Package 'httk'

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Title High-Throughput Toxicokinetics

Description Functions and data tables for simulation and statistical analysis of chemical toxicokinetics (``TK") as in 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 (e.g., one compartment) ``TK" models can be parameterized for several hundred chemicals and multiple species. These models are solved efficiently, often using compiled (C-based) code. A Monte Carlo sampler is included for simulating biological variability (Ring et al., 2017 <doi:10.1016/j.envint.2017.06.004>) and measurement limitations. 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 (e.g., 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)

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Suggests ggplot2, knitr, rmarkdown, R.rsp, GGally, gplots, scales, EnvStats, MASS, RColorBrewer, TeachingDemos, classInt, ks, stringr, reshape, reshape2, gdata, viridis, CensRegMod, gmodels, colorspace

License GPL-3
LazyData true

VignetteBuilder knitr, R.rsp

RoxygenNote 6.1.1

URL https:

//www.epa.gov/chemical-research/rapid-chemical-exposure-and-dose-research

BugReports https://github.com/USEPA/CompTox-ExpoCast-httk

2 R topics documented:

NeedsCompilation yes Author John Wambaugh [aut, cre], Robert Pearce [aut], Caroline Ring [aut], Greg Honda [aut], Mark Sfeir [aut], Jimena Davis [ctb], James Sluka [ctb], Nisha Sipes [ctb], Barbara Wetmore [ctb], Woodrow Setzer [ctb]

R topics documented:

httk-package	4
add_chemtable	5
age_dist_smooth	6
age_draw_smooth	7
armitage_estimate_sarea	8
armitage_eval	8
$\mathcal{E} = 1$	10
available_rblood2plasma	11
blood_mass_correct	12
blood_weight	12
bmiage	13
	14
bone_mass_age	14
	15
	15
calc_analytic_css_1comp	17
calc_analytic_css_3comp	18
calc_analytic_css_3compss	19
calc_analytic_css_pbtk	20
calc_css	21
calc_elimination_rate	23
calc_hepatic_clearance	25
calc_ionization	26
calc_mc_css	27
calc_mc_oral_equiv	31
calc_rblood2plasma	34
calc_stats	36
calc_total_clearance	37
calc_vdist	38
chem.invivo.PK.aggregate.data	40
chem.invivo.PK.data	40
	43
	46
	17
ckd_epi_eq	49
-1-1	50

draw_fup_clint	
estimate_gfr	
estimate_gfr_ped	
estimate_hematocrit	
export_pbtk_jarnac	
export_pbtk_sbml	
gen_age_height_weight	
gen_height_weight	
get_cheminfo	
get_chem_id	9
get_gfr_category	0
get_httk_params	
get_lit_cheminfo	
get_lit_css	3
get_lit_oral_equiv	5
get_physchem_param	6
get_rblood2plasma	7
get_weight_class	
hematocrit_infants	
honda.ivive	
howgate	
httkpop	
httkpop_bio	
httkpop_direct_resample	
httkpop_direct_resample_inner	
httkpop_generate	
httkpop_virtual_indiv	
in.list	
is.httk	
is_in_inclusive	
johnson	
kidney_mass_children	
liver_mass_children	
load_sipes2017	
lump_tissues	
lung_mass_children	
mcnally_dt	
monte_carlo	
-	
Obach2008	
onlyp	
pancreas_mass_children	
parameterize_1comp	
parameterize_3comp	
parameterize_pbtk	
parameterize_schmitt	
parameterize_steadystate	
pc.data	
pharma	
physiology.data	
predict_partitioning_schmitt	
rfun 10	13

4 httk-package

r_left_censored_norm
sipes2017
sipes2017.table
skeletal_muscle_mass
skeletal_muscle_mass_children
skin_mass_bosgra
solve_1comp
solve_3comp
solve_pbtk
spleen_mass_children
spline_heightweight
spline_hematocrit
spline_serumcreat
Tables.Rdata.stamp
tissue.data
tissue_masses_flows
tissue_scale
ToxCast2015subset
wambaugh2019
wambaugh2019.nhanes
wambaugh2019.raw
wambaugh2019.seem3
well_param
Wetmore.data
Wetmore2012
wfl

132

httk-package

High-Throughput Toxicokinetics

Description

Index

Functions and data tables for simulation and statistical analysis of chemical toxicokinetics ("TK") as in 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 (e.g., one compartment) "TK" models can be parameterized for several hundred chemicals and multiple species. These models are solved efficiently, often using compiled (C-based) code. A Monte Carlo sampler is included for simulating biological variability (Ring et al., 2017 <doi:10.1016/j.envint.2017.06.004>) and measurement limitations. 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 (e.g., Tox21, Tox-Cast) 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

add_chemtable 5

See Also

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

Pearce et al. (2017): httk: R Package for High-Throughput Toxicokinetics

Wetmore et al. (2015): Incorporating High-Throughput Exposure Predictions With Dosimetry-Adjusted In Vitro Bioactivity to Inform Chemical Toxicity Testing

Wambaugh et al. (2015): Toxicokinetic Triage for Environmental Chemicals

Pearce et al. (2017): Evaluation and calibration of high-throughput predictions of chemical distribution to tissues

Ring et al. (2017): Identifying populations sensitive to environmental chemicals by simulating toxicokinetic variability

Sipes et al. (2017): An Intuitive Approach for Predicting Potential Human Health Risk with the Tox21 10k Library

Wambaugh et al. (2018): Evaluating In Vitro-In Vivo Extrapolation of Toxicokinetics

Honda et al. (2019): Using the concordance of in vitro and in vivo data to evaluate extrapolation assumptionss

EPA's 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 = F)
```

Arguments

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

6 age_dist_smooth

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.

Value

data.frame

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

Author(s)

John Wambaugh

Examples

```
my.new.data <- as.data.frame(c("A","B","C"),stringsAsFactors=FALSE)</pre>
my.new.data <- cbind(my.new.data,as.data.frame(c("111-11-2","222-22-0","333-33-5"),</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 \leftarrow 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","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",
                                     MW="MW",
                                     logP="LogP",
                                     Funbound.plasma="Fup",
                                     Clint="CLint"),
                                     species="Human",
                                     reference="MyPaper 2015")
parameterize_steadystate(chem.name="C")
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
```

age_draw_smooth 7

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

References

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

Description

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

Usage

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

Arguments

g Gender. Either 'Male' or 'Female'.

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

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

nsamp Number of ages to draw.

 ${\tt agelim_months} \quad {\tt Two-element\ numeric\ vector\ giving\ the\ minimum\ and\ maximum\ ages\ in\ months}$

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

8 armitage_eval

```
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 == T (default) give nonzero surface area (sarea, m^2) option.bottom == T (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

tcdata

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)==T

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 on well number (m^3)

