nhanes(this.cas)) httk.table[this.index,"NHANES"] <- "Y"
    if (is.tox21(this.cas)) httk.table[this.index,"Tox21"] <- "Y"
    if (is.expocast(this.cas)) httk.table[this.index,"ToxCast"] <- "Y"
    if (is.expocast(this.cas)) httk.table[this.index,"ExpoCast"] <- "Y"
    if (is.httk(this.cas,model="PBTK")) httk.table[this.index,"Rat"] <- "Y"
    if (is.httk(this.cas,species="rat")) httk.table[this.index,"Rat"] <- "Y"</pre>
```

invitro_mc

Draw in vitro TK parameters including uncertainty and variability.

# Description

Given a CAS in the HTTK data set, a virtual population from HTTK-Pop, some user specifications on the assumed distributions of Funbound.plasma and Clint, draw "individual" values of Funbound.plasma and Clint from those distributions.

invitro_mc 127

#### Usage

```
invitro_mc(
  parameters.dt = NULL,
  samples,
  fup.meas.cv = 0.4,
  clint.meas.cv = 0.3,
  fup.pop.cv = 0.3,
  clint.pop.cv = 0.3,
  poormetab = TRUE,
  fup.lod = 0.01,
  fup.censored.dist = FALSE,
  adjusted.Funbound.plasma = TRUE,
  clint.pvalue.threshold = 0.05,
  minimum.Funbound.plasma = 1e-04
)
```

#### **Arguments**

parameters.dt A data table of physiological parameters

samples The number of samples to draw.

fup.meas.cv Coefficient of variation of distribution of measured Funbound.plasma values.

clint.meas.cv Coefficient of variation of distribution of measured Clint values.

fup.pop.cv Coefficient of variation of distribution of population Funbound.plasma values.

clint.pop.cv Coefficient of variation of distribution of population Clint values.

poormetab Logical. Whether to include poor metabolizers in the Clint distribution or not.

fup.lod The average limit of detection for Funbound.plasma, below which distribution

will be censored if fup.censored.dist is TRUE. Default 0.01.

fup.censored.dist

Logical. Whether to draw Funbound.plasma from a censored distribution or

not.

adjusted.Funbound.plasma

Uses adjusted Funbound.plasma when set to TRUE.

clint.pvalue.threshold

Hepatic clearance for chemicals where the in vitro clearance assay result has a

p-values greater than the threshold are set to zero.

minimum.Funbound.plasma

Monte Carlo draws less than this value are set equal to this value (default is

0.0001 – half the lowest measured Fup in our dataset).

parameters A list of chemical-specific model parameters containing at least Funbound.plasma,

Clint, and Fhep.assay.correction.

#### Value

A data.table with three columns: Funbound.plasma and Clint, containing the sampled values, and Fhep.assay.correction, containing the value for fraction unbound in hepatocyte assay.

#### Author(s)

Caroline Ring and John Wambaugh

128 is.httk

#### References

Wambaugh, John F., et al. "Assessing Toxicokinetic Uncertainty and Variability in Risk Prioritization." Toxicological Sciences (2019).

is.httk	Convenience Boolean (yes/no) function to identify chemical member-
	ship and treatment within the httk project.

#### **Description**

Allows easy identification of whether or not a chemical CAS is included in various aspects of the httk research project (by model type and species of interest). While it is our intent to keep these lists up-to-date, the information here is only for convenience and should not be considered definitive.

#### Usage

```
is.httk(chem.cas, species = "Human", model = "3compartmentss")
```

# **Arguments**

chem. cas The Chemical Abstracts Service Resgistry Number (CAS-RN) corresponding to

the chemical of interest.

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

model Model used in calculation, 'pbtk' for the multiple compartment model, '1com-

partment' for the one compartment model, '3compartment' for three compartment model, '3compartmentss' for the three compartment model without partition coefficients, or 'schmitt' for chemicals with logP and fraction unbound

(used in predict_partitioning_schmitt).

#### **Details**

Tox21: Toxicology in the 21st Century (Tox21) is a U.S. federal High Throughput Screening (HTS) collaboration among EPA, NIH, including National Center for Advancing Translational Sciences and the National Toxicology Program at the National Institute of Environmental Health Sciences, and the Food and Drug Administration. (Bucher et al., 2008)

ToxCast: The Toxicity Forecaster (ToxCast) is a HTS screening project led by the U.S. EPA to perform additional testing of a subset of Tox21 chemicals. (Judson et al. 2010)

ExpoCast: ExpoCast (Exposure Forecaster) is an U.S. EPA research project to generate tenetative exposure estimates (e.g., mg/kg BW/day) for thousands of chemicals that have little other information using models and informatics. (Wambaugh et al. 2014)

NHANES: The U.S. Centers for Disease Control (CDC) National Health and Nutrition Examination Survery (NHANES) is an on-going survey to characterize the health and biometrics (e.g., weight, height) of the U.S. population. One set of measurments includes the quantification of xenobiotic chemicals in various samples (blood, serum, urine) of the thousands of surveyed individuals. (CDC, 2014)

#### Value

logical A Boolean (1/0) value that is TRUE if the chemical is included in the httk project with a given modeling scheme (PBTK) and a given species

is.httk 129

#### Author(s)

John Wambaugh

#### References

Bucher, J. R. (2008). Guest Editorial: NTP: New Initiatives, New Alignment. Environ Health Perspect 116(1).

Judson, R. S., Houck, K. A., Kavlock, R. J., Knudsen, T. B., Martin, M. T., Mortensen, H. M., Reif, D. M., Rotroff, D. M., Shah, I., Richard, A. M. and Dix, D. J. (2010). In Vitro Screening of Environmental Chemicals for Targeted Testing Prioritization: The ToxCast Project. Environmental Health Perspectives 118(4), 485-492.

Wambaugh, J. F., Wang, A., Dionisio, K. L., Frame, A., Egeghy, P., Judson, R. and Setzer, R. W. (2014). High Throughput Heuristics for Prioritizing Human Exposure to Environmental Chemicals. Environmental Science & Technology, 10.1021/es503583j.

CDC (2014). National Health and Nutrition Examination Survey. Available at: https://www.cdc.gov/nchs/nhanes.htm.

#### See Also

in.list for determining chemical membership in several other key lists

# **Examples**

```
httk.table <- get_cheminfo(info=c("CAS","Compound"))</pre>
httk.table[,"Rat"] <- ""</pre>
httk.table[,"NHANES"] <- ""</pre>
httk.table[,"Tox21"] <- ""
httk.table[,"ToxCast"] <- ""</pre>
httk.table[,"ExpoCast"] <- ""</pre>
\verb|httk.table[,"PBTK"] <- ""
# To make this example run quickly, this loop is only over the first five
# chemicals. To build a table with all available chemicals use:
# for (this.cas in httk.table$CAS)
for (this.cas in httk.table$CAS[1:5])
  this.index <- httk.table$CAS==this.cas
  if (is.nhanes(this.cas)) httk.table[this.index,"NHANES"] <- "Y"</pre>
  if (is.tox21(this.cas)) httk.table[this.index,"Tox21"] <- "Y"</pre>
  if (is.toxcast(this.cas)) httk.table[this.index,"ToxCast"] <- "Y"</pre>
  if (is.expocast(this.cas)) httk.table[this.index,"ExpoCast"] <- "Y"</pre>
  if (is.httk(this.cas,model="PBTK")) httk.table[this.index,"PBTK"] <- "Y"</pre>
  if (is.httk(this.cas,species="rat")) httk.table[this.index,"Rat"] <- "Y"</pre>
```

johnson

is_in_inclusive	Checks whether a value, or all values in a vector, is within inclusive limits
-----------------	-------------------------------------------------------------------------------

# Description

Checks whether a value, or all values in a vector, is within inclusive limits

# Usage

```
is_in_inclusive(x, lims)
```

# **Arguments**

x A numeric value, or vector of values.

lims A two-element vector of (min, max) values for the inclusive limits. If x is a

vector, lims may also be a two-column matrix with nrow=length(x) where the first column is lower limits and the second column is upper limits. If x is a vector and lims is a two-element vector, then each element of x will be checked against the same limits. If x is a vector and lims is a matrix, then each element of x will be checked against the limits given by the corresponding row of lims.

#### Value

A logical vector the same length as x, indicating whether each element of x is within the inclusive limits given by lims.

# Author(s)

Caroline Ring

### References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

johnson Johnson 2006

# Description

This data set is only used in Vignette 5.

#### Usage

johnson

# Format

A data.table containing 60 rows and 11 columns.

kapraun2019 131

#### Author(s)

Caroline Ring

#### References

Johnson, Trevor N., Amin Rostami-Hodjegan, and Geoffrey T. Tucker. "Prediction of the clearance of eleven drugs and associated variability in neonates, infants and children." Clinical pharmacokinetics 45.9 (2006): 931-956.

kapraun2019

Kapraun et al. 2019 data

# **Description**

A list object containing time-varying parameters for the human maternal-fetal HTTK model. List elements contain scalar coefficients for the polynomial, logistic, Gompertz, and other functions of time describing blood flow rates, tissue volumes, hematocrits, and other anatomical/physiological quantities that change in the human mother and her fetus during pregnancy and gestation.

# Usage

kapraun2019

# **Format**

list

### Author(s)

Dustin F. Kapraun

#### Source

Kapraun et al. 2019 Fetal PBTK Model

#### References

Kapraun DF, Wambaugh JF, Setzer RW, Judson RS (2019). "Empirical models for anatomical and physiological changes in a human mother and fetus during pregnancy and gestation." *PLOS ONE*, **14**(5), 1–56. doi: 10.1371/journal.pone.0215906.

liver_mass_children

# Description

For individuals under age 18, predict kidney mass from weight, height, and gender. using equations from Ogiu et al. 1997

### Usage

```
kidney_mass_children(weight, height, gender)
```

#### **Arguments**

weight Vector of weights in kg. height Vector of heights in cm.

gender Vector of genders (either 'Male' or 'Female').

#### Value

A vector of kidney masses in kg.

### Author(s)

Caroline Ring

#### References

Ogiu, Nobuko, et al. "A statistical analysis of the internal organ weights of normal Japanese people." Health physics 72.3 (1997): 368-383.

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

# **Description**

For individuals under 18, predict the liver mass from height, weight, and gender, using equations from Ogiu et al. 1997

# Usage

```
liver_mass_children(height, weight, gender)
```

# **Arguments**

height Vector of heights in cm.
weight Vector of weights in kg.

gender Vector of genders (either 'Male' or 'Female').

load_dawson2021 133

#### Value

A vector of liver masses in kg.

#### Author(s)

Caroline Ring

#### References

Ogiu, Nobuko, et al. "A statistical analysis of the internal organ weights of normal Japanese people." Health physics 72.3 (1997): 368-383.

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

load_dawson2021

Load data from Dawson et al. 2021.

#### **Description**

This function returns an updated version of chem.physical_and_invitro.data that includes data predicted with Random Forest QSAR models developed and presented in Dawson et al. 2021, included in dawson2021.

#### Usage

load_dawson2021(overwrite = FALSE, exclude_oad = TRUE, target.env = .GlobalEnv)

#### **Arguments**

overwrite
Only matters if load.image=FALSE. If overwrite=TRUE then existing data in chem.physical_and_invitro.data will be replaced by any data/predictions in Dawson et al. (2021) that is for the same chemical and property. If overwrite=FALSE (DEFAULT) then new data for the same chemical and property are ignored. Funbound.plasma values of 0 (below limit of detection) are overwritten either way.

exclude_oad

Include the chemicals only within the applicability domain. If exlude_oad=TRUE

(DEFAULT) chemicals outside the applicability domain do not have their pre-

dicted values loaded.

target.env The environment where the new chem.physical_and_invitro.data is loaded. De-

faults to global environment.

# Value

data.frame An updated version of chem.physical_and_invitro.data.

# Author(s)

Sarah E. Davidson

load_pradeep2020

#### References

Dawson DE, Ingle BL, Phillips KA, Nichols JW, Wambaugh JF, Tornero-Velez R (2021). "Designing QSARs for Parameters of High-Throughput Toxicokinetic Models Using Open-Source Descriptors." *Environmental Science* & *Technology*, **55**(9), 6505–6517. doi: 10.1021/acs.est.0c06117, PMID: 33856768.

# **Examples**

```
## Not run:
chem.physical_and_invitro.data <- load_dawson2021()
chem.physical_and_invitro.data <- load_dawson2021(overwrite=TRUE)
## End(Not run)</pre>
```

load_pradeep2020

Load data from Pradeep et al. 2020.

# Description

This function returns an updated version of chem.physical_and_invitro.data that includes data predicted with Support Vector Machine and Random Forest models developed and presented in Pradeep et al. 2020, included in pradeep 2020.

# Usage

```
load_pradeep2020(overwrite = FALSE, target.env = .GlobalEnv)
```

# **Arguments**

overwrite

Only matters if load.image=FALSE. If overwrite=TRUE then existing data in chem.physical_and_invitro.data will be replaced by any data/predictions in Pradeep et al. (2020) that is for the same chemical and property. If overwrite=FALSE (DEFAULT) then new data for the same chemical and property are ignored. Funbound.plasma values of 0 (below limit of detection) are overwritten either way.

target.env

The environment where the new chem.physical_and_invitro.data is loaded. Defaults to global environment.

# Value

data.frame

An updated version of chem.physical_and_invitro.data.

# Author(s)

Sarah E. Davidson

load_sipes2017 135

#### References

Pradeep P, Patlewicz G, Pearce R, Wambaugh J, Wetmore B, Judson R (2020). "Using chemical structure information to develop predictive models for in vitro toxicokinetic parameters to inform high-throughput risk-assessment." Computational Toxicology, 16, 100136. ISSN 2468-1113, doi: 10.1016/j.comtox.2020.100136, https://www.sciencedirect.com/science/article/pii/ S2468111320300463.

#### **Examples**

```
## Not run:
chem.physical_and_invitro.data <- load_pradeep2020()</pre>
chem.physical_and_invitro.data <- load_pradeep2020(overwrite=TRUE)</pre>
## End(Not run)
```

load_sipes2017

Load data from Sipes et al 2017.

### **Description**

This function returns an updated version of chem.physical_and_invitro.data that includes data predicted with Simulations Plus' ADMET predictor that was used in Sipes et al. 2017, included in admet.data.

# Usage

```
load_sipes2017(overwrite = FALSE, target.env = .GlobalEnv)
```

### **Arguments**

overwrite

Only matters if load.image=FALSE. If overwrite=TRUE then existing data in chem.physical_and_invitro.data will be replaced by any data/predictions in Sipes et al. (2017) that is for the same chemical and property. If overwrite=FALSE (DEFAULT) then new data for the same chemical and property are ignored. Funbound.plasma values of 0 (below limit of detection) are overwritten either way.

The environment where the new chem.physical_and_invitro.data is loaded. Detarget.env faults to global environment.

#### Value

data.frame

An updated version of chem.physical_and_invitro.data.

### Author(s)

Robert Pearce and John Wambaugh

### References

Sipes, Nisha S., et al. "An intuitive approach for predicting potential human health risk with the Tox21 10k library." Environmental Science & Technology 51.18 (2017): 10786-10796.

lump_tissues

#### **Examples**

```
num.chems <- length(get_cheminfo())
load_sipes2017()

#We should have the ADMet Predicted chemicals from Sipes et al. (2017),
#this one is a good test since the logP is nearly 10
calc_css(chem.cas="26040-51-7")

#Let's see how many chemicals we have now with the Sipes (2017) data loaded:
length(get_cheminfo())

#Now let us reset
reset_httk()

# We should be back to our original number:
num.chems == length(get_cheminfo())</pre>
```

lump_tissues

Lump tissue parameters

# **Description**

This function takes the parameters from predict_partitioning_schmitt and lumps the partition coefficients along with the volumes and flows based on the given tissue list. It is useful in Monte Carlo simulation of individual partition coefficients when calculating the rest of body partition coefficient.

# Usage

```
lump_tissues(
  Ktissue2pu.in,
  parameters = NULL,
  tissuelist = NULL,
  species = "Human",
  tissue.vols = NULL,
  tissue.flows = NULL,
  model = "pbtk",
  suppress.messages = FALSE
)
```

# **Arguments**

Ktissue2pu.in List of partition coefficients from predict_partitioning_schmitt.

parameters A list of physiological parameters including flows and volumes for tissues in

tissuelist

tissuelist Manually specifies compartment names and tissues, which override the standard

compartment names and tissues that are usually specified in a model's associated modelinfo file. Remaining tissues in the model's associated alltissues listing

are lumped in the rest of the body.

lump_tissues 137

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

tissue.vols A list of volumes for tissues in tissuelist tissue.flows A list of flows for tissues in tissuelist

model Specify which model (and therefore which tissues) are being considered

suppress.messages

Whether or not the output message is suppressed.

#### **Details**

This function returns the flows, volumes, and partition coefficients for the lumped tissues specified in tissue list Ktissue2plasma – tissue to free plasma concentration partition coefficients for every tissue specified by Schmitt (2008) (the tissue.data table) tissuelist – a list of character vectors, the name of each entry in the list is its own compartment. The tissues in the alltissues vector are the Schmitt (2008) tissues that are to be considered in the lumping process. The tissuelist can also be manually specified for alternate lumping schemes: for example, tissuelist<-list(Rapid=c("Brain","Kidney")) specifies the flow.col and vol.col in the tissuedata.table.

#### Value

Krbc2pu Ratio of concentration of chemical in red blood cells to unbound concentration

in plasma.

Krest2pu Ratio of concentration of chemical in rest of body tissue to unbound concentra-

tion in plasma.

Vrestc Volume of the rest of the body per kg body weight, L/kg BW.

Vliverc Volume of the liver per kg body weight, L/kg BW.

Qtotal.liverf Fraction of cardiac output flowing to the gut and liver, i.e. out of the liver.

Qgutf Fraction of cardiac output flowing to the gut.

Qkidneyf Fraction of cardiac output flowing to the kidneys.

### Author(s)

John Wambaugh and Robert Pearce

#### References

Pearce, Robert G., et al. "Evaluation and calibration of high-throughput predictions of chemical distribution to tissues." Journal of pharmacokinetics and pharmacodynamics 44.6 (2017): 549-565.

#### **Examples**