Value

tcdata, A data table with well_number, sarea (surface area, m^2), cell_yield (# cells), v_working (m^3), v_total (m^3) per well

Author(s)

Greg Honda

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.

armitage_eval 9

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 = F,
    this.option.swat2 = F, 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)
```

Arguments

```
For vector or single value, CAS number
casrn.vector
nomconc.vector For vector or single value, micromolar nominal concentration (e.g. AC50 value)
this.well_number
                  For single value, plate format default is 384, used if is.na(tcdata)==T
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.
                  Surface area per well (m^2)
this.sarea
                  Total volume per well (m<sup>3</sup>)
this.v_total
this.v_working Working volume per well (m^3)
this.cell_yield
                  Number of cells per well
this.Tsys
                  System temperature (oC)
this.Tref
                  Reference temperature (K)
this.option.kbsa2
                  Use alternative bovine-serum-albumin partitioning model
this.option.swat2
                  Use alternative water solubility correction
this.pseudooct Pseudo-octanol cell storage lipid content
this.memblip
                  Membrane lipid content of cells
this.nlom
                  Structural protein conent of cells
this.P_nlom
                  Proportionality constant to octanol structural protein
this.P_dom
                  Proportionality constant to octnaol dom
this.P_cells
                  Proportionality constant to octanol storage lipid
this.csalt
                  Ionic strength of buffer, mol/L
this.celldensity
                  Cell density kg/L, g/mL
this.cellmass
                  Mass per cell, ng/cell
this.f_oc
                  1, everything assumed to be like proteins
```

Value

tcdata

10 armitage_input

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

```
temp <- armitage_eval(casrn.vector = c("80-05-7", "81-81-2"), this.FBSf = 0.1, this.well_number = 384, nomconc = 10) print(temp$cfree.invitro)
```

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)

Usage

```
armitage_input
```

Format

A data frame with 53940 rows and 10 variables:

MP

MW

casrn

compound_name

gkaw

gkow

gswat

Author(s)

Greg Honda

Source

```
http://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.

```
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.

Usage

```
available_rblood2plasma(chem.cas = NULL, chem.name = NULL,
   species = "Human", adjusted.Funbound.plasma = T,
   suppress.messages = F)
```

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.

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

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 ~~

Author(s)

Robert Pearce

Examples

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

12 blood_weight

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.

blood_weight

Predict blood mass.

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

bmiage 13

References

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

L, M, S LMS parameters; see https://www.cdc.gov/growthcharts/percentile_data_files.

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

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

```
http://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

14 bone_mass_age

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.

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.

brain_mass 15

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.

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.

Usage

```
calc_analytic_css(chem.name = NULL, chem.cas = NULL,
  parameters = NULL, daily.dose = 1, output.units = "uM",
  model = "pbtk", concentration = "plasma", suppress.messages = F,
  recalc.blood2plasma = F, tissue = NULL, restrictive.clearance = T,
  bioactive.free.invivo = F, IVIVE = 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.
parameters	Chemical parameters from parameterize_pbtk (for model = 'pbtk'), parameterize_3comp (for model = '3compartment), parmeterize_1comp(for model = '1compartment') or parameterize_steadystate (for model = '3compartmentss'), overrides chem.name and chem.cas.
daily.dose	Total daily dose, mg/kg BW.
output.units	Units for returned concentrations, defaults to uM (specify units = " uM ") but can also be mg/L .

16 calc_analytic_css

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.

concentration

Desired concentration type, 'blood', 'tissue', 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 metabolism (feater metabolism due to repid off hinding)

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.

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.

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

Details

Concentrations are calculated for the specified 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 concentration

Author(s)

Robert Pearce, John Wambaugh, and Greg Honda

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,
  parameters = NULL, hourly.dose = 1/24, concentration = "plasma",
  suppress.messages = F, recalc.blood2plasma = F, tissue = NULL,
  restrictive.clearance = T, bioactive.free.invivo = F, ...)
```

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.

Chemical parameters from parameterize_pbtk (for model = 'pbtk'), parameterize_3comp (for model = '3compartment), parmeterize_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 concentration in uM 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.

Usage

```
calc_analytic_css_3comp(chem.name = NULL, chem.cas = NULL,
  parameters = NULL, hourly.dose = 1/24, concentration = "plasma",
  suppress.messages = F, recalc.blood2plasma = F, tissue = NULL,
  restrictive.clearance = T, 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.

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

terize_3comp (for model = '3compartment), parmeterize_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 concentration in uM units

Author(s)

Robert Pearce and John Wambaugh

```
calc_analytic_css_3compss
```

Calculate the analytic steady state concentration for the three oompartment 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,
  parameters = NULL, hourly.dose = 1/24, concentration = "plasma",
  suppress.messages = F, recalc.blood2plasma = F, tissue = NULL,
  restrictive.clearance = T, 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.

Chemical parameters from parameterize_pbtk (for model = 'pbtk'), parameterize_3comp (for model = '3compartment), parmeterize_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 concentration in uM units

Author(s)

Robert Pearce and John Wambaugh

```
calc_analytic_css_pbtk
```

Calculate the analytic steady state concentration for model pbtk.

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,
  parameters = NULL, hourly.dose = 1/24, concentration = "plasma",
  suppress.messages = F, recalc.blood2plasma = F, tissue = NULL,
  restrictive.clearance = T, 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.

Chemical parameters from parameterize_pbtk (for model = 'pbtk'), parameterize_3comp (for model = '3compartment), parmeterize_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', 'tissue', or default 'plasma'.

calc_css 21

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 concentration in uM 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.

Usage

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

Arguments

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

22 calc_css

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.

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

suppress.messages

Whether or not to suppress messages.

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.

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

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 average concentration comes within 100 * p percent of the true

steady state concentration.

Author(s)

Robert Pearce, John Wambaugh

calc_elimination_rate 23

Examples

```
calc_css(chem.name='Bisphenol-A',doses.per.day=5,f=.001,output.units='mg/L')
## Not run:
parms <- parameterize_3comp(chem.name='Bisphenol-A')</pre>
parms$Funbound.plasma <- .07
calc_css(parms,concentration='blood',model='3compartment')
library("ggplot2")
out <- solve_pbtk(chem.name = "Bisphenol A", days = 50, doses.per.day = 3)</pre>
plot.data <- as.data.frame(out)</pre>
css <- calc_analytic_css(chem.name = "Bisphenol A")</pre>
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)
days <- NULL
avg <- NULL
max <- NULL
for(this.cas in get_cheminfo()){
css.info <- calc_css(chem.cas = this.cas, doses.per.day = 1,suppress.messages=T)</pre>
days[[this.cas]] <- css.info[["the.day"]]</pre>
avg[[this.cas]] <- css.info[["avg"]]</pre>
max[[this.cas]] <- css.info[["max"]]</pre>
days.data <- as.data.frame(days)</pre>
hist <- ggplot(days.data, aes(days)) +
geom_histogram(fill = "blue", binwidth = 1/6) + scale_x_log10() +
ylab("Number of Chemicals") + xlab("Days") + theme(axis.text =
element_text(size = 16), axis.title = element_text(size = 16))
print(hist)
avg.max.data <- as.data.frame(cbind(avg, max))</pre>
avg.vs.max <- ggplot(avg.max.data, aes(avg, max)) + geom_point() +</pre>
geom_abline() + scale_x_log10() + scale_y_log10() +
xlab("Average Concentration at Steady State (uM)") +
ylab("Max Concentration at Steady State (uM)") +
theme(axis.text = element_text(size = 16),
axis.title = element_text(size = 16))
print(avg.vs.max)
## End(Not run)
```

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.

24 calc_elimination_rate

Usage

```
calc_elimination_rate(chem.cas = NULL, chem.name = NULL,
  parameters = NULL, species = "Human", suppress.messages = F,
  default.to.human = F, restrictive.clearance = T,
  adjusted.Funbound.plasma = T, regression = T,
  well.stirred.correction = T, 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.

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.

Value

Elimination rate

Units of 1/h.

Author(s)

John Wambaugh

calc_hepatic_clearance 25

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_hepatic_clearance

Calculate the hepatic clearance.

Description

This function calculates the hepatic clearance in plasma for a well-stirred model or other type if specified.

Usage

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

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.

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.

26 calc_ionization

Value

```
Hepatic Clearance
```

Units of L/h/kg BW.

Author(s)

John Wambaugh and Robert Pearce

Examples

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

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, parameters = NULL,
pH = NULL, pKa_Donor = NA, pKa_Accept = NA)
```

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.

 $parameters \hspace{0.5cm} \textbf{Chemical parameters from a parameterize_MODEL function, overrides chem.name}$

and chem.cas.

pH where ionization is evaluated.

pKa_Donor Compound H dissociation equilibirum constant(s). Overwrites chem.name and

chem.cas.

pKa_Accept Compound H association equilibirum constant(s). Overwrites chem.name and

chem.cas.

Details

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

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

```
calc_ionization(chem.name='bisphenola',pH=7.4)
calc_ionization(pKa_Donor=8,pKa_Accept=c(1,4),pH=9)
```

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, parameters = NULL,
  daily.dose = 1, which.quantile = 0.95, species = "Human",
  output.units = "mg/L", suppress.messages = F,
  model = "3compartmentss", censored.params = list(Funbound.plasma =
  list(cv = 0.3, lod = 0.01)), vary.params = list(BW = 0.3, Vliverc =
  0.3, Qgfrc = 0.3, Qtotal.liverc = 0.3, million.cells.per.gliver = 0.3,
```

```
Clint = 0.3), fup.meas.cv = 0.4, clint.meas.cv = 0.3, fup.pop.cv = 0.3, clint.pop.cv = 0.3, samples = 1000, return.samples = F, default.to.human = F, tissue = NULL, well.stirred.correction = T, adjusted.Funbound.plasma = T, regression = T, clint.pvalue.threshold = 0.05, restrictive.clearance = T, bioactive.free.invivo = FALSE, concentration = "plasma", IVIVE = NULL, httkpop = T, poormetab = T, fup.censored.dist = FALSE, fup.lod = 0.01, 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"), physiology.matrix = NULL, parameter.matrix = NULL, ...)
```

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.

parameters Parameters from parameterize_steadystate. Not used with httkpop model.

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

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.

output.units Plasma concentration units, either uM or default mg/L. 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 '3compartmentss' is used.

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.

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.

samples Number of samples generated in calculating quantiles.

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

instead of the selected quantile.

default.to.human

Substitutes missing rat values with human values if true.

tissue Desired steady state tissue conentration.

well.stirred.correction

If TRUE (default) then the well-stirred correction (Rowland et al., 1973) is used in the calculation of hepatic clearance for the models that do not include flows for first-pass metabolism (currently, 1compartment and 3compartmentss). This assumes clearance relative to amount unbound in whole blood instead of plasma, but converted for use with plasma concentration.

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.

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.

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.

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

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.

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'.

poormetab TRUE (include poor metabolizers) or FALSE (exclude poor metabolizers)

fup.censored.dist

 $Logical. \ \ Whether \ to \ draw \ \ {\tt Funbound.plasma} \ \ from \ a \ censored \ distribution \ or$

fup.lod The average limit of detection for Funbound.plasma. if fup.censor == TRUE, the Funbound.plasma distribution will be censored below lod/2. Default value

iα Λ Λ1

method The population-generation method to use. Either "virtual individuals" or "di-

rect resampling" (default). Short names may be used: "d" or "dr" for "direct

resampling", and "v" or "vi" for "virtual individuals".

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 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 Diseas 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', 'Non-Hispanic

to include all races and ethnicities in their proportions in the NHANES data.

User-supplied vector must contain one or more of these strings.

physiology.matrix

A data table generated by httkpop_generate().

parameter.matrix

A data table generated by get_httk_params().

Additional parameters passed to calc_analytic_css

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.

calc_mc_oral_equiv 31

Honda6 Target Tissue Non-restrictive Total Max Conc.

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

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.

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

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.

32 calc_mc_oral_equiv

Usage

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

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 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.

calc_mc_oral_equiv 33

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

34 calc_rblood2plasma

	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

Description

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

calc_rblood2plasma 35

Usage

```
calc_rblood2plasma(chem.cas = NULL, chem.name = NULL, params = NULL,
hematocrit = NULL, default.to.human = F, species = "Human",
adjusted.Funbound.plasma = T, suppress.messages = F)
```

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.

params Parameters from parameterize_schmitt.

hematocrit Overwrites default hematocrit value in calculating Rblood2plasma.

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.

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 substitues human fraction unbound and tissue volumes.

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).