```
pcs <- predict_partitioning_schmitt(chem.name='bisphenola')
tissuelist <- list(liver=c("liver"),kidney=c("kidney"),lung=c("lung"),gut=c("gut")
,muscle.bone=c('muscle','bone'))
lump_tissues(pcs,tissuelist=tissuelist)</pre>
```

mcnally_dt

lung_mass_children

Predict lung mass for children

# **Description**

For individuals under 18, predict the liver mass from height, weight, and gender, using equations from Ogiu et al. 1997

#### Usage

```
lung_mass_children(height, weight, gender)
```

#### **Arguments**

height Vector of heights in cm.
weight Vector of weights in kg.

gender Vector of genders (either 'Male' or 'Female').

#### Value

A vector of lung masses in kg.

### Author(s)

Caroline Ring

#### References

Ogiu, Nobuko, et al. "A statistical analysis of the internal organ weights of normal Japanese people." Health physics 72.3 (1997): 368-383.

Price, Paul S., et al. "Modeling interindividual variation in physiological factors used in PBPK models of humans." Critical reviews in toxicology 33.5 (2003): 469-503.

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

mcnally_dt

Reference tissue masses and flows from tables in McNally et al. 2014.

# Description

Reference tissue masses, flows, and marginal distributions from McNally et al. 2014.

# Usage

mcnally_dt

#### **Format**

```
A data.table with variables:
```

tissue Body tissue

gender Gender: Male or Female

mass_ref Reference mass in kg, from Reference Man

mass_cv Coefficient of variation for mass

mass_dist Distribution for mass: Normal or Log-normal

flow_ref Reference flow in L/h, from Reference Man

flow_cv Coefficient of variation for flow (all normally distributed)

height_ref Reference heights (by gender)

CO_ref Reference cardiac output by gender

flow_frac Fraction of CO flowing to each tissue: flow_ref/CO_ref

# Author(s)

Caroline Ring

#### Source

McNally K, Cotton R, Hogg A, Loizou G. "PopGen: A virtual human population generator." Toxicology 315, 70-85, 2004.

#### References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

metabolism_data_Linakis2020

Metabolism data involved in Linakis 2020 vignette analysis.

# **Description**

Metabolism data involved in Linakis 2020 vignette analysis.

# Usage

metabolism_data_Linakis2020

### **Format**

A data frame containing x rows and y columns.

# Author(s)

Matt Linakis

140 monte_carlo

#### Source

Matt Linakis

#### References

DSStox database (https://www.epa.gov/ncct/dsstox

monte_carlo

Monte Carlo for pharmacokinetic models

#### **Description**

This function performs Monte Carlo to assess uncertainty and variability for toxicokinetic models.

# Usage

```
monte_carlo(
  parameters,
  cv.params = NULL,
  censored.params = NULL,
  samples = 1000
)
```

#### **Arguments**

parameters

These parameters that are also listed in either cv.params or censored.params are sampled using Monte Carlo.

cv.params

The parameters listed in cv.params are sampled from a normal distribution that is truncated at zero. This argument should be a list of coefficients of variation (cv) for the normal distribution. Each entry in the list is named for a parameter in "parameters". New values are sampled with mean equal to the value in "parameters" and standard deviation equal to the mean times the cv.

censored.params

The parameters listed in censored.params are sampled from a normal distribution that is censored for values less than the limit of detection (specified separately for each parameter). This argument should be a list of sub-lists. Each sublist is named for a parameter in "params" and contains two elements: "cv" (coefficient of variation) and "LOD" (limit of detection), below which parameter values are censored. New values are sampled with mean equal to the value in "params" and standard deviation equal to the mean times the cv. Censored values are sampled on a uniform distribution between 0 and the limit of detection.

samples

This argument is the number of samples to be generated for calculating quantiles.

#### Value

A data.table with a row for each individual in the sample and a column for each parater in the model.

#### Author(s)

John Wambaugh

nhanes_mec_svy 141

#### References

Pearce, Robert G., et al. "Httk: R package for high-throughput toxicokinetics." Journal of statistical software 79.4 (2017): 1.

# **Examples**

```
#Example based on Pearce et al. (2017):
# Set up means:
params <- parameterize_pbtk(chem.name="zoxamide")</pre>
# Nothing changes:
monte_carlo(params)
vary.params <- NULL
for (this.param in names(params)[!(names(params) %in%
  c("Funbound.plasma", "pKa_Donor", "pKa_Accept" )) &
  !is.na(as.numeric(params))]) vary.params[this.param] <- 0.2</pre>
# Most everything varies with CV of 0.2:
monte_carlo(
  parameters=params,
  cv.params = vary.params)
censored.params <- list(Funbound.plasma = list(cv = 0.2, lod = 0.01))</pre>
# Fup is censored below 0.01:
monte_carlo(
  parameters=params,
  cv.params = vary.params,
  censored.params = censored.params)
```

nhanes_mec_svy

Pre-processed NHANES data.

# **Description**

NHANES data on demographics, anthropometrics, and some laboratory measures, cleaned and combined into a single data set.

#### Usage

```
nhanes_mec_svy
```

#### **Format**

A survey.design2 object, including masked cluster and strata. Variables are available as a data.table by nhanes_mec_svy\$variables. Variables are as described in NHANES Demographics and Examination documentation, with the exception of:

wtmec6yr 6-year sample weights for combining 3 cycles, computed by dividing 2-year sample weights by 3.

bmxhtlenavg Average of height and recumbent length if both were measured; if only one was measured, takes value of the one that was measured.

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logbmxwt Natural log of measured body weight.

logbmxhtlenavg Natural log of bmxhtlenavg.

weight_class One of Underweight, Normal, Overweight, or Obese. Assigned using methods in get_weight_class.

# Author(s)

Caroline Ring

#### **Source**

https://wwwn.cdc.gov/nchs/nhanes/Default.aspx

#### References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

0bach2008

Published Pharmacokinetic Parameters from Obach et al. 2008

#### **Description**

This data set is used in Vignette 4 for steady state concentration.

# Usage

0bach2008

# **Format**

A data.frame containing 670 rows and 8 columns.

#### References

Obach, R. Scott, Franco Lombardo, and Nigel J. Waters. "Trend analysis of a database of intravenous pharmacokinetic parameters in humans for 670 drug compounds." Drug Metabolism and Disposition 36.7 (2008): 1385-1405.

onlyp 143

onlyp

NHANES Exposure Data

# **Description**

This data set is only used in Vignette 6.

# Usage

onlyp

#### **Format**

A data.table containing 1060 rows and 5 columns.

# Author(s)

Caroline Ring

#### References

Wambaugh, John F., et al. "High throughput heuristics for prioritizing human exposure to environmental chemicals." Environmental science & technology 48.21 (2014): 12760-12767.

```
pancreas_mass_children
```

Predict pancreas mass for children

# Description

For individuals under 18, predict the pancreas mass from height, weight, and gender, using equations from Ogiu et al.

# Usage

```
pancreas_mass_children(height, weight, gender)
```

# **Arguments**

height Vector of heights in cm.
weight Vector of weights in kg.

gender Vector of genders (either 'Male' or 'Female').

# Value

A vector of pancreas masses in kg.

#### Author(s)

Caroline Ring

144 parameterize_1comp

#### References

Ogiu, Nobuko, et al. "A statistical analysis of the internal organ weights of normal Japanese people." Health physics 72.3 (1997): 368-383.

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

parameterize_1comp

Parameterize_1comp

#### **Description**

This function initializes the parameters needed in the function solve 1comp.

# Usage

```
parameterize_1comp(
  chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
  species = "Human",
  default.to.human = FALSE,
  adjusted.Funbound.plasma = TRUE,
  regression = TRUE,
  restrictive.clearance = TRUE,
  well.stirred.correction = TRUE,
  suppress.messages = FALSE,
  clint.pvalue.threshold = 0.05,
  minimum.Funbound.plasma = 1e-04
)
```

### Arguments

chem. cas

Chemical Abstract Services Registry Number (CAS-RN) – the chemical must be identified by either CAS, name, or DTXISD

chem. name

Chemical name (spaces and capitalization ignored) – the chemical must be iden-

tified by either CAS, name, or DTXISD

dtxsid EPA's DSSTox Structure ID (https://comptox.epa.gov/dashboard) - the

chemical must be identified by either CAS, name, or DTXSIDs

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

default.to.human

Substitutes missing rat values with human values if true.

adjusted.Funbound.plasma

Uses adjusted Funbound.plasma when set to TRUE along with volume of distri-

bution calculated with this value.

regression Whether or not to use the regressions in calculating partition coefficients in vol-

ume of distribution calculation.

restrictive.clearance

In calculating elimination rate and hepatic bioavailability, protein binding is not taken into account (set to 1) in liver clearance if FALSE.

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well.stirred.correction

Uses correction in calculation of hepatic clearance for well-stirred model if TRUE. This assumes clearance relative to amount unbound in whole blood instead of plasma, but converted to use with plasma concentration.

suppress.messages

Whether or not to suppress messages.

clint.pvalue.threshold

Hepatic clearance for chemicals where the in vitro clearance assay result has a p-value greater than the threshold are set to zero.

minimum.Funbound.plasma

Monte Carlo draws less than this value are set equal to this value (default is 0.0001 – half the lowest measured Fup in our dataset).

### Value

Vdist Volume of distribution, units of L/kg BW.

Fgutabs Fraction of the oral dose absorbed, i.e. the fraction of the dose that enters the

gutlumen.

Fhep.assay.correction

The fraction of chemical unbound in hepatocyte assay using the method of Kil-

ford et al. (2008)

kelim Elimination rate, units of 1/h.

hematocrit Percent volume of red blood cells in the blood.

kgutabs Rate chemical is absorbed, 1/h.

million.cells.per.gliver

Millions cells per gram of liver tissue.

MW Molecular Weight, g/mol.

Rblood2plasma The ratio of the concentration of the chemical in the blood to the concentration

in the plasma. Not used in calculations but included for the conversion of plasma

outputs.

hepatic.bioavailability

Fraction of dose remaining after first pass clearance, calculated from the cor-

rected well-stirred model.

BW Body Weight, kg.

### Author(s)

John Wambaugh and Robert Pearce

### References

Pearce, Robert G., et al. "Httk: R package for high-throughput toxicokinetics." Journal of statistical software 79.4 (2017): 1.

Kilford, P. J., Gertz, M., Houston, J. B. and Galetin, A. (2008). Hepatocellular binding of drugs: correction for unbound fraction in hepatocyte incubations using microsomal binding or drug lipophilicity data. Drug Metabolism and Disposition 36(7), 1194-7, 10.1124/dmd.108.020834.

parameterize_3comp

## **Examples**

parameterize_3comp

Parameterize_3comp

# Description

This function initializes the parameters needed in the function solve_3comp.

## Usage

```
parameterize_3comp(
  chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
  species = "Human",
  default.to.human = FALSE,
  force.human.clint.fup = FALSE,
  clint.pvalue.threshold = 0.05,
  adjusted.Funbound.plasma = TRUE,
  regression = TRUE,
  suppress.messages = FALSE,
  restrictive.clearance = TRUE,
  minimum.Funbound.plasma = 1e-04
)
```

### **Arguments**

chem.cas	Chemical Abstract Services Registry Number (CAS-RN) – the chemical must be identified by either CAS, name, or DTXISD
chem.name	Chemical name (spaces and capitalization ignored) – the chemical must be identified by either CAS, name, or DTXISD
dtxsid	EPA's 'DSSTox Structure ID (https://comptox.epa.gov/dashboard) – the chemical must be identified by either CAS, name, or DTXSIDs
species	Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").
default.to.human	
	Substitutes missing animal values with human values if true.

force.human.clint.fup

Forces use of human values for hepatic intrinsic clearance and fraction of unbound plasma if true.

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clint.pvalue.threshold

Hepatic clearances with clearance assays having p-values greater than the threshold are set to zero.

adjusted.Funbound.plasma

Returns adjusted Funbound.plasma when set to TRUE along with parition coefficients calculated with this value.

regression Whether or not to use the regressions in calculating partition coefficients. suppress.messages

Whether or not the output message is suppressed.

restrictive.clearance

In calculating hepatic.bioavailability, protein binding is not taken into account (set to 1) in liver clearance if FALSE.

minimum.Funbound.plasma

Monte Carlo draws less than this value are set equal to this value (default is 0.0001 – half the lowest measured Fup in our dataset).

#### Value

BW Body Weight, kg.

Clmetabolismc Hepatic Clearance, L/h/kg BW.

Fgutabs Fraction of the oral dose absorbed, i.e. the fraction of the dose that enters the

gutlumen.

Funbound.plasma

Fraction of plasma that is not bound.

Fhep.assay.correction

The fraction of chemical unbound in hepatocyte assay using the method of Kil-

ford et al. (2008)

hematocrit Percent volume of red blood cells in the blood.

Kgut2pu Ratio of concentration of chemical in gut tissue to unbound concentration in

plasma.

Kliver2pu Ratio of concentration of chemical in liver tissue to unbound concentration in

plasma.

Krbc2pu Ratio of concentration of chemical in red blood cells to unbound concentration

in plasma.

Krest2pu Ratio of concentration of chemical in rest of body tissue to unbound concentra-

tion in plasma.

million.cells.per.gliver

Millions cells per gram of liver tissue.

MW Molecular Weight, g/mol.

Ocardiace Cardiac Output, L/h/kg BW^3/4.

Qgfrc Glomerular Filtration Rate, L/h/kg BW^3/4, volume of fluid filtered from kidney

and excreted.

Qgutf Fraction of cardiac output flowing to the gut.
Qliverf Fraction of cardiac output flowing to the liver.

Rblood2plasma The ratio of the concentration of the chemical in the blood to the concentration

in the plasma.

Vgutc Volume of the gut per kg body weight, L/kg BW.
Vliverc Volume of the liver per kg body weight, L/kg BW.

Vrestc Volume of the rest of the body per kg body weight, L/kg BW.

### Author(s)

Robert Pearce and John Wambaugh

#### References

Pearce, Robert G., et al. "Httk: R package for high-throughput toxicokinetics." Journal of statistical software 79.4 (2017): 1.

Kilford, P. J., Gertz, M., Houston, J. B. and Galetin, A. (2008). Hepatocellular binding of drugs: correction for unbound fraction in hepatocyte incubations using microsomal binding or drug lipophilicity data. Drug Metabolism and Disposition 36(7), 1194-7, 10.1124/dmd.108.020834.

### **Examples**

```
parameterize_fetal_pbtk
```

Parameterize_fetal_PBTK

## **Description**

This function initializes the parameters needed in the functions solve_fetal_pbtk by calling solve_pbtk and adding additional parameters.

## Usage