Examples

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

36 calc_stats

calc_stats	Calculate the statistics.
carc_stats	Calculate the 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_stats(days, chem.name = NULL, chem.cas = NULL,
  parameters = NULL, stats = c("AUC", "peak", "mean"),
  species = "Human", exclude.fup.zero = F, daily.dose = 1,
  dose = NULL, doses.per.day = NULL, output.units = "uM",
  concentration = "plasma", model = "pbtk", default.to.human = F,
  suppress.messages = F, ...)
```

Arguments

Length of the simulation. days Name of desired chemical. chem.name chem.cas CAS number of desired chemical. Chemical parameters from parameterize_pbtk function, overrides chem.name parameters and chem.cas. stats Desired values (either 'AUC', 'mean', 'peak', or a vector containing any combination). Species desired (either "Rat", "Rabbit", "Dog", "Mouse", or default "Human"). species exclude.fup.zero Whether or not to exclude chemicals with a fraction of unbound plasma equal to zero or include them with a value of 0.005, only used when chem.name, chem.cas, and parameters are not specified. daily.dose Total daily dose, mg/kg BW. Amount of a single dose, mg/kg BW. Overwrites daily.dose. dose doses.per.day Number of doses per day. Desired units (either "mg/L", "mg", "umol", or default "uM"). output.units concentration Desired concentration type, 'blood' or default 'plasma'. 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).

suppress.messages

Whether to suppress output message.

... Arguments passed to solve function.

calc_total_clearance 37

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

peak The highest concentration.

Author(s)

John Wambaugh and Robert Pearce

Examples

```
calc_stats(chem.name='Bisphenol-A',days=100,stats='mean',model='3compartment')
calc_stats(chem.name='Bisphenol-A',days=100,stats=c('peak','mean'),species='Rat')
## Not run:
all.peak.stats <- calc_stats(days=10, doses.per.day = 3, stats = "peak")
## End(Not run)
triclosan.stats <- calc_stats(days=10, chem.name = "triclosan")</pre>
```

calc_total_clearance Calculate the total clearance.

Description

This function calculates the total clearance rate for a one compartment model 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,
  parameters = NULL, species = "Human", suppress.messages = F,
  default.to.human = F, well.stirred.correction = T,
  restrictive.clearance = T, adjusted.Funbound.plasma = T, ...)
```

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.

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

38 calc_vdist

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.

Usage

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

calc_vdist 39

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.

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

Units of L/ kg BW.
```

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")
```

40 chem.invivo.PK.data

```
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

chem.invivo.PK.data 41

Source

Wambaugh et al. 2018 Toxicological Sciences, in press

References

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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

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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.

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42 chem.invivo.PK.data

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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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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

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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

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46 chem.lists

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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

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.

Usage

chem.physical_and_invitro.data

Format

A data.frame containing 565 rows and 33 columns.

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 49

Andersen, M. E. (2015). "Incorporating High-Throughput Exposure Predictions with Dosimetry-Adjusted In Vitro Bioactivity to Inform Chemical Toxicity Testing" Toxicological Sciences, kfv171.

ckd_epi_eq

CKD-EPI equation for GFR.

Description

Predict GFR from serum creatinine, gender, race, and age.

Usage

```
ckd_epi_eq(scr, gender, reth, age_years)
```

Arguments

scr Vector of serum creatinine values in mg/dL.

gender Vector of genders (either 'Male' or 'Female').

reth Vector of races/ethnicities.

age_years Vector of ages in years.

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

50 convert_httk

n
7

Description

Converts HTTK-Pop virtual population into parameters relevant to an HTTK model.

Usage

```
convert_httk(indiv.model.bio, model, this.chem = NULL,
  parameters = NULL, adjusted.Funbound.plasma = T, regression = T,
  well.stirred.correction = T, restrictive.clearance = T,
  concentration = "plasma", clint.pvalue.threshold = 0.05)
```

Arguments

indiv.model.bio

A data.table containing the physiological parameters as expected by HTTK (from httkpop_bio) and Funbound.plasma and Clint values (from draw_fup_clint).

model

Which HTTK model to use. One of '1compartment', '3compartmentss', '3com-

partment', or 'pbtk'.

this.chem

CAS number for the chemical in the HTTK data set (see get_cheminfo) for

which parameters are to be generated.

parameters

A list of chemical-specific model parameters containing at least Funbound.plasma, Clint, and Fhep.assay.correction.

adjusted.Funbound.plasma

Uses adjusted Funbound.plasma when set to TRUE.

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 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.

concentration Blood, plasma, or tissue 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.

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

draw_fup_clint 51

draw_fup_clint	Draw Funbound.plasma and Clint from censored or non-censored distributions.

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.

Usage

```
draw_fup_clint(this.chem = NULL, parameters = NULL, nsamp,
  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 = T,
  clint.pvalue.threshold = 0.05, minimum.Funbound.plasma = 1e-04)
```

Arguments

this.chem	The CAS number of one of the HTTK chemicals (see get_cheminfo).	
parameters	A list of chemical-specific model parameters containing at least Funbound.plasma, Clint, and Fhep.assay.correction.	
nsamp	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).

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.

52 estimate_gfr_ped

Author(s)

Caroline Ring and John Wambaugh

estimate_gfr

Predict GFR.

Description

First predict serum creatinine using smoothing spline, then predict GFR using CKD-EPI equation.

Usage

```
estimate_gfr(gfrtmp.dt)
```

Arguments

gfrtmp.dt

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

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

```
{\tt estimate\_gfr\_ped(BSA)}
```

Arguments

BSA

Vector of body surface areas in m².

Value

Vector of GFRs in mL/min/1.73m².

estimate_hematocrit 53

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

54 export_pbtk_jarnac

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.

Details

digits

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

Desired number of decimal places to round the parameters.

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.

Author(s)

Robert Pearce

Examples

```
## Not run:
export_pbtk_jarnac(chem.name='Nicotine',initial.amounts=list(Agutlumen=1),filename='PBTKmodel.jan')
## End(Not run)
```

export_pbtk_sbml 55

export	nhtk	chm1	
export	DDLK	SDIIIT	

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.

Author(s)

Robert Pearce

Examples

```
## Not run:
export_pbtk_sbml(chem.name='Nicotine',initial.amounts=list(Agutlumen=1),filename='PBTKmodel.xml')
## End(Not run)
```

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 popula-

tion. Defaultis c('Mexican American','Other Hispanic','Non-Hispanic White','Non-Hispa

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

User-supplied vector must contain one or more of these strings.

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 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

gen_height_weight 57

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_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.

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

58 get_cheminfo

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 paramterized for a given species.

Usage

```
get_cheminfo(info = "CAS", species = "Human", exclude.fup.zero = NA,
fup.lod.default = 0.005, model = "3compartmentss",
    default.to.human = F)
```

Arguments

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

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

"Funbound.plasma", "DSSTox_Substance_Id", "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").

exclude.fup.zero

Whether or not to exclude chemicals with a fraction of unbound plasma equal to zero or include them with a value of fup.lod.default. Defaults to FALSE for

'3compartmentss' and TRUE for pk models and schmitt.

fup.lod.default

model

Default value used for fraction of unbound plasma for chemicals where measured value was below the limit of detection. Default value is 0.0005.

sured value was below the fiffit of detection. Default value is 0.0003

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

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.

Value

info

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

Author(s)

John Wambaugh and Robert Pearce

get_chem_id 59

Examples

```
## Not run:
# 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_cheminfo(info="all")
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",
"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)
## End(Not run)
```

get_chem_id

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 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)
```

60 get_gfr_category

Arguments

chem.cas CAS regstry number

chem.name Chemical name

dtxsid DSSTox Substance identifier

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_httk_params 61

get_httk_params	Converts the HTTK-Pop population data table to a table of the param-
	eters 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

```
get_httk_params(indiv_dt, chemcas = NULL, parameters = NULL, model,
poormetab, fup.censored.dist = FALSE, fup.meas.cv = 0.4,
clint.meas.cv = 0.3, fup.pop.cv = 0.1, clint.pop.cv = 0.1,
fup.lod = 0.01, adjusted.Funbound.plasma = T, regression = T,
well.stirred.correction = T, restrictive.clearance = T,
concentration = "plasma", clint.pvalue.threshold = 0.05)
```

Arguments

indiv_dt	A data table generated by httkpop_generate().	
chemcas	The CAS number of one of the HTTK chemicals (see <pre>get_cheminfo</pre>). Defaults to NULL.	
parameters	A list of chemical-specific model parameters containing at least Funbound.plasma, Clint, and Fhep.assay.correction, otherwise defaults to NULL.	
mode1	One of the HTTK models: "1compartment", "3compartmentss", "3compartment", or "pbtk".	
poormetab	TRUE (include poor metabolizers) or FALSE (exclude poor metabolizers)	
fup.censored.dist		
	Logical. Whether to draw Funbound.plasma from a censored distribution or not.	
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.	
fup.lod	The average limit of detection for Funbound.plasma. if fup.censor == TRUE, the Funbound.plasma distribution will be censored below lod/2. Default value is 0.01.	
adjusted.Funbound.plasma		

Uses adjusted Funbound.plasma when set to TRUE.

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

If TRUE (default) then the well-stirred correction (Rowland et al., 1973) is used in the calculation of hepatic clearance for the models that do not include flows for first-pass metabolism (currently, 1compartment and 3compartmentss). This assumes clearance relative to amount unbound in whole blood instead of plasma, but converted for use with plasma concentration.

62 get_lit_cheminfo

```
restrictive.clearance
```

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

Blood, plasma, or tissue concentration. 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.

Value

A data.table whose columns correspond to the parameters of the HTTK model specified in model, and whose rows correspond to the individuals (rows) of indiv_dt.

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)</pre>
httk_param <- get_httk_params(indiv_dt=indiv_examp,</pre>
chemcas="80-05-7",
model="1compartment",
poormetab=TRUE,
fup.censored.dist=TRUE)
```

get_lit_cheminfo

Get literature Chemical Information.

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")
```

Arguments

info A single character vector (or collection of character vectors) from "Compound", "CAS", "MW", "Raw.F ","p.val","Concentration..uM.","Css_lower_5th_perc.mg.L.","Css_median_perc.mg.L.","Css_up

and "Species".

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

get_lit_css 63

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

```
## Not run:
get_lit_cheminfo()
get_lit_cheminfo(info=c('CAS','MW'))
## End(Not run)
```

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.

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 = F)
```

64 get_lit_css

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, $1\ \mathrm{or}\ 10$

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.

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_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 65

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 = F, 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.

 $output.\,units \qquad Units of dose, default of 'mg' for \,mg/kg \,BW/\,day \,or \,'mol' \,for \,mol/\,kg \,BW/\,day.$

clearance.assay.conc

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

uM.

... Additional parameters passed to get_lit_css.

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 retrives 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)
```

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

Value

The paramters, either a single value, a named list for a single chemical, or a list of lists

Examples

```
get_physchem_param(param = 'logP', chem.CAS = '80-05-7') get_physchem_param(param = c('logP', 'MW'), chem.CAS = c('80-05-7', '81-81-2'))
```

get_rblood2plasma 67

σet	rblood2plasma	
20.	I DIOUGEDIASIIIA	

Get ratio of the blood concentration to the plasma concentration.

Description

This function retrieves the in vivo ratio of the blood concentration to the plasma concentration.

Usage

```
get_rblood2plasma(chem.name = NULL, chem.cas = NULL,
  species = "Human", default.to.human = F)
```

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.

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

default.to.human

Substitutes missing animal values with human values if true.

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 ~~

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)
```

68 hematocrit_infants

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

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

honda.ivive 69

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

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.

70 httkpop

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.

Examples

```
honda.ivive(method = "Honda1", tissue = NULL)
```

howgate

Howgate 2006

Description

This data set is only used in Vignette 5.

Usage

howgate

Format

An object of class data. table (inherits from data. frame) with 24 rows and 11 columns.

httkpop

httkpop: Virtual population generator for HTTK.

Description

The httkpop package generates virtual population physiologies for use in population TK.

The httkpop package generates virtual population physiologies for use in population TK.

Main function to generate a population

If you just want to generate a table of (chemical-independent) population physiology parameters, use httkpop_generate.

If you just want to generate a table of (chemical-independent) population physiology parameters, use httkpop_generate.

httkpop_bio 71

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 get_httk_params.

Author(s)

Caroline Ring

Caroline Ring

References

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

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

httkpop_bio

Convert HTTK-Pop-generated parameters to HTTK physiological parameters

Description

Convert HTTK-Pop-generated parameters to HTTK physiological parameters

Usage

httkpop_bio(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

Description

Generate a virtual population by directly resampling the NHANES data.

Usage

```
httkpop_direct_resample(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"))
```

Arguments

The desired number of individuals in the virtual population. nsamp need not be nsamp

provided if gendernum is provided.

Optional: A named list giving the numbers of male and female individuals gendernum

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

User-supplied vector must contain one or more of these strings.

The kidney function categories to include in the population. Default is c('Normal', 'Kidney Diseas gfr_category

to include all kidney function levels.

reths Optional: a character vector giving the races/ethnicities to include in the popula-

tion. Defaultis c('Mexican American','Other Hispanic','Non-Hispanic White','Non-Hispa

to include all races and ethnicities in their proportions in the NHANES data.

User-supplied vector must contain one or more of these strings.

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)
```

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 popula-

tion. Defaultis c('Mexican American','Other Hispanic','Non-Hispanic White','Non-Hispanic

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

 $Optional: The weight categories to include in the population. \ Default is \verb|c('Underweight', 'Normal')| and the population of the popul$

User-supplied vector must contain one or more of these strings.

Value

A data.table where each row represents an individual, and each column represents a demographic, anthropometric, or physiological parameter.

74 httkpop_generate

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, 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"))
```

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.

httkpop_generate 75

weight_category

Optional: The weight categories to include in the population. Default is c('Underweight', 'Normal

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 Diseas

to include all kidney function levels.

reths Optional: a character vector giving the races/ethnicities to include in the popula-

tion. Defaultis c('Mexican American','Other Hispanic','Non-Hispanic White','Non-Hispa

to include all races and ethnicities in their proportions in the NHANES data.

User-supplied vector must contain one or more of these strings.

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

Examples

```
## Not run:
#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'))
## End(Not run)
```

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 = 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"))
```

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

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 Diseas

to include all kidney function levels.

reths Optional: a character vector giving the races/ethnicities to include in the popula-

tion. Defaultis c('Mexican American','Other Hispanic','Non-Hispanic White','Non-Hispa

to include all races and ethnicities in their proportions in the NHANES data.