```
parameterize_fetal_pbtk(
  chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
  species = "Human",
  fetal_fup_adjustment = TRUE,
  return.kapraun2019 = TRUE,
  suppress.messages = FALSE,
  ...
)
```

## **Arguments**

chem.cas	Either the chemical name or the CAS number must be specified.
chem.name	Either the chemical name or the CAS number must be specified.
dtxsid	$EPA's \ DSSTox \ Structure \ ID \ (http://comptox.epa.gov/dashboard) \ the \ chemical \ must be \ identified \ by \ either \ CAS, \ name, \ or \ DTXSIDs$
species	Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human"). Currently only a narrow human model is supported.

fetal_fup_adjustment

Logical indicator of whether to use an adjusted estimate for fetal fup based on the fetal:maternal plasma protein binding ratios presented in McNamara and Alcorn's 2002 study "Protein Binding Predictions in Infants." Defaults to TRUE.

return.kapraun2019

If TRUE (default) the empirical parameters for the Kapraun et al. (2019) maternal-fetal growth parameters are provided.

suppress.messages

Whether or not the output message is suppressed.

... Arguments passed to parameterize_pbtk.

### Value

pre_pregnant_BW

Body Weight before pregnancy, kg.

Clmetabolismc Hepatic Clearance, L/h/kg BW.

Fgutabs Fraction of the oral dose absorbed, i.e. the fraction of the dose that enters the

gutlumen.

Funbound.plasma

Fraction of plasma that is not bound.

Fhep.assay.correction

The fraction of chemical unbound in hepatocyte assay using the method of Kil-

ford et al. (2008)

hematocrit Percent volume of red blood cells in the blood.

Kgut2pu Ratio of concentration of chemical in gut tissue to unbound concentration in

plasma.

kgutabs Rate that chemical enters the gut from gutlumen, 1/h.

Kkidney2pu Ratio of concentration of chemical in kidney tissue to unbound concentration in

plasma.

Kliver2pu Ratio of concentration of chemical in liver tissue to unbound concentration in

plasma.

Klung2pu Ratio of concentration of chemical in lung tissue to unbound concentration in

plasma.

Krbc2pu Ratio of concentration of chemical in red blood cells to unbound concentration

in plasma.

Krest2pu Ratio of concentration of chemical in rest of body tissue to unbound concentra-

tion in plasma.

million.cells.per.gliver

Millions cells per gram of liver tissue.

MW Molecular Weight, g/mol.

Qgfrc Glomerular Filtration Rate, L/h/kg BW^3/4, volume of fluid filtered from kidney

and excreted.

in the plasma from available_rblood2plasma.

Vgutc Volume of the gut per kg body weight, L/kg BW.

Vkidneyc Volume of the kidneys per kg body weight, L/kg BW.
Vliverc Volume of the liver per kg body weight, L/kg BW.

Vlungc	Volume of the lungs per kg body weight, L/kg BW.
Vthyroidc	Volume of the thyroid per kg body weight, L/kg BW.
Kfgut2pu	Ratio of concentration of chemical in fetal gut tissue to unbound concentration in plasma.
Kfkidney2pu	Ratio of concentration of chemical in fetal kidney tissue to unbound concentration in plasma.
Kfliver2pu	Ratio of concentration of chemical in fetal liver tissue to unbound concentration in plasma.
Kflung2pu	Ratio of concentration of chemical in fetal lung tissue to unbound concentration in plasma.
Kfrest2pu	Ratio of concentration of chemical in fetal rest of body tissue to unbound concentration in plasma.
Kfbrain2pu	Ratio of concentration of chemical in fetal brain tissue to unbound concentration in plasma.
Kthyroid2pu	Ratio of concentration of chemical in fetal thyroid tissue to unbound concentration in plasma.
Kfthyroid2pu	Ratio of concentration of chemical in fetal thyroid tissue to unbound concentration in plasma.
Kplacenta2pu	Ratio of concentration of chemical in placental tissue to unbound concentration in maternal plasma.
Kfplacenta2pu	Ratio of concentration of chemical in placental tissue to unbound concentration in fetal plasma.

## Author(s)

Robert Pearce, Mark Sfeir, John Wambaugh, and Dustin Kapraun Mark Sfeir, Dustin Kapraun, John Wambaugh

### References

Kilford, P. J., Gertz, M., Houston, J. B. and Galetin, A. (2008). Hepatocellular binding of drugs: correction for unbound fraction in hepatocyte incubations using microsomal binding or drug lipophilicity data. Drug Metabolism and Disposition 36(7), 1194-7, 10.1124/dmd.108.020834.

McNamara PJ, Alcorn J. Protein binding predictions in infants. AAPS PharmSci. 2002;4(1):E4. doi: 10.1208/ps040104. PMID: 12049488.

# **Examples**

```
parameters <- parameterize_fetal_pbtk(chem.cas='80-05-7')
parameters <- parameterize_fetal_pbtk(chem.name='Bisphenol-A',species='Rat')</pre>
```

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```
parameterize_gas_pbtk
```

### **Description**

This function initializes the parameters needed in the function solve_gas_pbtk

### Usage

```
parameterize_gas_pbtk(
  chem.cas = NULL,
  chem.name = NULL
  dtxsid = NULL,
  species = "Human",
  default.to.human = FALSE,
 tissuelist = list(liver = c("liver"), kidney = c("kidney"), lung = c("lung"), gut =
    c("gut")),
  force.human.clint.fup = FALSE,
  clint.pvalue.threshold = 0.05,
  adjusted.Funbound.plasma = TRUE,
  regression = TRUE,
  vmax = 0,
  km = 1,
  exercise = FALSE,
  fR = 12.
  VT = 0.75,
  VD = 0.15,
  suppress.messages = FALSE,
  minimum.Funbound.plasma = 1e-04,
)
```

# **Arguments**

chem.cas Either the chemical name or the CAS number must be specified. chem.name Either the chemical name or the CAS number must be specified. EPA's DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chemdtxsid ical must be identified by either CAS, name, or DTXSIDs species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human"). default.to.human Substitutes missing animal values with human values if true (hepatic intrinsic clearance or fraction of unbound plasma). tissuelist Specifies compartment names and tissues groupings. Remaining tissues in tissue.data are lumped in the rest of the body. However, solve_pbtk only works with the default parameters.

force.human.clint.fup

Forces use of human values for hepatic intrinsic clearance and fraction of unbound plasma if true.

clint.pvalue.threshold

Hepatic clearance for chemicals where the in vitro clearance assay result has a p-values greater than the threshold are set to zero.

adjusted.Funbound.plasma

Returns adjusted Funbound.plasma when set to TRUE along with parition coef-

ficients calculated with this value.

regression Whether or not to use the regressions in calculating partition coefficients.

vmax Michaelis-Menten vmax value in reactions/min

km Michaelis-Menten concentration of half-maximal reaction velocity in desired

output concentration units.

exercise Logical indicator of whether to simulate an exercise-induced heightened respi-

ration rate

fR Respiratory frequency (breaths/minute), used especially to adjust breathing rate

in the case of exercise. This parameter, along with VT and VD (below) gives another option for calculating Qalv (Alveolar ventilation) in case pulmonary

ventilation rate is not known

VT Tidal volume (L), to be modulated especially as part of simulating the state of

exercise

VD Anatomical dead space (L), to be modulated especially as part of simulating the

state of exercise

suppress.messages

Whether or not the output message is suppressed.

minimum.Funbound.plasma

Monte Carlo draws less than this value are set equal to this value (default is

0.0001 – half the lowest measured Fup in our dataset).

... Other parameters

### Value

BW Body Weight, kg.

Clint Hepatic intrinsic clearance, uL/min/10^6 cells

Clint.dist Distribution of hepatic intrinsic clearance values (median, lower 95th, upper

95th, p value)

Clmetabolismc Hepatic Clearance, L/h/kg BW.

Fgutabs Fraction of the oral dose absorbed, i.e. the fraction of the dose that enters the

gut lumen.

Fhep.assay.correction

The fraction of chemical unbound in hepatocyte assay using the method of Kil-

ford et al. (2008)

Funbound.plasma

Fraction of chemical unbound to plasma.

Funbound.plasma.adjustment

Fraction unbound to plasma adjusted as described in Pearce et al. 2017

Funbound.plasma.dist

Distribution of fraction unbound to plasma (median, lower 95th, upper 95th)

hematocrit Percent volume of red blood cells in the blood.

Kblood2air Ratio of concentration of chemical in blood to air

Kgut2pu Ratio of concentration of chemical in gut tissue to unbound concentration in

plasma.

kgutabs Rate that chemical enters the gut from gutlumen, 1/h.

Kkidney2pu Ratio of concentration of chemical in kidney tissue to unbound concentration in

plasma.

Kliver2pu Ratio of concentration of chemical in liver tissue to unbound concentration in

plasma.

Klung2pu Ratio of concentration of chemical in lung tissue to unbound concentration in

plasma.

km Michaelis-Menten concentration of half-maximal activity

Kmuc2air Mucus to air partition coefficient

Krbc2pu Ratio of concentration of chemical in red blood cells to unbound concentration

in plasma.

Krest2pu Ratio of concentration of chemical in rest of body tissue to unbound concentra-

tion in plasma.

kUrtc Unscaled upper respiratory tract uptake parameter (L/h/kg^0.75)

liver.density Density of liver in g/mL

MA phospholipid:water distribution coefficient, membrane affinity

million.cells.per.gliver

Millions cells per gram of liver tissue.

MW Molecular Weight, g/mol.

pKa_Accept compound H association equilibrium constant(s)
pKa_Donor compound H dissociation equilibrium constant(s)

Pow octanol:water partition coefficient (not log transformed)

Qalvc Unscaled alveolar ventilation rate ( $L/h/kg^0.75$ )

Qcardiacc Cardiac Output, L/h/kg BW^3/4.

Qgfrc Glomerular Filtration Rate, L/h/kg BW^0.75, volume of fluid filtered from kid-

ney and excreted.

Qgutf Fraction of cardiac output flowing to the gut.

Qkidneyf Fraction of cardiac output flowing to the kidneys.

Qliverf Fraction of cardiac output flowing to the liver.

Qlungf Fraction of cardiac output flowing to lung tissue.

Qrestf Fraction of blood flow to rest of body

Rblood2plasma The ratio of the concentration of the chemical in the blood to the concentration

in the plasma from available_rblood2plasma.

Vartc Volume of the arteries per kg body weight, L/kg BW.

Vgutc Volume of the gut per kg body weight, L/kg BW.

Vkidneyc Volume of the kidneys per kg body weight, L/kg BW.

Vliverc Volume of the liver per kg body weight, L/kg BW.

Vlungc Volume of the lungs per kg body weight, L/kg BW.

vmax Michaelis-Menten maximum reaction velocity (1/min)

Vmucc Unscaled mucosal volume (L/kg BW^0.75

Vrestc Volume of the rest of the body per kg body weight, L/kg BW.

Vvenc Volume of the veins per kg body weight, L/kg BW.

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### Author(s)

Matt Linakis, Robert Pearce, John Wambaugh

#### References

Linakis, Matthew W., et al. "Development and Evaluation of a High Throughput Inhalation Model for Organic Chemicals", submitted

Kilford, P. J., Gertz, M., Houston, J. B. and Galetin, A. (2008). Hepatocellular binding of drugs: correction for unbound fraction in hepatocyte incubations using microsomal binding or drug lipophilicity data. Drug Metabolism and Disposition 36(7), 1194-7, 10.1124/dmd.108.020834.

### **Examples**

parameterize_pbtk

Parameterize_PBTK

## **Description**

This function initializes the parameters needed in the functions solve_pbtk, calc_css, and others using the multiple compartment model.

## Usage

```
parameterize_pbtk(
  chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
  species = "Human",
  default.to.human = FALSE,
  tissuelist = list(liver = c("liver"), kidney = c("kidney"), lung = c("lung"), gut =
       c("gut")),
  force.human.clint.fup = FALSE,
  clint.pvalue.threshold = 0.05,
  adjusted.Funbound.plasma = TRUE,
  regression = TRUE,
  suppress.messages = FALSE,
  restrictive.clearance = TRUE,
```

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```
minimum.Funbound.plasma = 1e-04
)
```

### **Arguments**

chem.cas Chemical Abstract Services Registry Number (CAS-RN) – the chemical must

be identified by either CAS, name, or DTXISD

chem. name Chemical name (spaces and capitalization ignored) – the chemical must be iden-

tified by either CAS, name, or DTXISD

dtxsid EPA's 'DSSTox Structure ID (https://comptox.epa.gov/dashboard) - the

chemical must be identified by either CAS, name, or DTXSIDs

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

default.to.human

Substitutes missing animal values with human values if true (hepatic intrinsic

clearance or fraction of unbound plasma).

tissuelist Specifies compartment names and tissues groupings. Remaining tissues in tis-

sue.data are lumped in the rest of the body. However, solve_pbtk only works

with the default parameters.

force.human.clint.fup

Forces use of human values for hepatic intrinsic clearance and fraction of un-

bound plasma if true.

clint.pvalue.threshold

Hepatic clearance for chemicals where the in vitro clearance assay result has a

p-values greater than the threshold are set to zero.

adjusted.Funbound.plasma

Returns adjusted Funbound.plasma when set to TRUE along with parition coef-

ficients calculated with this value.

regression Whether or not to use the regressions in calculating partition coefficients.

suppress.messages

Whether or not the output message is suppressed.

restrictive.clearance

In calculating hepatic.bioavailability, protein binding is not taken into account

(set to 1) in liver clearance if FALSE.

minimum.Funbound.plasma

Monte Carlo draws less than this value are set equal to this value (default is

0.0001 – half the lowest measured Fup in our dataset).

### Value

BW Body Weight, kg.

Clmetabolismc Hepatic Clearance, L/h/kg BW.

Fgutabs Fraction of the oral dose absorbed, i.e. the fraction of the dose that enters the

gutlumen.

Funbound.plasma

Fraction of plasma that is not bound.

Fhep.assay.correction

The fraction of chemical unbound in hepatocyte assay using the method of Kil-

ford et al. (2008)

hematocrit Percent volume of red blood cells in the blood.

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Kgut2pu Ratio of concentration of chemical in gut tissue to unbound concentration in

plasma.

kgutabs Rate that chemical enters the gut from gutlumen, 1/h.

Kkidney2pu Ratio of concentration of chemical in kidney tissue to unbound concentration in

plasma.

Kliver2pu Ratio of concentration of chemical in liver tissue to unbound concentration in

plasma.

Klung2pu Ratio of concentration of chemical in lung tissue to unbound concentration in

plasma.

Krbc2pu Ratio of concentration of chemical in red blood cells to unbound concentration

in plasma.

Krest2pu Ratio of concentration of chemical in rest of body tissue to unbound concentra-

tion in plasma.

million.cells.per.gliver

Millions cells per gram of liver tissue.

MW Molecular Weight, g/mol.

Qcardiacc Cardiac Output, L/h/kg BW^3/4.

Qgfrc Glomerular Filtration Rate, L/h/kg BW^3/4, volume of fluid filtered from kidney

and excreted.

Qgutf Fraction of cardiac output flowing to the gut.

Qkidneyf Fraction of cardiac output flowing to the kidneys.

Oliverf Fraction of cardiac output flowing to the liver.

Rblood2plasma The ratio of the concentration of the chemical in the blood to the concentration

in the plasma from available_rblood2plasma.

Volume of the arteries per kg body weight, L/kg BW.

Vgutc

Volume of the gut per kg body weight, L/kg BW.

Vkidneyc

Volume of the kidneys per kg body weight, L/kg BW.

Vliverc

Volume of the liver per kg body weight, L/kg BW.

Vlungc

Volume of the lungs per kg body weight, L/kg BW.

Vrestc Volume of the rest of the body per kg body weight, L/kg BW.

Vvenc Volume of the veins per kg body weight, L/kg BW.

### Author(s)

John Wambaugh and Robert Pearce

### References

Pearce, Robert G., et al. "Httk: R package for high-throughput toxicokinetics." Journal of statistical software 79.4 (2017): 1.

Kilford, P. J., Gertz, M., Houston, J. B. and Galetin, A. (2008). Hepatocellular binding of drugs: correction for unbound fraction in hepatocyte incubations using microsomal binding or drug lipophilicity data. Drug Metabolism and Disposition 36(7), 1194-7, 10.1124/dmd.108.020834.

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### **Examples**

parameterize_schmitt Get the Parameters for Schmitt's Tissue Partition Coefficient Method

### **Description**

This function provides the necessary parameters to run predict_partitioning_schmitt, excluding the data in tissue.data.

## Usage

```
parameterize_schmitt(
  chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
  parameters = NULL,
  species = "Human",
  default.to.human = FALSE,
  force.human.fup = FALSE,
  suppress.messages = FALSE,
  minimum.Funbound.plasma = 1e-04
)
```

### **Arguments**

chem.cas	Chemical Abstract Services Registry Number (CAS-RN) – if parameters is not specified then the chemical must be identified by either CAS, name, or DTXISD
chem.name	Chemical name (spaces and capitalization ignored) – if parameters is not specified then the chemical must be identified by either CAS, name, or DTXISD
dtxsid	EPA's DSSTox Structure ID (https://comptox.epa.gov/dashboard) – if parameters is not specified then the chemical must be identified by either CAS, name, or DTXSIDs
parameters	Chemcial and physiological description parameters needed to run the Schmitt et al. (2008) model
species	Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

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default.to.human

Substitutes missing fraction of unbound plasma with human values if true.

force.human.fup

Returns human fraction of unbound plasma in calculation for rats if true. When species is specified as rabbit, dog, or mouse, the human unbound fraction is substituted.

suppress.messages

Whether or not the output message is suppressed.

minimum.Funbound.plasma

Monte Carlo draws less than this value are set equal to this value (default is 0.0001 – half the lowest measured Fup in our dataset).

#### Value

Funbound.plasma

corrected unbound fraction in plasma

unadjusted.Funbound.plasma

measured unbound fraction in plasma (0.005 if below limit of detection)

Pow octanol:water partition coefficient (not log transformed)

pKa_Donor compound H dissociation equilibrium constant(s)
pKa_Accept compound H association equilibrium constant(s)

MA phospholipid:water distribution coefficient, membrane affinity

Fprotein.plasma

protein fraction in plasma

plasma.pH pH of the plasma

### Author(s)

Robert Pearce and John Wambaugh

## References

Schmitt, Walter. "General approach for the calculation of tissue to plasma partition coefficients." Toxicology in Vitro 22.2 (2008): 457-467.

Schmitt, Walter. "Corrigendum to: General approach for the calculation of tissue to plasma partition coefficients" Toxicology in Vitro 22.6 (2008): 1666.

Peyret, Thomas, Patrick Poulin, and Kannan Krishnan. "A unified algorithm for predicting partition coefficients for PBPK modeling of drugs and environmental chemicals." Toxicology and applied pharmacology 249.3 (2010): 197-207.

Pearce, Robert G., et al. "Evaluation and calibration of high-throughput predictions of chemical distribution to tissues." Journal of pharmacokinetics and pharmacodynamics 44.6 (2017): 549-565.

#### **Examples**

```
parameterize_schmitt(chem.name='bisphenola')
```

```
parameterize_steadystate
```

Parameterize_SteadyState

## **Description**

This function initializes the parameters needed in the functions calc_mc_css, calc_mc_oral_equiv, and calc_analytic_css for the three compartment steady state model ('3compartmentss').

### Usage

```
parameterize_steadystate(
  chem.cas = NULL,
  chem.name = NULL,
  dtxsid = NULL,
  species = "Human",
  clint.pvalue.threshold = 0.05,
  default.to.human = FALSE,
  human.clint.fup = FALSE,
  adjusted.Funbound.plasma = TRUE,
  restrictive.clearance = TRUE,
  fup.lod.default = 0.005,
  suppress.messages = FALSE,
  minimum.Funbound.plasma = 1e-04
)
```

### **Arguments**

chem.cas	Chemical Abstract Services Registry Number (CAS-RN) – the chemical must be identified by either CAS, name, or DTXISD
chem.name	Chemical name (spaces and capitalization ignored) – the chemical must be identified by either CAS, name, or DTXISD
dtxsid	EPA's DSSTox Structure ID (https://comptox.epa.gov/dashboard) – the chemical must be identified by either CAS, name, or DTXSIDs
species	Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").
clint.pvalue.threshold	
	Hepatic clearances with clearance assays having p-values greater than the thresh-
	old are set to zero.
default.to.human	

Substitutes missing rat values with human values if true.

human.clint.fup

Uses human hepatic intrinsic clearance and fraction of unbound plasma in calculation of partition coefficients for rats if true.

adjusted.Funbound.plasma

Returns adjusted Funbound.plasma when set to TRUE.

restrictive.clearance

In calculating hepatic.bioavailability, protein binding is not taken into account (set to 1) in liver clearance if FALSE.

fup.lod.default

Default value used for fraction of unbound plasma for chemicals where measured value was below the limit of detection. Default value is 0.0005.

suppress.messages

Whether or not the output message is suppressed.

minimum.Funbound.plasma

Monte Carlo draws less than this value are set equal to this value (default is 0.0001 – half the lowest measured Fup in our dataset).

### Value

Clint Hepatic Intrinsic Clearance, uL/min/10⁶ cells.

Fgutabs Fraction of the oral dose absorbed, i.e. the fraction of the dose that enters the

gutlumen.

Funbound.plasma

Fraction of plasma that is not bound.

Qtotal.liverc Flow rate of blood exiting the liver, L/h/kg BW^3/4.

Qgfrc Glomerular Filtration Rate, L/h/kg BW^3/4, volume of fluid filtered from kidney

and excreted.

BW Body Weight, kg

MW Molecular Weight, g/mol

million.cells.per.gliver

Millions cells per gram of liver tissue.

Vliverc Volume of the liver per kg body weight, L/kg BW.

liver.density Liver tissue density, kg/L.

Fhep.assay.correction

The fraction of chemical unbound in hepatocyte assay using the method of Kil-

ford et al. (2008)

hepatic.bioavailability

Fraction of dose remaining after first pass clearance, calculated from the cor-

rected well-stirred model.

### Author(s)

John Wambaugh

### References

Pearce, Robert G., et al. "Httk: R package for high-throughput toxicokinetics." Journal of statistical software 79.4 (2017): 1.

Kilford, P. J., Gertz, M., Houston, J. B. and Galetin, A. (2008). Hepatocellular binding of drugs: correction for unbound fraction in hepatocyte incubations using microsomal binding or drug lipophilicity data. Drug Metabolism and Disposition 36(7), 1194-7, 10.1124/dmd.108.020834.

## **Examples**

```
parameters <- parameterize_steadystate(chem.name='Bisphenol-A',species='Rat')
parameters <- parameterize_steadystate(chem.cas='80-05-7')</pre>
```

pc.data 161

pc.data

Partition Coefficient Data

### **Description**

Measured rat in vivo partition coefficients and data for predicting them.

### Usage

pc.data

#### **Format**

A data.frame.

#### Author(s)

Jimena Davis and Robert Pearce

#### References

Schmitt, W., General approach for the calculation of tissue to plasma partition coefficients. Toxicology in Vitro, 2008. 22(2): p. 457-467.

Schmitt, W., Corrigendum to: "General approach for the calculation of tissue to plasma partition coefficients" [Toxicology in Vitro 22 (2008) 457-467]. Toxicology in Vitro, 2008. 22(6): p. 1666.

Poulin, P. and F.P. Theil, A priori prediction of tissue: plasma partition coefficients of drugs to facilitate the use of physiologically based pharmacokinetic models in drug discovery. Journal of pharmaceutical sciences, 2000. 89(1): p. 16-35.

Rodgers, T. and M. Rowland, Physiologically based pharmacokinetic modelling 2: predicting the tissue distribution of acids, very weak bases, neutrals and zwitterions. Journal of pharmaceutical sciences, 2006. 95(6): p. 1238-1257.

Rodgers, T., D. Leahy, and M. Rowland, Physiologically based pharmacokinetic modeling 1: predicting the tissue distribution of moderate-to-strong bases. Journal of pharmaceutical sciences, 2005. 94(6): p. 1259-1276.

Rodgers, T., D. Leahy, and M. Rowland, Tissue distribution of basic drugs: Accounting for enantiomeric, compound and regional differences amongst beta-blocking drugs in rat. Journal of pharmaceutical sciences, 2005. 94(6): p. 1237-1248.

Gueorguieva, I., et al., Development of a whole body physiologically based model to characterise the pharmacokinetics of benzodiazepines. 1: Estimation of rat tissue-plasma partition ratios. Journal of pharmacokinetics and pharmacodynamics, 2004. 31(4): p. 269-298.

Poulin, P., K. Schoenlein, and F.P. Theil, Prediction of adipose tissue: plasma partition coefficients for structurally unrelated drugs. Journal of pharmaceutical sciences, 2001. 90(4): p. 436-447.

Bjorkman, S., Prediction of the volume of distribution of a drug: which tissue-plasma partition coefficients are needed? Journal of pharmacy and pharmacology, 2002. 54(9): p. 1237-1245.

Yun, Y. and A. Edginton, Correlation-based prediction of tissue-to-plasma partition coefficients using readily available input parameters. Xenobiotica, 2013. 43(10): p. 839-852.

Uchimura, T., et al., Prediction of human blood-to-plasma drug concentration ratio. Biopharmaceutics & drug disposition, 2010. 31(5-6): p. 286-297.

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pearce2017regression Pearce et al. 2017 data

### **Description**

This table includes the adjusted and unadjusted regression parameter estimates for the chemical-specifc plasma protein unbound fraction (fup) in 12 different tissue types.

## Usage

pearce2017regression

#### **Format**

data.frame

#### **Details**

Predictions were made with regression models, as reported in Pearce et al. (2017).

## Author(s)

Robert G. Pearce

### **Source**

Pearce et al. 2017 Regression Models

# References

Pearce, Robert G., et al. "Evaluation and calibration of high-throughput predictions of chemical distribution to tissues." Journal of pharmacokinetics and pharmacodynamics 44.6 (2017): 549-565.

pharma DRUGS\NORMAN: Pharmaceutical List with EU, Swiss, US Consumption Data

## **Description**

SWISSPHARMA is a list of pharmaceuticals with consumption data from Switzerland, France, Germany and the USA, used for a suspect screening/exposure modelling approach described in Singer et al 2016, DOI: 10.1021/acs.est.5b03332. The original data is available on the NORMAN Suspect List Exchange.

### Usage

pharma

### **Format**

An object of class data. frame with 954 rows and 14 columns.

physiology.data 163

#### **Source**

https://comptox.epa.gov/dashboard/chemical_lists/swisspharma

#### References

Wambaugh et al. (2019) "Assessing Toxicokinetic Uncertainty and Variability in Risk Prioritization", Toxicological Sciences, 172(2), 235-251.

physiology.data

Species-specific physiology parameters

## **Description**

This data set contains values from Davies and Morris (1993) necessary to paramaterize a toxicokinetic model for human, mouse, rat, dog, or rabbit. The temperature for each species are taken from Robertshaw et al. (2004), Gordon (1993), and Stammers(1926).

# Usage

physiology.data

### **Format**

A data.frame containing 11 rows and 7 columns.

# Author(s)

John Wambaugh and Nisha Sipes

### **Source**

Wambaugh, John F., et al. "Toxicokinetic triage for environmental chemicals." Toxicological Sciences (2015): 228-237.

### References

Davies, B. and Morris, T. (1993). Physiological Parameters in Laboratory Animals and Humans. Pharmaceutical Research 10(7), 1093-1095, 10.1023/a:1018943613122.

Environment, in Dukes' Physiology of Domestic Animals, 12th ed., Reece W.O., Ed. Copyright 2004 by Cornell University. Stammers (1926) The blood count and body temperature in normal rats Gordon (1993) Temperature Regulation in Laboratory Rodents

pradeep2020

pksim.pcs

Partition Coefficients from PK-Sim

### **Description**

Dallmann et al. (2018) made use of PK-Sim to predict chemical- and tissue- specific partition coefficients. The methods include both the default PK-Sim approach and PK-Sim Standard and Rodgers & Rowland (2006).

## Usage

pksim.pcs

#### **Format**

data.frame

#### **Source**

Kapraun et al. 2021 (submitted)

### References

Dallmann A, Ince I, Coboeken K, Eissing T, Hempel G (2018). "A physiologically based pharmacokinetic model for pregnant women to predict the pharmacokinetics of drugs metabolized via several enzymatic pathways." *Clinical pharmacokinetics*, **57**(6), 749–768.

pradeep2020

Pradeep et al. 2020

## **Description**

This table includes Support Vector Machine and Random Forest model predicted values for unbound fraction plasma protein (fup) and intrinsic hepatic clearance (clint) values for a subset of chemicals in the Tox21 library (see https://www.epa.gov/chemical-research/toxicology-testing-21st-century).

### Usage

pradeep2020

### **Format**

data.frame

### **Details**

Prediction were made with Support Vector Machine and Random Forest models, as reported in Pradeep et al. (2020).

#### **Source**

Pradeep et al. 2020 Chemical Structure Predictive Models for HTTK

# References

Pradeep P, Patlewicz G, Pearce R, Wambaugh J, Wetmore B, Judson R (2020). "Using chemical structure information to develop predictive models for in vitro toxicokinetic parameters to inform high-throughput risk-assessment." *Computational Toxicology*, **16**, 100136. ISSN 2468-1113, doi: 10.1016/j.comtox.2020.100136, https://www.sciencedirect.com/science/article/pii/S2468111320300463.

```
predict_partitioning_schmitt
```

Predict partition coefficients using the method from Schmitt (2008).

# Description

This function implements the method from Schmitt (2008) in predicting the tissue to unbound plasma partition coefficients for the tissues contained in the tissue.data table.

## Usage

```
predict_partitioning_schmitt(
  chem.name = NULL,
  chem.cas = NULL,
  dtxsid = NULL,
  species = "Human",
  model = "pbtk",
  default.to.human = FALSE,
  parameters = NULL,
  alpha = 0.001,
  adjusted.Funbound.plasma = TRUE,
  regression = TRUE,
 regression.list = c("brain", "adipose", "gut", "heart", "kidney", "liver", "lung",
    "muscle", "skin", "spleen", "bone"),
  tissues = NULL,
  minimum.Funbound.plasma = 1e-04,
  suppress.messages = FALSE
)
```

### **Arguments**

chem.name	Either the chemical name or the CAS number must be specified.
chem.cas	Either the chemical name or the CAS number must be specified.
dtxsid	$EPA's \ DSSTox \ Structure \ ID \ (https://comptox.epa.gov/dashboard) \ the \ chemical \ must \ be \ identified \ by \ either \ CAS, \ name, \ or \ DTXSIDs$
species	Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").
model	Model for which partition coefficients are needed (for example, "pbtk", "3compartment")

default.to.human

Substitutes missing animal values with human values if true (hepatic intrinsic

clearance or fraction of unbound plasma).

parameters Chemical parameters from parameterize_schmitt overrides chem.name, dtxsid,

and chem.cas.

alpha Ratio of Distribution coefficient D of totally charged species and that of the

neutral form

adjusted.Funbound.plasma

Whether or not to use Funbound.plasma adjustment.

regression Whether or not to use the regressions. Regressions are used by default.

regression.list

Tissues to use regressions on.

tissues Vector of desired partition coefficients. Returns all by default.

minimum.Funbound.plasma

Monte Carlo draws less than this value are set equal to this value (default is

0.0001 – half the lowest measured Fup in our dataset).

suppress.messages

Whether or not the output message is suppressed.

#### **Details**

A separate regression is used when adjusted. Funbound. plasma is FALSE.

A regression is used for membrane affinity when not provided. The regressions for correcting each tissue are performed on tissue plasma partition coefficients (Ktissue2pu * Funbound.plasma) calculated with the corrected Funbound.plasma value and divided by this value to get Ktissue2pu. Thus the regressions should be used with the corrected Funbound.plasma.

The red blood cell regression can be used but is not by default because of the span of the data used, reducing confidence in the regression for higher and lower predicted values.

Human tissue volumes are used for species other than Rat.

### Value

Returns tissue to unbound plasma partition coefficients for each tissue.

### Author(s)

Robert Pearce

### References

Schmitt, Walter. "General approach for the calculation of tissue to plasma partition coefficients." Toxicology in Vitro 22.2 (2008): 457-467.

Birnbaum, L., et al. "Physiological parameter values for PBPK models." International Life Sciences Institute, Risk Science Institute, Washington, DC (1994).

Pearce, Robert G., et al. "Evaluation and calibration of high-throughput predictions of chemical distribution to tissues." Journal of pharmacokinetics and pharmacodynamics 44.6 (2017): 549-565.

Yun, Y. E., and A. N. Edginton. "Correlation-based prediction of tissue-to-plasma partition coefficients using readily available input parameters." Xenobiotica 43.10 (2013): 839-852.

pregnonpregaucs 167

### **Examples**

```
predict_partitioning_schmitt(chem.name='ibuprofen',regression=FALSE)
```

pregnonpregaucs

AUCs for Pregnant and Non-Pregnant Women

### **Description**

Dallmann et al. (2018) includes compiled literature descriptions of toxicokinetic summary statistics, including time-integrated plasma concentrations (area under the curve or AUC) for drugs administered to a sample of subjects including both pregnant and non-pregnant women. The circumstances of the dosing varied slightly between drugs and are summarized in the table.

### Usage

pregnonpregaucs

#### **Format**

data.frame

#### **Source**

Kapraun et al. 2021 (submitted)

### References

Dallmann A, Ince I, Coboeken K, Eissing T, Hempel G (2018). "A physiologically based pharmacokinetic model for pregnant women to predict the pharmacokinetics of drugs metabolized via several enzymatic pathways." *Clinical pharmacokinetics*, **57**(6), 749–768.