User-supplied vector must contain one or more of these strings.

Value

A data.table where each row represents an individual, and each column represents a demographic, anthropometric, or physiological parameter.

in.list 77

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	Convenience Boolean (yes/no) functions to identify chemical member- ship in several key lists.
111.1130	

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.

78 is.httk

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: http://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"))</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 fifty
# chemicals. To build a table with all available chemicals use:
# for (this.cas in httk.table$CAS)
for (this.cas in httk.table$CAS[1:50])
  this.index <- httk.table$CAS==this.cas</pre>
  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>
```

is.httk 79

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

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

80 is_in_inclusive

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: http://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>
\verb|httk.table[,"Rat"] <- ""
httk.table[,"NHANES"] <- ""</pre>
httk.table[,"Tox21"] <- ""</pre>
httk.table[,"ToxCast"] <- ""</pre>
httk.table[,"ExpoCast"] <- ""
httk.table[,"PBTK"] <- ""</pre>
# To make this example run quickly, this loop is only over the first fifty
# chemicals. To build a table with all available chemicals use:
# for (this.cas in httk.table$CAS)
for (this.cas in httk.table$CAS[1:50])
  this.index <- httk.table$CAS==this.cas</pre>
  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>
}
```

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

Χ

A numeric value, or vector of values.

johnson 81

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

An object of class data.table (inherits from data.frame) with 60 rows and 11 columns.

kidney_mass_children Predict kidney mass for children.

Description

For individuals under age 18, predict kidney mass from weight, height, and gender. using equations from Ogiu et al.

```
kidney_mass_children(weight, height, gender)
```

82 load_sipes2017

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.

liver_mass_children Predic

Predict liver mass for children.

Description

For individuals under 18, predict the liver mass from height, weight, and gender, using equations from Ogiu et al.

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').

Value

A vector of liver masses in kg.

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.

```
load_sipes2017(load.image = T, overwrite = F,
  target.env = .GlobalEnv)
```

lump_tissues 83

Arguments

load.image If overwrite=TRUE (DEFAULT)) then the default HTTK chemical data plus the

any new data/predictions from Sipes et al. (2017) will be quickly loaded. This is the same as load.image=F, but much faster, however any other data added by

the user will be deleted.

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.

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)

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.

Examples

```
## Not run:
chem.physical_and_invitro.data <- load_sipes2017()
chem.physical_and_invitro.data <- load_sipes2017(overwrite=T)
## End(Not run)</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.

```
lump_tissues(Ktissue2pu.in, tissuelist = NULL, species = "Human")
```

84 lung_mass_children

Arguments

Ktissue2pu.in List of partition coefficients from predict_partitioning_schmitt.

tissuelist Specifies compartment names and tissues groupings. Remaining tissues in tis-

sue.data are lumped in the rest of the body.

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

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 a lumped tissue, the words in the vector are the Schmitt (2008) tissues that are to be lumped, for example: tissuelist<-list(Rapid=c("Brain","Kidney")) species 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

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>
```

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.

mcnally_dt 85

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.

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.

86 monte_carlo

References

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

monte_carlo

Monte Carlo for pharmacokinetic models

Description

This function performs Monte Carlo to assess uncertainty and variability for toxicokinetic models.

Usage

```
monte_carlo(params, which.quantile = 0.95, cv.params = NULL,
 censored.params = NULL, samples = 1000,
 name.model = "calc_analytic_css", output.col.model = NA,
 return.samples = F, ...)
```

Arguments

params

All parameters needed by the function indicated by the argument "name.model". These paramters that are also listed in either cv.params or censored.params are sampled using Monte Carlo.

which quantile This argument specifies which quantiles are to be calculated. It can be a vector or a single value. It defaults to the 0.95 quantile (95%).

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 "params". New values are sampled with mean equal to the value in "params" 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 paramter). 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 quan-

name.model

This argument is a character vector giving the name of the model to be sampled. Defaults to 'calc_analytic_css'.

output.col.model

If the evaluation of the function indicated by "model" returns a list, then model.output.col is the element from that list that is sampled and is used for calculating quantiles. Defaults to NA (i.e., the function returns a single value).

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

Additional arguments passed to name.model.

monte_carlo 87

Author(s)

John Wambaugh

Examples

```
#Example from httk jss paper:
## Not run:
library(ggplot2)
library(scales)
vary.params <- NULL
params <- parameterize_pbtk(chem.name = "Zoxamide")</pre>
for(this.param in names(subset(params,
names(params) != "Funbound.plasma"))) vary.params[this.param] <- .2</pre>
censored.params <- list(Funbound.plasma = list(cv = 0.2, lod = 0.01))</pre>
set.seed(1)
out <- monte_carlo(params, cv.params = vary.params,</pre>
censored.params = censored.params, return.samples = T,
model = "pbtk", suppress.messages = T)
zoxamide <- ggplot(as.data.frame(out), aes(out)) +</pre>
geom_histogram(fill="blue", binwidth=1/6) + scale_x_log10() +
ylab("Number of Samples") + xlab("Steady State Concentration (uM)") +
theme(axis.text = element_text(size = 16),
axis.title = element_text(size = 16))
print(zoxamide)
# Fig 1 in Wambaugh et al. (2015) SimCYP vs. our predictions:
vary.params <- list(BW=0.3)</pre>
vary.params[["Vliverc"]]<-0.3</pre>
vary.params[["Qgfrc"]]<-0.3</pre>
vary.params[["Qtotal.liverc"]]<-0.3</pre>
vary.params[["million.cells.per.gliver"]]<-0.3</pre>
vary.params[["Clint"]]<-0.3</pre>
censored.params<-list(Funbound.plasma=list(cv=0.3,lod=0.01))</pre>
pValues <- get_cheminfo(c("Compound","CAS","Clint.pValue"))</pre>
pValues.rat <- get_cheminfo(c("Compound","CAS","Clint.pValue"),species="Rat")</pre>
Wetmore.table <- NULL
for (this.CAS in get_cheminfo(model="3compartmentss")){
  if (this.CAS %in% get_wetmore_cheminfo()){
    print(this.CAS)
    these.params <- parameterize_steadystate(chem.cas=this.CAS)</pre>
    if (these.params[["Funbound.plasma"]] == 0.0)
      these.params[["Funbound.plasma"]] <- 0.005</pre>
    }
    these.params[["Fhep.assay.correction"]] <- 1</pre>
    vLiver.human.values <- monte_carlo(these.params,</pre>
                                          cv.params=vary.params,
                                          censored.params=censored.params,
                                          which.quantile=c(0.05, 0.5, 0.95),
```