```
propagate_invitrouv_1comp
```

Propagates uncertainty and variability in in vitro HTTK data into one compartment model parameters

# Description

Propagates uncertainty and variability in in vitro HTTK data into one compartment model parameters

# Usage

```
propagate_invitrouv_1comp(parameters.dt, ...)
```

# **Arguments**

```
parameters.dt The data table of parameters being used by the Monte Carlo sampler
... Additional arguments passed to calc_elimination_rate
```

#### Value

A data.table whose columns are the parameters of the HTTK model specified in model.

### Author(s)

John Wambaugh

```
propagate_invitrouv_3comp
```

Propagates uncertainty and variability in in vitro HTTK data into three compartment model parameters

## **Description**

Propagates uncertainty and variability in in vitro HTTK data into three compartment model parameters

# Usage

```
propagate_invitrouv_3comp(parameters.dt, ...)
```

### **Arguments**

```
parameters.dt The data table of parameters being used by the Monte Carlo sampler
... Additional arguments passed to calc_hep_clearance
```

### Value

A data.table whose columns are the parameters of the HTTK model specified in model.

### Author(s)

John Wambaugh

```
propagate_invitrouv_pbtk
```

Propagates uncertainty and variability in in vitro HTTK data into PBPK model parameters

## **Description**

Propagates uncertainty and variability in in vitro HTTK data into PBPK model parameters

## Usage

```
propagate_invitrouv_pbtk(parameters.dt, ...)
```

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### **Arguments**

parameters.dt The data table of parameters being used by the Monte Carlo sampler
... Additional arguments passed to calc_hep_clearance

### Value

A data.table whose columns are the parameters of the HTTK model specified in model.

### Author(s)

John Wambaugh

reset_httk

Reset HTTK to Default Data Tables

# Description

This function returns an updated version of chem.physical_and_invitro.data that includes data predicted with Simulations Plus' ADMET predictor that was used in Sipes et al. 2017, included in admet.data.

### Usage

```
reset_httk(target.env = .GlobalEnv)
```

## **Arguments**

target.env

The environment where the new chem.physical_and_invitro.data is loaded. Defaults to global environment.

### Value

data.frame

The package default version of chem.physical_and_invitro.data.

## Author(s)

John Wambaugh

# Examples

```
chem.physical_and_invitro.data <- load_sipes2017()
reset_httk()</pre>
```

r_left_censored_norm

rfun

Randomly draws from a one-dimensional KDE

### **Description**

Randomly draws from a one-dimensional KDE

### Usage

```
rfun(n, fhat)
```

## **Arguments**

n Number of samples to draw

fhat A list with elements x, w, and h (h is the KDE bandwidth).

## Value

A vector of n samples from the KDE fhat

## Author(s)

Caroline Ring

## References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

```
{\tt r\_left\_censored\_norm} \quad \textit{Returns draws from a normal distribution with a lower censoring limit} \\ \quad of lod (limit of detection)
```

# Description

Returns draws from a normal distribution with a lower censoring limit of lod (limit of detection)

# Usage

```
r_left_censored_norm(n, mean = 0, sd = 1, lod = 0.005, lower = 0, upper = 1)
```

# Arguments

n	Number of samples to take
mean	Mean of censored distribution. Default 0.
sd	Standard deviation of censored distribution. Default 1.
lod	Bound below which to censor. Default 0.005.
lower	Lower bound on censored distribution. Default 0.
upper	Upper bound on censored distribution. Default 1.

scale_dosing 171

### Value

A vector of samples from the specified censored distribution.

scale_dosing

Scale mg/kg body weight doses according to body weight and units

## **Description**

This function transforms the dose (in mg/kg) into the appropriate units. It handles single doses, matrices of doses, or daily repeated doses at varying intervals. Gut absorption is also factored in through the parameter Fgutabs, and scaling is currently avoided in the inhalation exposure case with a scale factor of 1

# Usage

```
scale_dosing(
  dosing,
  parameters,
  route,
  input.units = NULL,
  output.units = "uM",
  vol = NULL
)
```

### **Arguments**

dosing	List of dosing metrics used in simulation, which must include the general entries with names "initial.dose", "doses.per.day", "daily.dose", and "dosing.matrix". The "dosing.matrix" is used for more precise dose regimen specification, and is a matrix consisting of two columns or rows named "time" and "dose" containing the time and amount, in mg/kg BW, of each dose. The minimal usage case involves all entries but "initial.dose" set to NULL in value.
parameters	Chemical parameters from parameterize_pbtk function, overrides chem.name and chem.cas.
route	String specification of route of exposure for simulation: "oral", "iv", "inhalation",
input.units	Units of the dose values being scaled. (Default is NULL.) Currently supported units "mg/L", "ug/L", "ug/mL", "uM", "umol/L", "ug/dL", "ug/g", "nmol/L", "nM", and "ppmw" (supported input.units subject to change).
output.units	Desired units (either "mg/L", "mg", "umol", or default "uM").
vol	Volume for the target tissue of interest. NOTE: Volume should not be in units of per BW, i.e. "kg".

# Value

A list of numeric values for doses converted to output.units, potentially (depending on argument dosing) including:

initial.dose The first dose given dosing.matrix A 2xN matrix where the first column is dose time and the second is dose amount for N doses The total cumulative daily dose

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### Author(s)

John Wambaugh and Sarah E. Davidson

set_httk_precision

set_httk_precision

## **Description**

Although the ODE solver and other functions return very precise numbers, we cannot (or at least do not spend enough computing time to) be sure of the precioion to an arbitrary level. This function both limits the number of signficant figures reported and truncates the numerical precision.

# Usage

```
set_httk_precision(in.num, sig.fig = 4, num.prec = 9)
```

## **Arguments**

in.num The numeric variable (or assembly of numerics) to be processed.

sig.fig The number of significant figures reported. Defaults to 4.

num.prec The precision maintained, digits below 10^num.prec are dropped. Defaults to 9.

### Value

numeric values

### Author(s)

John Wambaugh

sipes2017

Sipes et al. 2017 data

# Description

This table includes in silico predicted chemical-specifc plasma protein unbound fraction (fup) and intrinsic hepatic clearance values for the entire Tox21 library (see <a href="https://www.epa.gov/chemical-research/toxicology-testing-21st-century-tox21">https://www.epa.gov/chemical-research/toxicology-testing-21st-century-tox21</a>). Predictions were made with Simulations Plus ADMET predictor, as reported in Sipes et al. (2017).

# Usage

sipes2017

## Format

data.frame

skeletal_muscle_mass 173

### Author(s)

Nisha Sipes

#### Source

ADMET, Simulations Plus

#### References

Sipes, Nisha S., et al. "An Intuitive Approach for Predicting Potential Human Health Risk with the Tox21 10k Library." Environmental Science & Technology 51.18 (2017): 10786-10796.

### **Description**

Predict skeletal muscle mass from age, height, and gender.

### Usage

```
skeletal_muscle_mass(smm, age_years, height, gender)
```

### **Arguments**

smm Vector of allometrically-scaled skeletal muscle masses.

age_years Vector of ages in years. height Vector of heights in cm.

gender Vector of genders, either 'Male' or 'Female.'

## **Details**

For individuals over age 18, use allometrically-scaled muscle mass with an age-based scaling factor, to account for loss of muscle mass with age (Janssen et al. 2000). For individuals under age 18, use skeletal_muscle_mass_children.

# Value

Vector of skeletal muscle masses in kg.

## Author(s)

Caroline Ring

# References

Janssen, Ian, et al. "Skeletal muscle mass and distribution in 468 men and women aged 18-88 yer." Journal of Applied Physiology 89.1 (2000): 81-88

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

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#### See Also

skeletal_muscle_mass_children

```
skeletal_muscle_mass_children
```

Predict skeletal muscle mass for children

### **Description**

For individuals under age 18, predict skeletal muscle mass from gender and age, using a nonlinear equation from Webber and Barr (2012)

# Usage

```
skeletal_muscle_mass_children(gender, age_years)
```

## **Arguments**

gender Vector of genders (either 'Male' or 'Female').

age_years Vector of ages in years.

#### Value

Vector of skeletal muscle masses in kg.

# Author(s)

Caroline Ring

#### References

Webber, Colin E., and Ronald D. Barr. "Age-and gender-dependent values of skeletal muscle mass in healthy children and adolescents." Journal of cachexia, sarcopenia and muscle 3.1 (2012): 25-29.

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

skin_mass_bosgra

Predict skin mass

# Description

Using equation from Bosgra et al. 2012, predict skin mass from body surface area.

### Usage

```
skin_mass_bosgra(BSA)
```

### **Arguments**

BSA

Vector of body surface areas in cm².

### Value

Vector of skin masses in kg.

#### Author(s)

Caroline Ring

#### References

Bosgra, Sieto, et al. "An improved model to predict physiologically based model parameters and their inter-individual variability from anthropometry." Critical reviews in toxicology 42.9 (2012): 751-767.

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

solve_1comp

Solve one compartment TK model

### **Description**

This function solves for the amount or concentration of a chemical in plasma for a one compartment model as a function of time based on the dose and dosing frequency.

### Usage

```
solve_1comp(
 chem.name = NULL,
 chem.cas = NULL,
 dtxsid = NULL,
  times = NULL,
 parameters = NULL,
 days = 10,
  tsteps = 4,
 daily.dose = NULL,
 dose = NULL,
 doses.per.day = NULL,
  initial.values = NULL,
 plots = FALSE,
  suppress.messages = FALSE,
  species = "Human",
  iv.dose = FALSE,
  input.units = "mg/kg",
 output.units = NULL,
 method = "lsoda",
 rtol = 1e-08,
 atol = 1e-12,
 default.to.human = FALSE,
 recalc.blood2plasma = FALSE,
 recalc.clearance = FALSE,
 dosing.matrix = NULL,
```

```
adjusted.Funbound.plasma = TRUE,
regression = TRUE,
restrictive.clearance = TRUE,
minimum.Funbound.plasma = 1e-04,
monitor.vars = NULL,
...
)
```

### **Arguments**

chem. name Either the chemical name, CAS number, or the parameters must be specified. Either the chemical name, CAS number, or the parameters must be specified.

dtxsid EPA's 'DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chem-

ical must be identified by either CAS, name, or DTXSIDs

times Optional time sequence for specified number of days.

parameters Chemical parameters from parameterize_1comp function, overrides chem.name

and chem.cas.

days Length of the simulation.

tsteps The number time steps per hour.

daily.dose Total daily dose, default is mg/kg BW.

dose Amount of a single dose, default is mg/kg BW.

doses.per.day Number of doses per day.

initial.values Vector containing the initial concentrations or amounts of the chemical in spec-

ified tissues with units corresponding to output.units. Defaults are zero.

plots Plots all outputs if true.

suppress.messages

Whether or not the output message is suppressed.

species Species desired (either "Rat", "Rabbit", "Dog", or default "Human").

iv. dose Simulates a single i.v. dose if true.

input units Input units of interest assigned to dosing, defaults to "mg/kg" BW.

output.units A named vector of output units expected for the model results. Default, NULL,

returns model results in units specified in the 'modelinfo' file. See table below

for details.

method Method used by integrator (deSolve).

rtol Argument passed to integrator (deSolve).

atol Argument passed to integrator (deSolve).

default.to.human

Substitutes missing rat values with human values if true.

recalc.blood2plasma

Whether or not to recalculate the blood:plasma chemical concentration ratio

recalc.clearance

Whether or not to recalculate the elimination rate.

dosing.matrix Vector of dosing times or a matrix consisting of two columns or rows named

"dose" and "time" containing the time and amount, in mg/kg BW by default, of

each dose.

adjusted.Funbound.plasma

Uses adjusted Funbound.plasma when set to TRUE along with volume of distribution calculated with this value.

regression

Whether or not to use the regressions in calculating partition coefficients in volume of distribution calculation.

restrictive.clearance

In calculating elimination rate, protein binding is not taken into account (set to 1) in liver clearance if FALSE.

minimum.Funbound.plasma

Monte Carlo draws less than this value are set equal to this value (default is 0.0001 – half the lowest measured Fup in our dataset).

monitor.vars

Which variables are returned as a function of time. Defaults value of NULL provides "Agutlumen", "Ccompartment", "Ametabolized", "AUC"

... Additional arguments passed to the integrator.

# **Details**

Note that the model parameters have units of hours while the model output is in days.

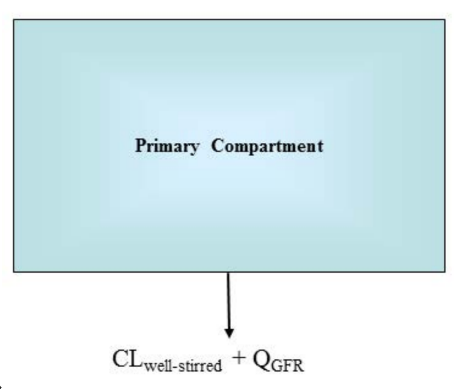
Default value of NULL for doses.per.day solves for a single dose.

When species is specified as rabbit, dog, or mouse, the function uses the appropriate physiological data(volumes and flows) but substitutes human fraction unbound, partition coefficients, and intrinsic hepatic clearance.

AUC is area under plasma concentration curve.

Model Figure





altalt

# Value

A matrix with a column for time(in days) and a column for the compartment and the area under the curve (concentration only).

# Author(s)

Robert Pearce

# References

Pearce, Robert G., et al. "Httk: R package for high-throughput toxicokinetics." Journal of statistical software 79.4 (2017): 1.

## **Examples**

```
solve_1comp(chem.name='Bisphenol-A',days=1)
params <- parameterize_1comp(chem.cas="80-05-7")
solve_1comp(parameters=params)</pre>
```

solve_3comp

Solve_3comp

## **Description**

This function solves for the amounts or concentrations of a chemical in different tissues as functions of time based on the dose and dosing frequency. It uses a three compartment model with partition coefficients.

## Usage

```
solve_3comp(
  chem.name = NULL,
  chem.cas = NULL,
  dtxsid = NULL,
  times = NULL,
  parameters = NULL,
  days = 10,
  tsteps = 4,
  daily.dose = NULL,
  dose = NULL,
  doses.per.day = NULL,
  initial.values = NULL,
  plots = FALSE,
  suppress.messages = FALSE,
  species = "Human",
  iv.dose = FALSE,
  input.units = "mg/kg",
  output.units = NULL,
  method = "lsoda",
  rtol = 1e-08,
  atol = 1e-12,
  default.to.human = FALSE,
  recalc.blood2plasma = FALSE,
  recalc.clearance = FALSE,
  dosing.matrix = NULL,
  adjusted.Funbound.plasma = TRUE,
  regression = TRUE,
  restrictive.clearance = TRUE,
  minimum.Funbound.plasma = 1e-04,
  monitor.vars = NULL,
)
```

### **Arguments**

dtxsid

chem.name Either the chemical name, CAS number, or the parameters must be specified. Either the chemical name, CAS number, or the parameters must be specified. chem.cas

EPA's 'DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chemical must be identified by either CAS, name, or DTXSIDs

Optional time sequence for specified number of days. The dosing sequence times

begins at the beginning of times.

Chemical parameters from parameterize_3comp function, overrides chem.name parameters

and chem.cas.

days Length of the simulation.

tsteps The number time steps per hour. daily.dose Total daily dose, mg/kg BW.

dose Amount of a single dose, mg/kg BW.

doses.per.day Number of doses per day.

Vector containing the initial concentrations or amounts of the chemical in specinitial.values

ified tissues with units corresponding to output.units. Defaults are zero.

plots Plots all outputs if true.

suppress.messages

Whether or not the output message is suppressed.

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

iv.dose Simulates a single i.v. dose if true.

Input units of interest assigned to dosing, defaults to mg/kg BW input.units

output.units A named vector of output units expected for the model results. Default, NULL,

returns model results in units specified in the 'modelinfo' file. See table below

for details.

method Method used by integrator (deSolve). Argument passed to integrator (deSolve). rtol atol Argument passed to integrator (deSolve).

default.to.human

Substitutes missing animal values with human values if true (hepatic intrinsic

clearance or fraction of unbound plasma).

recalc.blood2plasma

Recalculates the ratio of the amount of chemical in the blood to plasma using the input parameters, calculated with hematocrit, Funbound.plasma, and Krbc2pu.

recalc.clearance

Recalculates the the hepatic clearance (Clmetabolism) with new million.cells.per.gliver

parameter.

dosing.matrix Vector of dosing times or a matrix consisting of two columns or rows named

"dose" and "time" containing the time and amount, in mg/kg BW, of each dose.

adjusted.Funbound.plasma

Uses adjusted Funbound.plasma when set to TRUE along with partition coeffi-

cients calculated with this value.

Whether or not to use the regressions in calculating partition coefficients. regression

restrictive.clearance

Protein binding not taken into account (set to 1) in liver clearance if FALSE.

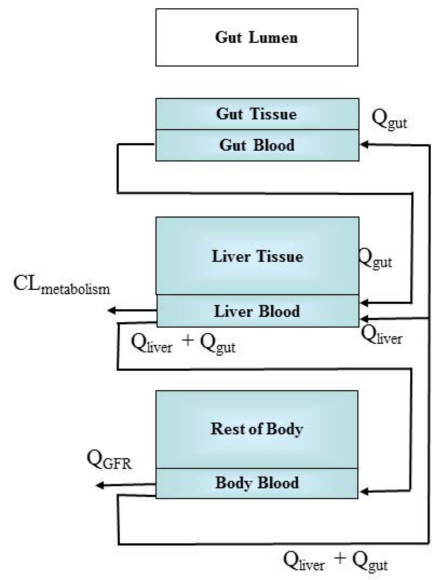
181 solve_3comp minimum.Funbound.plasma Monte Carlo draws less than this value are set equal to this value (default is 0.0001 – half the lowest measured Fup in our dataset). monitor.vars Which variables are returned as a function of time. Defaults value of NULL provides "Cliver", "Csyscomp", "Atubules", "Ametabolized", "AUC" Additional arguments passed to the integrator. **Details** Note that the model parameters have units of hours while the model output is in days. Default of NULL for doses.per.day solves for a single dose.

The compartments used in this model are the gutlumen, gut, liver, and rest-of-body, with the plasma

equivalent to the liver plasma.

Model Figure

solve_3comp



altalt

When species is specified as rabbit, dog, or mouse, the function uses the appropriate physiological data(volumes and flows) but substitues human fraction unbound, partition coefficients, and intrinsic hepatic clearance.

# Value

A matrix of class deSolve with a column for time(in days) and each compartment, the plasma concentration, area under the curve, and a row for each time point.

# Author(s)

John Wambaugh and Robert Pearce

solve_fetal_pbtk 183

#### References

Pearce, Robert G., et al. "Httk: R package for high-throughput toxicokinetics." Journal of statistical software 79.4 (2017): 1.

### **Examples**

```
solve_3comp(chem.name='Bisphenol-A',doses.per.day=2,daily.dose=.5,days=1,tsteps=2)
params <-parameterize_3comp(chem.cas="80-05-7")
solve_3comp(parameters=params)</pre>
```

solve_fetal_pbtk

Solve_fetal_PBTK

# **Description**

This function solves for the amounts or concentrations in uM of a chemical in different tissues of a maternofetal system as functions of time based on the dose and dosing frequency.

# Usage

```
solve_fetal_pbtk(
  chem.name = NULL,
 chem.cas = NULL,
 dtxsid = NULL,
  times = seq(13 * 7, 40 * 7, 1),
 parameters = NULL,
 days = NULL,
  species = "human",
  tsteps = 4,
 dose = NULL,
 dosing.matrix = NULL,
 daily.dose = NULL,
  doses.per.day = NULL,
  initial.values = NULL,
 plots = FALSE,
  suppress.messages = FALSE,
  iv.dose = FALSE,
  input.units = "mg/kg",
 output.units = NULL,
 method = "lsoda",
 rtol = 1e-08,
 atol = 1e-12,
 default.to.human = FALSE,
 recalc.blood2plasma = FALSE,
  recalc.clearance = FALSE,
  adjusted.Funbound.plasma = TRUE,
  regression = TRUE,
  restrictive.clearance = TRUE,
```

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```
minimum.Funbound.plasma = 1e-04,
monitor.vars = NULL,
...
)
```

### **Arguments**

chem. name Either the chemical name, CAS number, or the parameters must be specified.

chem. cas Either the chemical name, CAS number, or the parameters must be specified.

dtxsid EPA's DSSTox Structure ID (http://comptox.epa.gov/dashboard) the chem-

ical must be identified by either CAS, name, or DTXSIDs

times Optional time sequence in days. Dosing sequence begins at the beginning of

times. Default is from 13th week of pregnancy to 40th due to data constraints.

parameters Chemical parameters from parameterize_fetal_pbtk function, overrides chem.name

and chem.cas.

days Length of the simulation.

species Included for compatibility with other functions, but the model will not run for

non-human species (default "Human").

tsteps The number time steps per hour. Default of 4.

dose Amount of a single dose, mg/kg BW.

dosing.matrix A matrix of either one column (or row) with a set of dosing times or with two

columns (or rows) correspondingly named "dose" and "time" containing the time

and amount, in mg/kg BW, of each dose.

daily.dose Total daily dose, mg.
doses.per.day Number of doses per day.

initial.values Vector containing the initial concentrations or amounts of the chemical in spec-

ified tissues with units corresponding to compartment.units. Defaults are zero.

plots Plots all outputs if true.

suppress.messages

Whether or not the output message is suppressed.

iv. dose Simulates a single i.v. dose if true.

input.units Input units of interest assigned to dosing, defaults to mg/kg BW

output.units A named vector of output units expected for the model results. Default, NULL,

returns model results in units specified in the 'modelinfo' file. See table below

for details.

method Method used by integrator (deSolve).

rtol Argument passed to integrator (deSolve).

Argument passed to integrator (deSolve).

default.to.human

Substitutes missing animal values with human values if true (hepatic intrinsic

clearance or fraction of unbound plasma).

recalc.blood2plasma

Recalculates the ratio of the amount of chemical in the blood to plasma using the input parameters, calculated with hematocrit, Funbound.plasma, and Krbc2pu.

recalc.clearance

Recalculates the the hepatic clearance (Clmetabolism) with new million.cells.per.gliver parameter.

solve_fetal_pbtk 185

adjusted.Funbound.plasma

Uses adjusted Funbound.plasma when set to TRUE along with partition coefficients calculated with this value.

regression Whether or not to use the regressions in calculating partition coefficients.

restrictive.clearance

Protein binding not taken into account (set to 1) in liver clearance if FALSE.

minimum.Funbound.plasma

Monte Carlo draws less than this value are set equal to this value (default is 0.0001 – half the lowest measured Fup in our dataset).

monitor.vars Which variables to track by default

... Additional arguments passed to the integrator.

#### **Details**

The stage of pregnancy simulated here begins by default at the 13th week due to a relative lack of data to support parameterization prior, in line with the recommendations of Kapraun et al. 2019 ("Empirical models for anatomical and physiological..."), and ends at the 40th week of pregnancy.

Note that the model parameters have units of hours while the model output is in days. Dose is in mg, not scaled for body weight.

Default NULL value for doses.per.day solves for a single dose.

The maternal compartments used in this model are the gut lumen, gut, liver, venous blood, arterial blood, lung, adipose tissue, kidney, thyroid, and rest of body. A placenta is modeled as a joint organ shared by mother and fetus, through which chemical exchange can occur with the fetus. Fetal compartments include arterial blood, venous blood, kidney, thyroid, liver, lung, gut, brain, and rest of body.

The extra compartments include the amounts or concentrations metabolized by the liver and excreted by the kidneys through the tubules.

AUC is the area under the curve of the plasma concentration.

This gestational model is only parameterized for humans.

# Value

A matrix of class deSolve with a column for time(in days), each compartment, the area under the curve, and plasma concentration and a row for each time point.

# Author(s)

John Wambaugh, Mark Sfeir, and Dustin Kapraun

### **Examples**

```
out = solve_fetal_pbtk(chem.name = 'bisphenol a', daily.dose = 1,
doses.per.day = 3, plots = TRUE)
```

solve_gas_pbtk

solve_gas_pbtk

# **Description**

This function solves for the amounts or concentrations of a chemical in different tissues as functions of time as a result of inhalation exposure to an ideal gas.

# Usage

```
solve_gas_pbtk(
 chem.name = NULL,
  chem.cas = NULL,
 dtxsid = NULL,
 parameters = NULL,
  times = NULL,
 days = 10,
  tsteps = 4,
 daily.dose = NULL,
 doses.per.day = NULL,
 dose = NULL,
 dosing.matrix = NULL,
 forcings = NULL,
 exp.start.time = 0,
 exp.conc = 1,
 period = 24,
 exp.duration = 12,
 initial.values = NULL,
 plots = FALSE,
  suppress.messages = FALSE,
  species = "Human",
  iv.dose = FALSE,
  input.units = "ppmv",
 output.units = NULL,
 method = "lsoda",
 rtol = 1e-08,
 atol = 1e-12,
 default.to.human = FALSE,
 recalc.blood2plasma = FALSE,
  recalc.clearance = FALSE,
 adjusted.Funbound.plasma = TRUE,
 regression = TRUE,
 restrictive.clearance = TRUE,
 minimum.Funbound.plasma = 1e-04,
 monitor.vars = NULL,
 vmax = 0,
 km = 1,
 exercise = FALSE,
  fR = 12,
 VT = 0.75,
 VD = 0.15,
```

)

### **Arguments**

chem. name Either the chemical name, CAS number, or the parameters must be specified.

chem. cas Either the chemical name, CAS number, or the parameters must be specified.

dtxsid EPA's DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chem-

ical must be identified by either CAS, name, or DTXSIDs

parameters Chemical parameters from parameterize_gas_pbtk (or other bespoke) function,

overrides chem.name and chem.cas.

times Optional time sequence for specified number of days. Dosing sequence begins

at the beginning of times.

days Length of the simulation.

tsteps The number of time steps per hour.

daily.dose Total daily dose

doses.per.day Number of doses per day.

Amount of a single dose

dosing.matrix Vector of dosing times or a matrix consisting of two columns or rows named

"dose" and "time" containing the time and amount of each dose.

forcings Manual input of 'forcings' data series argument for ode integrator. If left un-

specified, 'forcings' defaults to NULL, and then other input parameters (see exp.start.time, exp.conc, exp.duration, and period) provide the necessary infor-

mation to assemble a forcings data series.

exp. start.time Start time in specifying forcing exposure series, default 0.

exp.conc Specified inhalation exposure concentration for use in assembling "forcings"

data series argument for integrator. Defaults to units of ppmv.

period For use in assembling forcing function data series 'forcings' argument, specified

in hours

exp. duration For use in assembling forcing function data series 'forcings' argument, specified

in hours

initial.values Vector containing the initial concentrations or amounts of the chemical in spec-

ified tissues with units corresponding to those specified for the model outputs.

Default values are zero.

plots Plots all outputs if true.

suppress.messages

Whether or not the output message is suppressed.

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

iv. dose Simulates a single i.v. dose if true.

input.units Input units of interest assigned to dosing, including forcings. Defaults to "ppmv"

as applied to the default forcings scheme.

output.units A named vector of output units expected for the model results. Default, NULL,

returns model results in units specified in the 'modelinfo' file. See table below

for details.

method Method used by integrator (deSolve).

rtol Argument passed to integrator (deSolve).

atol Argument passed to integrator (deSolve).

default.to.human

Substitutes missing animal values with human values if true (hepatic intrinsic clearance or fraction of unbound plasma).

recalc.blood2plasma

Recalculates the ratio of the amount of chemical in the blood to plasma using the input parameters, calculated with hematocrit, Funbound.plasma, and Krbc2pu.

recalc.clearance

Recalculates the hepatic clearance (Clmetabolism) with new million.cells.per.gliver parameter.

adjusted.Funbound.plasma

Uses adjusted Funbound.plasma when set to TRUE along with partition coefficients calculated with this value.

regression Whether or not to use the regressions in calculating partition coefficients.

restrictive.clearance

Protein binding not taken into account (set to 1) in liver clearance if FALSE.

minimum.Funbound.plasma

Monte Carlo draws less than this value are set equal to this value (default is 0.0001 – half the lowest measured Fup in our dataset).

monitor.vars Which variables are returned as a function of time. Defaults value of NULL pro-

vides "Cgut", "Cliver", "Cven", "Clung", "Cart", "Crest", "Ckidney", "Cplasma", "Calv", "Cendexh", "Cmixexh", "Cmuc", "Atubules", "Ametabolized", "AUC"

vmax Michaelis-Menten vmax value in reactions/min

km Michaelis-Menten concentration of half-maximal reaction velocity in desired

output concentration units.

exercise Logical indicator of whether to simulate an exercise-induced heightened respi-

ration rate

fR Respiratory frequency (breaths/minute), used especially to adjust breathing rate

in the case of exercise. This parameter, along with VT and VD (below) gives another option for calculating Qalv (Alveolar ventilation) in case pulmonary

ventilation rate is not known

VT Tidal volume (L), to be modulated especially as part of simulating the state of

exercise

VD Anatomical dead space (L), to be modulated especially as part of simulating the

state of exercise

... Additional arguments passed to the integrator.

# **Details**

The default dosing scheme involves a specification of the start time of exposure (exp.start.time), the concentration of gas inhaled (exp.conc), the period of a cycle of exposure and non-exposure (period), the duration of the exposure during that period (exp.duration), and the total days simulated. Together, these arguments determine the "forcings" passed to the ODE integrator. Forcings can also be specified manually, or effectively turned off by setting exposure concentration to zero, if the user prefers to simulate dosing by other means.

The "forcings" object is configured to be passed to the integrator with, at the most, a basic unit conversion among ppmv, mg/L, and uM. No scaling by BW is set to be performed on the forcings series.

Note that the model parameters have units of hours while the model output is in days.

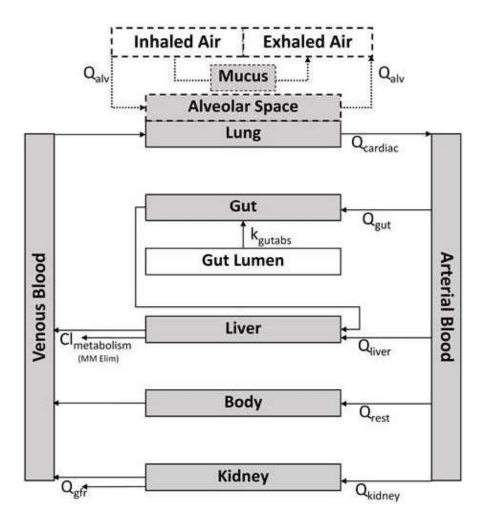
Default NULL value for doses.per.day solves for a single dose.

The compartments used in this model are the gut lumen, gut, liver, kidneys, veins, arteries, lungs, and the rest of the body.

The extra compartments include the amounts or concentrations metabolized by the liver and excreted by the kidneys through the tubules.

AUC is the area under the curve of the plasma concentration.

Model Figure from (Linakis et al. 2020):



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Model parameters are named according to the following convention:

prefix	suffic	Meaning
K		Partition coefficient for tissue to free plasma \ tab unitless
V		Volume

units

Q Flow L/h
Rate 1/h
c Parameter is proportional to body weight 1 / kg for volumes and 1/kg^(3/4) for flows

When species is specified but chemical-specific in vitro data are not available, the function uses the appropriate physiological data (volumes and flows) but default.to.human = TRUE must be used to substitute human fraction unbound, partition coefficients, and intrinsic hepatic clearance.

#### Value

A matrix of class deSolve with a column for time(in days), each compartment, the area under the curve, and plasma concentration and a row for each time point.

### Author(s)

Matt Linakis, John Wambaugh, Mark Sfeir, Miyuki Breen

#### References

Linakis MW, Sayre RR, Pearce RG, Sfeir MA, Sipes NS, Pangburn HA, Gearhart JM, Wambaugh JF (2020). "Development and evaluation of a high-throughput inhalation model for organic chemicals." *Journal of exposure science* & environmental epidemiology, **30**(5), 866–877.

Pearce RG, Setzer RW, Strope CL, Wambaugh JF, Sipes NS (2017). "Httk: R package for high-throughput toxicokinetics." *Journal of statistical software*, **79**(4), 1.

### **Examples**

```
solve_gas_pbtk(chem.name = 'pyrene', exp.conc = 1, period = 24, expduration = 24)

out <- solve_gas_pbtk(chem.name='pyrene',exp.conc = 0, doses.per.day = 2,
daily.dose = 3, input.units = "umol", plots=TRUE,initial.values=c(Aven=20))

out <- solve_gas_pbtk(chem.name = 'pyrene',exp.conc = 3, period = 24,
exp.duration = 6, exercise = TRUE)

params <- parameterize_gas_pbtk(chem.cas="80-05-7")
solve_gas_pbtk(parameters=params)

# Note that different model compartments for this model have different units
# and that the final units can be controlled with the output.units argument:
head(solve_gas_pbtk(chem.name="lindane"))
# Convert all compartment units to mg/L:
head(solve_gas_pbtk(chem.name="lindane",output.units="mg/L"))
# Convert just the plasma to mg/L:
head(solve_gas_pbtk(chem.name="lindane",output.units=list(Cplasma="mg/L")))</pre>
```

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solve_model

Solve_model

### **Description**

solve_model is designed to accept systematized metadata (provided by the model.list defined in the modelinfo files) for a given toxicokinetic model, including names of variables, parameterization functions, and key units, and use it along with chemical information to prepare an ode system for numerical solution over time of the amounts or concentrations of chemical in different bodily compartments of a given species (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

# Usage

```
solve_model(
  chem.name = NULL,
  chem.cas = NULL,
 dtxsid = NULL,
  times = NULL,
 parameters = NULL,
 model = NULL,
 route = "oral",
 dosing = NULL,
  days = 10,
  tsteps = 4,
  initial.values = NULL,
  initial.value.units = NULL,
 plots = FALSE,
 monitor.vars = NULL,
  suppress.messages = FALSE,
  species = "Human",
  input.units = "mg/kg",
 output.units = NULL,
 method = "lsoda",
  rtol = 1e-08,
  atol = 1e-12,
 recalc.blood2plasma = FALSE,
 recalc.clearance = FALSE,
 adjusted.Funbound.plasma = TRUE,
 minimum.Funbound.plasma = 1e-04,
 parameterize.arg.list = list(default.to.human = FALSE, clint.pvalue.threshold = 0.05,
    restrictive.clearance = TRUE, regression = TRUE),
)
```

### **Arguments**

chem. name Either the chemical name, CAS number, or the parameters must be specified.

chem. cas Either the chemical name, CAS number, or the parameters must be specified.

dtxsid EPA's DSSTox Structure ID (http://comptox.epa.gov/dashboard) the chemical must be identified by either CAS, name, or DTXSIDs

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times Optional time sequence for specified number of days. Dosing sequence begins

at the beginning of times.

parameters List of chemical parameters, as output by parameterize_pbtk function. Over-

rides chem.name and chem.cas.

model Specified model to use in simulation: "pbtk", "3compartment", "3compartmentss",

"1compartment", "schmitt", ...

route String specification of route of exposure for simulation: "oral", "iv", "inhala-

tion", ...

dosing List of dosing metrics used in simulation, which includes the namesake en-

tries of a model's associated dosing.params. In the case of most httk models, these should include "initial.dose", "doses.per.day", "daily.dose", and "dosing.matrix". The "dosing.matrix" is used for more precise dose regimen specification, and is a matrix consisting of two columns or rows named "time" and "dose" containing the time and amount of each dose. If none of the namesake entries of the dosing list is set to a non-NULL value, solve_model uses a default dose of 1 mg/kg BW along with the dose type (add/multiply) specified for a

given route (e.g. add the dose to gut lumen for oral route)

days Simulated period. Default 10 days.

tsteps The number of time steps per hour. Default of 4.

initial.values Vector of numeric values containing the initial concentrations or amounts of the

chemical in specified tissues with units corresponding to those specified for the

model outputs. Default values are zero.

initial.value.units

Vector of character strings containing the units corresponding to 'initial.values' specified for the model outputs. Default is assuming the units match expected

compartment units for the model.

plots Plots all outputs if true.

monitor.vars Which variables are returned as a function of time. Default values of NULL

looks up variables specified in modelinfo_MODEL.R

suppress.messages

Whether or not the output messages are suppressed.

species Species desired (models have been designed to be parameterized for some sub-

set of the following species: "Rat", "Rabbit", "Dog", "Mouse", or default "Hu-

man").

input units Input units of interest assigned to dosing. Defaults to mg/kg BW, in line with the

default dosing scheme of a one-time dose of 1 mg/kg in which no other dosing

parameters are specified.

output units Output units of interest for the compiled components. Defaults to NULL, and

will provide values in model units if unspecified.

method Method used by integrator (deSolve).

rtol Argument passed to integrator (deSolve).

atol Argument passed to integrator (deSolve).

recalc.blood2plasma

Recalculates the ratio of the amount of chemical in the blood to plasma using the input parameters, calculated with hematocrit, Funbound.plasma, and Krbc2pu.

recalc.clearance

Recalculates the the hepatic clearance (Clmetabolism) with new million.cells.per.gliver parameter.