88 nhanes_mec_svy

```
output.units="mg/L",
                                         model='3compartmentss',
                                         suppress.messages=T,
                                         well.stirred.correction=F,
                                         Funbound.plasma.correction=F)
    percentiles <- c("5","50","95")</pre>
    for (this.index in 1:3)
    {
      this.row <- as.data.frame(get_wetmore_css(chem.cas=this.CAS,</pre>
                                 which.guantile=as.numeric(percentiles[this.index])/100))
      this.row <- cbind(this.row, as.data.frame(vLiver.human.values[this.index]))</pre>
      this.row <- cbind(this.row, as.data.frame(percentiles[this.index]))</pre>
      this.row <- cbind(this.row, as.data.frame("Human"))</pre>
      this.row <- cbind(this.row, as.data.frame(this.CAS))</pre>
      this.row <- cbind(this.row, as.data.frame(pValues[pValues$CAS==this.CAS,</pre>
                                                  "Human.Clint.pValue"]<0.05))
      colnames(this.row) <- c("Wetmore", "Predicted", "Percentile", "Species",</pre>
                               "CAS", "Systematic")
      if (is.na(this.row["Systematic"])) this.row["Systematic"] <- F</pre>
      Wetmore.table <- Wetmore.table <- rbind(Wetmore.table,this.row)</pre>
    }
 }
}
scientific_10 <- function(x) {</pre>
  out <- gsub("1e", "10^", scientific_format()(x))</pre>
 out <- gsub("\+","",out)
 out <- gsub("10^01","10",out)
 out <- parse(text=gsub("10^00","1",out))</pre>
Fig1 <- ggplot(Wetmore.table, aes(Predicted, Wetmore, group = CAS)) +
  geom_line() +
  geom_point(aes(colour=factor(Percentile), shape=factor(Percentile))) +
  scale_colour_discrete(name="Percentile") +
  scale_shape_manual(name="Percentile", values=c("5"=21, "50"=22,"95"=24)) +
  scale_x_log10(expression(paste(C[ss],"Predicted(mg/L)with Refined Assumptions")),
                label=scientific_10) +
  scale_y_log10(expression(paste(C[ss]," Wetmore ",italic("et al.")," (2012) (mg/L)")),
                label=scientific_10) +
  geom_abline(intercept = 0, slope = 1,linetype="dashed")+
  theme_bw()+
  theme(legend.position="bottom", text = element_text(size=18))
print(Fig1)
Fig1a.fit <- lm(log(Wetmore) ~ log(Predicted)*Percentile, Wetmore.table)
## End(**Not run**)
## End(Not run)
```

Obach2008 89

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.

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

http://www.cdc.gov/nhanes/nhanes_questionnaires.htm

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

Obach2008

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

NHANES Exposure Data

Description

This data set is only used in Vignette 6.

Usage

onlyp

Format

An object of class data. table (inherits from data. frame) with 1060 rows and 5 columns.

```
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.

parameterize_1comp 91

parameterize_1comp

Description

This function initializes the parameters needed in the function solve_1comp.

Usage

```
parameterize_1comp(chem.cas = NULL, chem.name = NULL,
  species = "Human", default.to.human = F,
  adjusted.Funbound.plasma = T, regression = T,
  restrictive.clearance = T, well.stirred.correction = T,
  suppress.messages = F, clint.pvalue.threshold = 0.05,
  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.

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 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 and hepatic bioavailability, protein binding is not taken into account (set to 1) in liver clearance if FALSE.

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-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).

Value

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.

92 parameterize_3comp

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

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,
  species = "Human", default.to.human = F, force.human.clint.fup = F,
  clint.pvalue.threshold = 0.05, adjusted.Funbound.plasma = T,
  regression = T, suppress.messages = F,
  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.

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

default.to.human

Substitutes missing animal values with human values if true.

parameterize_3comp 93

force.human.clint.fup

Forces use of human values for hepatic intrinsic clearance and fraction of unbound plasma if true.

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.

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.

94 parameterize_pbtk

Author(s)

Robert Pearce and 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.

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,
   species = "Human", default.to.human = F, tissuelist = list(liver =
   c("liver"), kidney = c("kidney"), lung = c("lung"), gut = c("gut")),
   force.human.clint.fup = F, clint.pvalue.threshold = 0.05,
   adjusted.Funbound.plasma = T, regression = T,
   suppress.messages = F, 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.

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 unbound plasma if true.

parameterize_pbtk 95

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 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.

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.

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.

Qliverf Fraction of cardiac output flowing to the liver.

96 parameterize_schmitt

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.
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

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

Description

This function provides the necessary parameters to run predict_partitioning_schmitt, excluding the data in tissue.data.

```
parameterize_schmitt(chem.cas = NULL, chem.name = NULL,
   species = "Human", default.to.human = F, force.human.fup = F,
   suppress.messages = F, minimum.Funbound.plasma = 1e-04)
```

parameterize_schmitt 97

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.

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

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.

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).

Details

When species is specified as rabbit, dog, or mouse, the human unbound fraction is substituted.

force.human.fup calculates Funbound.plasma.corrected with the human lipid fractional volume in plasma.

Value

Funbound.plasma

corrected unbound fraction in plasma

unadjusted.Funbound.plasma

measured unbound fraction in plasma (0.005 if below limit of detection)

Pow octonol:water partition coefficient (not log transformed)

pKa_Donor compound H dissociation equilibirum constant(s)
pKa_Accept compound H association equilibirum 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

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,
    species = "Human", clint.pvalue.threshold = 0.05,
    default.to.human = F, human.clint.fup = F,
    adjusted.Funbound.plasma = T, restrictive.clearance = T,
    fup.lod.default = 0.005, suppress.messages = F,
    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.

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 threshold 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).

pc.data 99

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

Examples

```
parameters <- parameterize_steadystate(chem.name='Bisphenol-A',species='Rat')
parameters <- parameterize_steadystate(chem.cas='80-05-7')</pre>
```

pc.data

Partition Coefficient Data

Description

Measured rat in vivo partition coefficients and data for predicting them.

Usage

pc.data

Format

A data.frame.

100 pharma

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.

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

physiology.data 101

Format

An object of class data. frame with 954 rows and 14 columns.

Source

https://comptox.epa.gov/dashboard/chemical_lists/swisspharma

References

Wambaugh et al. "Assessing Toxicokinetic Uncertainty and Variability in Risk Prioritization ", submitted.