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adjusted.Funbound.plasma

Uses adjusted Funbound.plasma when set to TRUE along with partition coefficients calculated with this value.

minimum.Funbound.plasma

Monte Carlo draws less than this value are set equal to this value (default is 0.0001 – half the lowest measured Fup in our dataset)

parameterize.arg.list

Additional parameterized passed to the model parameterization function.

. . . Additional arguments passed to the integrator.

default.to.human

Substitutes missing animal values with human values if true (hepatic intrinsic clearance or fraction of unbound plasma).

regression Whether or not to use the regressions in calculating partition coefficients. restrictive.clearance

Protein binding not taken into account (set to 1) in liver clearance if FALSE.

### **Details**

Dosing values with certain acceptable associated input.units (like mg/kg BW) are configured to undergo a unit conversion. All model simulations are intended to run with units as specifed by "compartment.units" in the model.list (as defined by the modelinfo files).

The 'dosing' argument includes all parameters needed to describe exposure in terms of route of administration, frequency, and quantity short of scenarios that require use of a more precise forcing function. If the dosing argument's namesake entries are left NULL, solve_model defaults to a single-time dose of 1 mg/kg BW according to the given dosing route and associated type (either add/multiply, for example we typically add a dose to gut lumen when oral route is specified).

AUC is the area under the curve of the plasma concentration.

Model parameters are named according to the following convention:

prefix	suffix	Meaning	units
K		Partition coefficient for tissue to free plasma \ tab unitless	
V		Volume	L
Q		Flow	L/h
k		Rate	1/h
	c	Parameter is proportional to body weight	1 / kg for volumes and 1/kg^(3/4) for flows

When species is specified but chemical-specific in vitro data are not available, the function uses the appropriate physiological data (volumes and flows) but default.to.human = TRUE must be used to substitute human fraction unbound, partition coefficients, and intrinsic hepatic clearance.

### Value

A matrix of class deSolve with a column for time(in days), each compartment, the area under the curve, and plasma concentration and a row for each time point.

### Author(s)

John Wambaugh, Robert Pearce, Miyuki Breen, Mark Sfeir, and Sarah E. Davidson

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### References

Pearce, Robert G., et al. "Httk: R package for high-throughput toxicokinetics." Journal of statistical software 79.4 (2017): 1.

solve_pbtk

Solve_PBTK

# **Description**

This function solves for the amounts or concentrations in uM of a chemical in different tissues as functions of time based on the dose and dosing frequency.

# Usage

```
solve_pbtk(
 chem.name = NULL,
 chem.cas = NULL,
 dtxsid = NULL,
  times = NULL,
 parameters = NULL,
 days = 10,
  tsteps = 4,
 daily.dose = NULL,
 dose = NULL,
 doses.per.day = NULL,
 initial.values = NULL,
 plots = FALSE,
 suppress.messages = FALSE,
  species = "Human",
  iv.dose = FALSE,
  input.units = "mg/kg",
 output.units = NULL,
 method = "lsoda",
 rtol = 1e-08,
 atol = 1e-12,
 default.to.human = FALSE,
 recalc.blood2plasma = FALSE,
  recalc.clearance = FALSE,
 dosing.matrix = NULL,
 adjusted.Funbound.plasma = TRUE,
 regression = TRUE,
 restrictive.clearance = TRUE,
 minimum.Funbound.plasma = 1e-04,
 monitor.vars = NULL,
)
```

# **Arguments**

chem.name

Either the chemical name, CAS number, or the parameters must be specified.

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chem. cas Either the chemical name, CAS number, or the parameters must be specified.

dtxsid EPA's DSSTox Structure ID (https://comptox.epa.gov/dashboard) the chem-

ical must be identified by either CAS, name, or DTXSIDs

times Optional time sequence for specified number of days. Dosing sequence begins

at the beginning of times.

parameters Chemical parameters from parameterize_pbtk function, overrides chem.name

and chem.cas.

days Length of the simulation.

tsteps The number of time steps per hour.
daily.dose Total daily dose, defaults to mg/kg BW.

dose Amount of a single dose, defaults to mg/kg BW.

doses.per.day Number of doses per day.

initial.values Vector containing the initial concentrations or amounts of the chemical in spec-

ified tissues with units corresponding to output.units. Defaults are zero.

plots Plots all outputs if true.

suppress.messages

Whether or not the output message is suppressed.

species Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human").

iv. dose Simulates a single i.v. dose if true.

input.units Input units of interest assigned to dosing, defaults to mg/kg BW

output.units A named vector of output units expected for the model results. Default, NULL,

returns model results in units specified in the 'modelinfo' file. See table below

for details.

method Method used by integrator (deSolve).

rtol Argument passed to integrator (deSolve).

atol Argument passed to integrator (deSolve).

default.to.human

Substitutes missing animal values with human values if true (hepatic intrinsic

clearance or fraction of unbound plasma).

recalc.blood2plasma

Recalculates the ratio of the amount of chemical in the blood to plasma using the

input parameters, calculated with hematocrit, Funbound.plasma, and Krbc2pu.

recalc.clearance

Recalculates the the hepatic clearance (Clmetabolism) with new million.cells.per.gliver

parameter.

dosing.matrix Vector of dosing times or a matrix consisting of two columns or rows named

"dose" and "time" containing the time and amount, in mg/kg BW, of each dose.

adjusted.Funbound.plasma

Uses adjusted Funbound.plasma when set to TRUE along with partition coeffi-

cients calculated with this value.

regression Whether or not to use the regressions in calculating partition coefficients.

restrictive.clearance

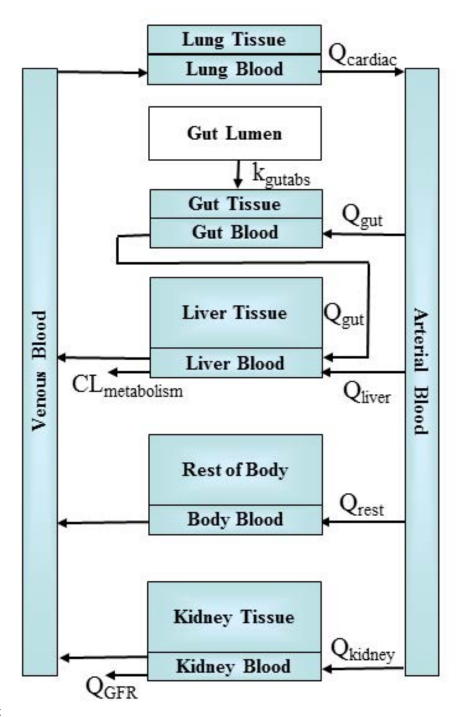
Protein binding not taken into account (set to 1) in liver clearance if FALSE.

minimum.Funbound.plasma

Monte Carlo draws less than this value are set equal to this value (default is 0.0001 – half the lowest measured Fup in our dataset).

Which variables are returned as a function of time. The default value of NULL monitor.vars provides "Cgut", "Cliver", "Cven", "Clung", "Cart", "Crest", "Ckidney", "Cplasma", "Atubules", "Ametabolized", and "AUC" Additional arguments passed to the integrator. . . . **Details** Note that the model parameters have units of hours while the model output is in days. Default NULL value for doses.per.day solves for a single dose. The compartments used in this model are the gutlumen, gut, liver, kidneys, veins, arteries, lungs, and the rest of the body. The extra compartments include the amounts or concentrations metabolized by the liver and excreted by the kidneys through the tubules. AUC is the area under the curve of the plasma concentration. Model Figure

solve_pbtk 197



altalt

When species is specified as rabbit, dog, or mouse, the function uses the appropriate physiological data(volumes and flows) but substitues human fraction unbound, partition coefficients, and intrinsic hepatic clearance.

# Value

A matrix of class deSolve with a column for time(in days), each compartment, the area under the curve, and plasma concentration and a row for each time point.

198 solve_pbtk

### Author(s)

John Wambaugh and Robert Pearce

#### References

Pearce, Robert G., et al. "Httk: R package for high-throughput toxicokinetics." Journal of statistical software 79.4 (2017): 1.

### **Examples**

```
# Multiple doses per day:
head(solve_pbtk(
  chem.name='Bisphenol-A',
  daily.dose=.5,
  days=5,
  doses.per.day=2,
  tsteps=2))
# Starting with an initial concentration:
out <- solve_pbtk(</pre>
  chem.name='bisphenola',
  dose=0,
  output.units="mg/L",
  initial.values=c(Agut=200))
# Working with parameters (rather than having solve_pbtk retrieve them):
params <- parameterize_pbtk(chem.cas="80-05-7")</pre>
head(solve_pbtk(parameters=params))
# We can change the parameters given to us by parameterize_pbtk:
params <- parameterize_pbtk(dtxsid="DTXSID4020406", species = "rat")</pre>
params["Funbound.plasma"] <- 0.1</pre>
out <- solve_pbtk(parameters=params)</pre>
# A fifty day simulation:
out <- solve_pbtk(</pre>
 chem.name = "Bisphenol A",
  days = 50,
  daily.dose=1,
  doses.per.day = 3)
plot.data <- as.data.frame(out)</pre>
css <- calc_analytic_css(chem.name = "Bisphenol A")</pre>
library("ggplot2")
c.vs.t <- ggplot(plot.data, aes(time, Cplasma)) +</pre>
  geom_line() +
  geom_hline(yintercept = css) +
  ylab("Plasma Concentration (uM)") +
  xlab("Day") +
  theme(
    axis.text = element_text(size = 16),
    axis.title = element_text(size = 16),
    plot.title = element_text(size = 17)) +
  ggtitle("Bisphenol A")
```

spleen_mass_children 199

```
print(c.vs.t)
```

spleen_mass_children Predict spleen mass for children

# **Description**

For individuals under 18, predict the spleen mass from height, weight, and gender, using equations from Ogiu et al. (1997)

# Usage

```
spleen_mass_children(height, weight, gender)
```

# **Arguments**

height Vector of heights in cm.

weight Vector of weights in kg.

gender Vector of genders (either 'Male' or 'Female').

# Value

A vector of spleen masses in kg.

# Author(s)

Caroline Ring

### References

Ogiu, Nobuko, et al. "A statistical analysis of the internal organ weights of normal Japanese people." Health physics 72.3 (1997): 368-383.

Price, Paul S., et al. "Modeling interindividual variation in physiological factors used in PBPK models of humans." Critical reviews in toxicology 33.5 (2003): 469-503.

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

200 spline_hematocrit

spline_heightweight	Smoothing splines for log height vs. age and log body weight vs. age, along with 2-D KDE residuals, by race and gender.
---------------------	-------------------------------------------------------------------------------------------------------------------------

# Description

#'Smoothing splines and KDE fits to joint distribution of height and weight residuals pre-calculated from NHANES height, weight, and age data by race/ethnicity and gender.

### Usage

spline_heightweight

#### **Format**

A data.table with 6 variables:

- g Gender: Male or Female
- r Race/ethnicity: Mexican American, Other Hispanic, Non-Hispanic White, Non-Hispanic Black, Other
- height_spline A list of smooth.spline objects, each giving a smoothed relationship between log height in cm and age in months
- weight_spline A list of smooth.spline objects, each giving a smoothed relationship between log body weight in kg and age in months
- hw_kde A list of kde objects; each is a 2-D KDE of the distribution of log height and log body weight residuals about the smoothing splines.

### Author(s)

Caroline Ring

### References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

spline_hematocrit	Smoothing splines for log hematocrit vs.	age in months, and KDE
	residuals, by race and gender.	

# Description

Smoothing splines and KDE residuals pre-calculated from NHANES hematocrit and age data by race/ethnicity and gender.

### Usage

spline_hematocrit

spline_serumcreat 201

#### **Format**

A data.table with 6 variables:

gender Gender: Male or Female

reth Race/ethnicity: Mexican American, Other Hispanic, Non-Hispanic White, Non-Hispanic Black, Other

hct_spline A list of smooth.spline objects, each giving a smoothed relationship between log hematocrit and age in months

hct_kde A list of kde objects; each is a KDE of the distribution of residuals about the smoothing spline.

### Author(s)

Caroline Ring

#### References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

spline_serumcreat Smoo

Smoothing splines for log serum creatinine vs. age in months, along with KDE residuals, by race and gender.

# **Description**

#'Smoothing splines and KDE residuals pre-calculated from NHANES serum creatinine and age data by race/ethnicity and gender.

# Usage

spline_serumcreat

### **Format**

A data.table with 6 variables:

gender Gender: Male or Female

reth Race/ethnicity: Mexican American, Other Hispanic, Non-Hispanic White, Non-Hispanic Black, Other

sc_spline A list of smooth.spline objects, each giving a smoothed relationship between log serum creatinine and age in months

sc_kde A list of kde objects; each is a KDE of the distribution of residuals about the smoothing spline.

### Author(s)

Caroline Ring

### References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

supptab1_Linakis2020 Supplementary output from Linakis 2020 vignette analysis.

# Description

Supplementary output from Linakis 2020 vignette analysis.

### Usage

supptab1_Linakis2020

#### **Format**

A data.frame containing x rows and y columns.

# Author(s)

Matt Linakis

### **Source**

Matt Linakis

### References

DSStox database (https://www.epa.gov/ncct/dsstox

supptab2_Linakis2020 More supplementary output from Linakis 2020 vignette analysis.

# Description

More supplementary output from Linakis 2020 vignette analysis.

# Usage

supptab2_Linakis2020

### **Format**

A data frame containing x rows and y columns.

### Author(s)

Matt Linakis

Tables.Rdata.stamp 203

#### Source

Matt Linakis

#### References

DSStox database (https://www.epa.gov/ncct/dsstox

Tables.Rdata.stamp

A timestamp of table creation

# **Description**

The Tables.RData file is separately created as part of building a new release of HTTK. This time stamp indicates the script used to build the file and when it was run.

# Usage

Tables.Rdata.stamp

### **Format**

An object of class character of length 1.

### Author(s)

John Wambaugh

tissue.data

Tissue composition and species-specific physiology parameters

# Description

This data set contains values from Schmitt (2008) and Ruark et al. (2014) describing the composition of specific tissues and from Birnbaum et al. (1994) describing volumes of and blood flows to those tissues, allowing parameterization of toxicokinetic models for human, mouse, rat, dog, or rabbit. Tissue volumes were calculated by converting the fractional mass of each tissue with its density (both from ICRP), lumping the remaining tissues into the rest-of-body, excluding the mass of the gastrointestinal contents

# Usage

tissue.data

### **Format**

A data.frame containing 13 rows and 20 columns.

### Author(s)

John Wambaugh, Robert Pearce, and Nisha Sipes

204 tissue_masses_flows

#### Source

Pearce et al. (2017), in preparation,

Wambaugh, John F., et al. "Toxicokinetic triage for environmental chemicals." Toxicological Sciences (2015): 228-237.

#### References

Birnbaum, L and Brown, R and Bischoff, K and Foran, J and Blancato, J and Clewell, H and Dedrick, R (1994). Physiological parameter values for PBPK model. International Life Sciences Institute, Risk Science Institute, Washington, DC

Ruark, Christopher D., et al. "Predicting passive and active tissue: plasma partition coefficients: Interindividual and interspecies variability." Journal of pharmaceutical sciences 103.7 (2014): 2189-2198.

Schmitt, W. (2008). General approach for the calculation of tissue to plasma partition coefficients. Toxicology in vitro: an international journal published in association with BIBRA 22(2), 457-67, 10.1016/j.tiv.2007.09.010.

ICRP. Report of the Task Group on Reference Man. ICRP Publication 23 1975

tissue_masses_flows

Given a data.table describing a virtual population by the NHANES quantities, generates HTTK physiological parameters for each individual.

### **Description**

Given a data.table describing a virtual population by the NHANES quantities, generates HTTK physiological parameters for each individual.

# Usage

```
tissue_masses_flows(tmf_dt)
```

### **Arguments**

tmf_dt

A data.table generated by gen_age_height_weight(), containing variables gender, reth, age_months, age_years, weight, and height.

### Value

The same data.table, with aditional variables describing tissue masses and flows.

# Author(s)

Caroline Ring

tissue_scale 205

#### References

Barter, Zoe E., et al. "Scaling factors for the extrapolation of in vivo metabolic drug clearance from in vitro data: reaching a consensus on values of human micro-somal protein and hepatocellularity per gram of liver." Current Drug Metabolism 8.1 (2007): 33-45.

Birnbaum, L., et al. "Physiological parameter values for PBPK models." International Life Sciences Institute, Risk Science Institute, Washington, DC (1994).

Geigy Pharmaceuticals, "Scientific Tables", 7th Edition, John Wiley and Sons (1970)

McNally, Kevin, et al. "PopGen: a virtual human population generator." Toxicology 315 (2014): 70-85.

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

tissue_scale

Allometric scaling.

# **Description**

Allometrically scale a tissue mass or flow based on height^3/4.

### Usage

```
tissue_scale(height_ref, height_indiv, tissue_mean_ref)
```

### **Arguments**

```
height_ref Reference height in cm.
height_indiv Individual height in cm.
tissue_mean_ref
```

Reference tissue mass or flow.

# Value

Allometrically scaled tissue mass or flow, in the same units as tissue_mean_ref.

### Author(s)

Caroline Ring

# References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

206 wambaugh2019

wambaugh2019

in vitro Toxicokinetic Data from Wambaugh et al. (2019)

### **Description**

These data are the new HTTK in vitro data for chemicals reported in Wambaugh et al. (2019) They are the processed values used to make the figures in that manuscript. These data summarize the results of Bayesian analysis of the in vitro toxicokinetic experiments conducted by Cyprotex to characterize fraction unbound in the presence of pooled human plasma protein and the intrnsic hepatic clearance of the chemical by pooled human hepatocytes.

### Usage

wambaugh2019

#### **Format**

A data frame with 496 rows and 17 variables:

**Compound** The name of the chemical

**CAS** The Chemical Abstracts Service Registry Number

**Human.Clint** Median of Bayesian credible interval for intrinsic hepatic clearance (uL/min/million hepatocytes)]

Human.Clint.pValue Probability that there is no clearance

**Human.Funbound.plasma** Median of Bayesian credibl interval for fraction of chemical free in the presence of plasma

**pKa_Accept** pH(s) at which hydrogen acceptor sites (if any) are at equilibrium

pKa_Donor pH(s) at which hydrogne donor sites (if any) are at equilibrium

DSSTox_Substance_Id Identifier for CompTox Chemical Dashboard

SMILES Simplified Molecular-Input Line-Entry System structure description

**Human.Clint.Low95** Lower 95th percentile of Bayesian credible interval for intrinsic hepatic clearance (uL/min/million hepatocytes)

**Human.Clint.High95** Uppper 95th percentile of Bayesian credible interval for intrinsic hepatic clearance (uL/min/million hepatocytes)

Human.Clint.Point Point estimate of intrinsic hepatic clearance (uL/min/million hepatocytes)

**Human.Funbound.plasma.Low95** Lower 95th percentile of Bayesian credible interval for fraction of chemical free in the presence of plasma

**Human.Funbound.plasma.High95** Upper 95th percentile of Bayesian credible interval for fraction of chemical free in the presence of plasma

**Human.Funbound.plasma.Point** Point estimate of the fraction of chemical free in the presence of plasma

MW Molecular weight (Daltons)

logP log base ten of octanol:water partiion coefficient

### Author(s)

John Wambaugh

wambaugh2019.nhanes 207

#### **Source**

Wambaugh et al. (2019)

#### References

Wambaugh et al. (2019) "Assessing Toxicokinetic Uncertainty and Variability in Risk Prioritization", Toxicological Sciences, 172(2), 235-251.

wambaugh 2019.nhanes

NHANES Chemical Intake Rates for chemicals in Wambaugh et al. (2019)

# **Description**

These data are a subset of the Bayesian inferrences reported by Ring et al. (2017) from the U.S. Centers for Disease Control and Prevention (CDC) National Health and Nutrition Examination Survey (NHANES). They reflect the populaton median intake rate (mg/kg body weight/day), with uncertainty.

### Usage

wambaugh2019.nhanes

### **Format**

A data frame with 20 rows and 4 variables:

**IP** The median of the Bayesian credible interval for median population intake rate (mg/kg bodyweight/day)

**IP.min** The lower 95th percentile of the Bayesian credible interval for median population intake rate (mg/kg bodyweight/day)

**IP.max** The upper 95th percentile of the Bayesian credible interval for median population intake rate (mg/kg bodyweight/day)

**CASRN** The Chemical Abstracts Service Registry Number

### Author(s)

John Wambaugh

### Source

Wambaugh et al. (2019)

# References

Ring, Caroline L., et al. "Identifying populations sensitive to evironmental chemicals by simulating toxicokinetic variability." Environment international 106 (2017): 105-118

Wambaugh et al. (2019) "Assessing Toxicokinetic Uncertainty and Variability in Risk Prioritization", Toxicological Sciences, 172(2), 235-251.

208 wambaugh2019.raw

wambaugh2019.raw Raw Bayesian in vitro Toxicokinetic Data Analysis from Wambaugh et al. (2019)

### **Description**

These data are the new HTTK in vitro data for chemicals reported in Wambaugh et al. (2019) They are the output of different Bayesian models evaluated to compare using a single protein concentration vs. the new three concentration titration protocol. These data summarize the results of Bayesian analysis of the in vitro toxicokinetic experiments conducted by Cyprotex to characterize fraction unbound in the presence of pooled human plasma protein and the intrnsic hepatic clearance of the chemical by pooled human hepatocytes. This file includes replicates (different Compound-Name id's but same chemical')

### Usage

wambaugh2019.raw

#### **Format**

A data frame with 530 rows and 28 variables:

**DTXSID** Identifier for CompTox Chemical Dashboard

Name The name of the chemical

**CAS** The Chemical Abstracts Service Registry Number

CompoundName Sample name provided by EPA to Cyprotex

Fup.point Point estimate of the fraction of chemical free in the presence of plasma

**Base.Fup.Med** Median of Bayesian credible interval for fraction of chemical free in the presence of plasma for analysis of 100 physiological plasma protein data only (base model)

**Base.Fup.Low** Lower 95th percentile of Bayesian credible interval for fraction of chemical free in the presence of plasma for analysis of 100 physiological plasma protein data only (base model)

**Base.Fup.High** Upper 95th percentile of Bayesian credible interval for fraction of chemical free in the presence of plasma for analysis of 100 physiological plasma protein data only (base model)

**Affinity.Fup.Med** Median of Bayesian credible interval for fraction of chemical free in the presence of plasma for analysis of protein titration protocol data (affinity model)

**Affinity.Fup.Low** Lower 95th percentile of Bayesian credible interval for fraction of chemical free in the presence of plasma for analysis of protein titration protocol data (affinity model)

**Affinity.Fup.High** Upper 95th percentile of Bayesian credible interval for fraction of chemical free in the presence of plasma for analysis of protein titration protocol data (affinity model)

**Affinity.Kd.Med** Median of Bayesian credible interval for protein binding affinity from analysis of protein titration protocol data (affinity model)

**Affinity.Kd.Low** Lower 95th percentile of Bayesian credible interval for protein binding affinity from analysis of protein titration protocol data (affinity model)

**Affinity.Kd.High** Upper 95th percentile of Bayesian credible interval for protein binding affinity from analysis of protein titration protocol data (affinity model)

wambaugh2019.raw 209

**Decreases.Prob** Probability that the chemical concentration decreased systematically during hepatic clearance assay.

- **Saturates.Prob** Probability that the rate of chemical concentration decrease varied between the 1 and 10 uM hepatic clearance experiments.
- **Slope.1uM.Median** Estimated slope for chemcial concentration decrease in the 1 uM hepatic clearance assay.
- **Slope.10uM.Median** Estimated slope for chemcial concentration decrease in the 10 uM hepatic clearance assay.
- **CLint.1uM.Median** Median of Bayesian credible interval for intrinsic hepatic clearance at 1 uM initial chemical concentration (uL/min/million hepatocytes)]
- **CLint.1uM.Low95th** Lower 95th percentile of Bayesian credible interval for intrinsic hepatic clearance at 1 uM initial chemical concentration (uL/min/million hepatocytes)
- **CLint.1uM.High95th** Uppper 95th percentile of Bayesian credible interval for intrinsic hepatic clearance at 1 uM initial chemical concentration(uL/min/million hepatocytes)
- **CLint.10uM.Median** Median of Bayesian credible interval for intrinsic hepatic clearance at 10 uM initial chemical concentration (uL/min/million hepatocytes)]
- **CLint.10uM.Low95th** Lower 95th percentile of Bayesian credible interval for intrinsic hepatic clearance at 10 uM initial chemical concentration (uL/min/million hepatocytes)
- **CLint.10uM.High95th** Uppper 95th percentile of Bayesian credible interval for intrinsic hepatic clearance at 10 uM initial chemical concentration(uL/min/million hepatocytes)
- **CLint.1uM.Point** Point estimate of intrinsic hepatic clearance (uL/min/million hepatocytes) for 1 uM initial chemical concentration
- **CLint.10uM.Point** Point estimate of intrinsic hepatic clearance (uL/min/million hepatocytes) for 10 uM initial chemical concentration
- Fit Classification of clearance observed
- SMILES Simplified Molecular-Input Line-Entry System structure description

# Author(s)

John Wambaugh

### **Source**

Wambaugh et al. (2019)

### References

Wambaugh et al. (2019) "Assessing Toxicokinetic Uncertainty and Variability in Risk Prioritization", Toxicological Sciences, 172(2), 235-251.

210 wambaugh2019.tox21

wambaugh2019.seem3

ExpoCast SEEM3 Consensus Exposure Model Predictions for Chemical Intake Rates

### **Description**

These data are a subset of the Bayesian inferrences reported by Ring et al. (2019) for a consensus model of twelve exposue predictors. The predictors were calibrated based upon their ability to predict intake rates inferred National Health and Nutrition Examination Survey (NHANES). They reflect the populaton median intake rate (mg/kg body weight/day), with uncertainty.

### Usage

wambaugh2019.seem3

### **Format**

A data frame with 385 rows and 38 variables:

#### Author(s)

John Wambaugh

#### **Source**

Wambaugh et al. (2019)

### References

Ring, Caroline L., et al. "Consensus modeling of median chemical intake for the US population based on predictions of exposure pathways." Environmental science & technology 53.2 (2018): 719-732.

Wambaugh et al. (2019) "Assessing Toxicokinetic Uncertainty and Variability in Risk Prioritization", Toxicological Sciences, 172(2), 235-251.

wambaugh2019.tox21

Tox21 2015 Active Hit Calls (EPA)

# **Description**

The ToxCast and Tox21 research programs employ batteries of high-throughput assays to assess chemical bioactivity in vitro. Not every chemical is tested through every assay. Most assays are conducted in concentration response, and each corresponding assay endpoint is analyzed statistically to determine if there is a concentration-dependent response or "hit" using the ToxCast Pipeline. Most assay endpoint-chemical combinations are non-responsive. Here, only the hits are treated as potential indicators of bioactivity. This bioactivity does not have a direct toxicological interpretation. The October 2015 release (invitrodb_v2) of the ToxCast and Tox21 data were used for this analysis. This object contains just the chemicals in Wambaugh et al. (2019) and only the quantiles across all assays for the ACC.

wang2018 211

### **Usage**

wambaugh2019.tox21

#### **Format**

A data.table with 401 rows and 6 columns

### Author(s)

John Wambaugh

#### **Source**

ftp://newftp.epa.gov/COMPTOX/High_Throughput_Screening_Data/Previous_Data/ToxCast_
Data_Release_Oct_2015/

#### References

Kavlock, Robert, et al. "Update on EPA's ToxCast program: providing high-throughput decision support tools for chemical risk management." Chemical research in toxicology 25.7 (2012): 1287-1302.

Tice, Raymond R., et al. "Improving the human hazard characterization of chemicals: a Tox21 update." Environmental health perspectives 121.7 (2013): 756-765.

Richard, Ann M., et al. "ToxCast chemical landscape: paving the road to 21st century toxicology." Chemical research in toxicology 29.8 (2016): 1225-1251.

Filer, Dayne L., et al. "tcpl: the ToxCast pipeline for high-throughput screening data." Bioinformatics 33.4 (2016): 618-620.

Wambaugh, John F., et al. "Assessing Toxicokinetic Uncertainty and Variability in Risk Prioritization." Toxicological Sciences 172.2 (2019): 235-251.

wang2018

Wang et al. 2018 Wang et al. (2018) screened the blood of 75 pregnant women for the presence of environmental organic acids (EOAs) and identified mass spectral features corresponding to 453 chemical formulae of which 48 could be mapped to likely structures. Of the 48 with tentative structures the identity of six were confirmed with available chemical standards.

### **Description**

Wang et al. 2018 Wang et al. (2018) screened the blood of 75 pregnant women for the presence of environmental organic acids (EOAs) and identified mass spectral features corresponding to 453 chemical formulae of which 48 could be mapped to likely structures. Of the 48 with tentative structures the identity of six were confirmed with available chemical standards.

### Usage

wang2018

212 well_param

#### **Format**

data.frame

#### Source

Kapraun et al. 2021 (submitted)

### References

Wang A, Gerona RR, Schwartz JM, Lin T, Sirota M, Morello-Frosch R, Woodruff TJ (2018). "A Suspect Screening Method for Characterizing Multiple Chemical Exposures among a Demographically Diverse Population of Pregnant Women in San Francisco." *Environmental Health Perspectives*, **126**(7), 077009. doi: 10.1289/EHP2920, https://ehp.niehs.nih.gov/doi/abs/10.1289/EHP2920.

well_param

Microtiter Plate Well Descriptions for Armitage et al. (2014) Model

### **Description**

Microtiter Plate Well Descriptions for Armitage et al. (2014) model from Honda et al. (2019)

# Usage

well_param

### **Format**

A data frame / data table with 11 rows and 8 variables:

sysID Identifier for each multi-well plate system

well_desc Well description

well_number Number of wells on plate

**area_bottom** Area of well bottom in mm^2

cell_yield Number of cells

diam Diameter of well in mm

v total Total volume of well in uL)

v_working Working volume of well in uL

### Author(s)

Greg Honda

### **Source**

https://www.corning.com/catalog/cls/documents/application-notes/CLS-AN-209.pdf

# References

Armitage, J. M.; Wania, F.; Arnot, J. A. Environ. Sci. Technol. 2014, 48, 9770-9779. dx.doi.org/10.1021/es501955g Honda, Gregory S., et al. "Using the Concordance of In Vitro and In Vivo Data to Evaluate Extrapolation Assumptions", PloS ONE 14.5 (2019): e0217564.

Wetmore.data 213

Wetmore data

Published toxicokinetic predictions based on in vitro data

### **Description**

This data set gives the chemical specific predictions for serum concentration at steady state resulting from constant infusion exposure, as published in a series of papers from Barbara Wetmore's group at the Hamner Institutes for Life Sciences. Predictions include the median and 90% interval in uM and mg/L. Calculations were made using the 1 and 10 uM in vitro measured clearances.

### Usage

Wetmore.data

#### **Format**

A data frame containing 577 rows and 20 columns.

### **Source**

Wambaugh, John F., et al. "Toxicokinetic triage for environmental chemicals." Toxicological Sciences (2015): 228-237.

#### References

Wetmore, B.A., Wambaugh, J.F., Ferguson, S.S., Sochaski, M.A., Rotroff, D.M., Freeman, K., Clewell, H.J., Dix, D.H., Andersen, M.E., Houck, K.A., Allen, B., Judson, R.S., Sing, R., Kavlock, R.J., Richard, A.M., and Thomas, R.S., "Integration of Dosimetry, Exposure and High-Throughput Screening Data in Chemical Toxicity Assessment," Toxicological Sciences 125 157-174 (2012)

Wetmore, B.A., Wambaugh, J.F., Ferguson, S.S., Li, L., Clewell, H.J. III, Judson, R.S., Freeman, K., Bao, W, Sochaski, M.A., Chu T.-M., Black, M.B., Healy, E, Allen, B., Andersen M.E., Wolfinger, R.D., and Thomas R.S., "The Relative Impact of Incorporating Pharmacokinetics on Predicting in vivo Hazard and Mode-of-Action from High-Throughput in vitro Toxicity Assays" Toxicological Sciences, 132:327-346 (2013).

Wetmore, B. A., Wambaugh, J. F., Allen, B., Ferguson, S. S., Sochaski, M. A., Setzer, R. W., Houck, K. A., Strope, C. L., Cantwell, K., Judson, R. S., LeCluyse, E., Clewell, H.J. III, Thomas, R.S., and Andersen, M. E. (2015). "Incorporating High-Throughput Exposure Predictions with Dosimetry-Adjusted In Vitro Bioactivity to Inform Chemical Toxicity Testing" Toxicological Sciences, kfv171.

Wetmore2012

Published toxicokinetic predictions based on in vitro data from Wetmore et al. 2012.

# **Description**

This data set overlaps with Wetmore.data and is used only in Vignette 4 for steady state concentration. 214 wfl

### Usage

Wetmore2012

#### **Format**

A data.frame containing 13 rows and 15 columns.

#### References

Wetmore, B.A., Wambaugh, J.F., Ferguson, S.S., Sochaski, M.A., Rotroff, D.M., Freeman, K., Clewell, H.J., Dix, D.H., Andersen, M.E., Houck, K.A., Allen, B., Judson, R.S., Sing, R., Kavlock, R.J., Richard, A.M., and Thomas, R.S., "Integration of Dosimetry, Exposure and High-Throughput Screening Data in Chemical Toxicity Assessment," Toxicological Sciences 125 157-174 (2012)

wf1

WHO weight-for-length charts

### **Description**

Charts giving weight-for-length percentiles for boys and girls under age 2.

### Usage

wf1

### **Format**

A data.table object with variables

Sex 'Male' or 'Female'

Length length in cm

L, M, S LMS parameters; see https://www.cdc.gov/growthcharts/percentile_data_files.

P2.3, P5, P10, P25, P50, P75, P90, P95, and P97.7 weight percentiles

### **Details**

For infants under age 2, weight class depends on weight for length percentile. #'

**Underweight** <2.3rd percentile

Normal weight 2.3rd-97.7th percentile

**Obese** >=97.7th percentile

# Author(s)

Caroline Ring

### **Source**

https://www.cdc.gov/growthcharts/who/girls_weight_head_circumference.htm and https://www.cdc.gov/growthcharts/who/boys_weight_head_circumference.htm

# References

Ring, Caroline L., et al. "Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability." Environment International 106 (2017): 105-118

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