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

```
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,
   species = "Human", default.to.human = F, parameters = NULL,
   alpha = 0.001, adjusted.Funbound.plasma = T, regression = T,
   regression.list = c("brain", "adipose", "gut", "heart", "kidney",
   "liver", "lung", "muscle", "skin", "spleen", "bone"), tissues = NULL,
   minimum.Funbound.plasma = 1e-04)
```

Arguments

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

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).

parameters Chemical parameters from the parameterize_schmitt function, overrides chem.name

and chem.cas.

alpha Ratio of Distribution coefficient D of totally charged species and that of the

neutral form

 $\verb|adjusted.Funbound.plasma|\\$

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).

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.

rfun 103

Value

Returns tissue to unbound plasma partition coefficients for each tissue.

Author(s)

Robert Pearce

Examples

```
predict_partitioning_schmitt(chem.name='ibuprofen',regression=FALSE)
```

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

104 sipes2017

r_left_censored_norm	Returns draws from a normal distribution with a lower censoring limit of lod (limit of detection)
	ey tea (time ey derection)

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.	

Value

A vector of samples from the specified censored distribution.

sipes2017	Sipes et al. 2017 data	

Description

This table includes data predicted with Simulations Plus' ADMET predictor, used in load_sipes2017, that was used in Sipes et al. 2017. The column names are equivalent to those of chem.physical_and_invitro.data.

Usage

sipes2017

Format

data.frame

Author(s)

Nisha Sipes

Source

ADMET, Simulations Plus

sipes2017.table 105

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.

sipes2017.table

Physico-chemical properties and toxicokinetics, measured values and Sipes et al. (2017)

Description

This is an image of the chem.phys_and_invitro.data table that has had the Sipes et al. (2017) AD-MET predictions adfdded to it. The 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.

Usage

sipes2017.table

Format

A data.frame containing 9211 rows and 47 columns.

Author(s)

John Wambaugh

Source

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

References

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

EPI Suite, http://www.epa.gov/opptintr/exposure/pubs/episuite.htm

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106 sipes 2017.table

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Paixao, P., Gouveia, L. F., & Morais, J. A. (2012). Prediction of the human oral bioavailability by using in vitro and in silico drug related parameters in a physiologically based absorption model. International journal of pharmaceutics, 429(1), 84-98.

Pirovano, Alessandra, et al. "QSARs for estimating intrinsic hepatic clearance of organic chemicals in humans." Environmental toxicology and pharmacology 42 (2016): 190-197.

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.

Shibata, Y., Takahashi, H., Chiba, M. and Ishii, Y. (2002). Prediction of Hepatic Clearance and Availability by Cryopreserved Human Hepatocytes: An Application of Serum Incubation Method. Drug Metabolism and Disposition 30(8), 892-896, 10.1124/dmd.30.8.892.

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.

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Wetmore, B. A., Wambaugh, J. F., Ferguson, S. S., Li, L., Clewell, H. J., 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. (2013). Relative Impact of Incorporating Pharmacokinetics on Predicting In Vivo Hazard and Mode of Action from High-Throughput In Vitro Toxicity Assays. Toxicological Sciences 132(2), 327-346, 10.1093/toxsci/kft012.

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.

skeletal_muscle_mass 107

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.

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 J Cachexia Sarcopenia Muscle 2012 3:25-29.

Usage

```
skeletal_muscle_mass_children(gender, age_years)
```

Arguments

gender Vector of genders (either 'Male' or 'Female').

age_years Vector of ages in years.

108 solve_1comp

Value

Vector of skeletal muscle masses in kg.

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.

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.

```
solve_1comp(chem.name = NULL, chem.cas = NULL, times = NULL,
  parameters = NULL, daily.dose = 1, dose = NULL,
  doses.per.day = NULL, days = 10, tsteps = 4,
  suppress.messages = F, species = "Human", output.units = "uM",
  plots = F, initial.values = NULL, iv.dose = F, method = "lsoda",
  rtol = 1e-08, atol = 1e-12, default.to.human = F,
  dosing.matrix = NULL, recalc.elimination = F,
  adjusted.Funbound.plasma = T, regression = T,
  restrictive.clearance = T, well.stirred.correction = T,
  minimum.Funbound.plasma = 1e-04, ...)
```

solve_1comp 109

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.

times Optional time sequence for specified number of days.

parameters Chemical parameters from parameterize_1comp function, overrides chem.name

and chem.cas.

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

dose Amount of a single dose, mg/kg BW. Overwrites daily.dose.

doses.per.day Number of doses per day.

days Length of the simulation.

tsteps The number time steps per hour.

suppress.messages

Whether or not the output message is suppressed.

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

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

plots Plots all outputs if true.

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.

iv. dose Simulates a single i.v. dose if true.

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.

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.

recalc.elimination

Whether or not to recalculate the elimination rate.

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

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.

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).

. . . Additional arguments passed to the integrator.

solve_1comp

Details

Note that the model parameters have units of hours while the model output is in days.

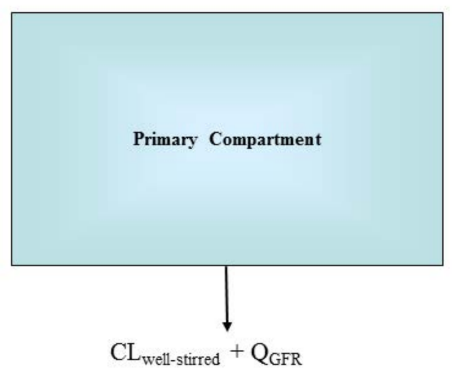
Default value of NULL for doses.per.day solves for a single dose.

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

AUC is area under plasma concentration curve.

Model Figure





altalt

solve_3comp 111

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. function does. ~~

Usage

```
solve_3comp(chem.name = NULL, chem.cas = NULL, times = NULL,
  parameters = NULL, days = 10, tsteps = 4, daily.dose = 1,
  dose = NULL, doses.per.day = NULL, initial.values = NULL,
  plots = F, suppress.messages = F, species = "Human", iv.dose = F,
  output.units = "uM", method = "lsoda", rtol = 1e-08,
  atol = 1e-12, default.to.human = F, recalc.blood2plasma = F,
  recalc.clearance = F, dosing.matrix = NULL,
  adjusted.Funbound.plasma = T, regression = T,
  restrictive.clearance = T, minimum.Funbound.plasma = 1e-04, ...)
```

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.
times	Optional time sequence for specified number of days. The dosing sequence begins at the beginning of times.
parameters	Chemical parameters from parameterize_3comp function, overrides chem.name and chem.cas.
days	Length of the simulation.

112 solve_3comp

tsteps The number time steps per hour. daily.dose Total daily dose, mg/kg BW.

dose Amount of a single dose, mg/kg BW. Overwrites daily.dose.

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.

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

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.

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 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).

. . . Additional arguments passed to the integrator.

Details

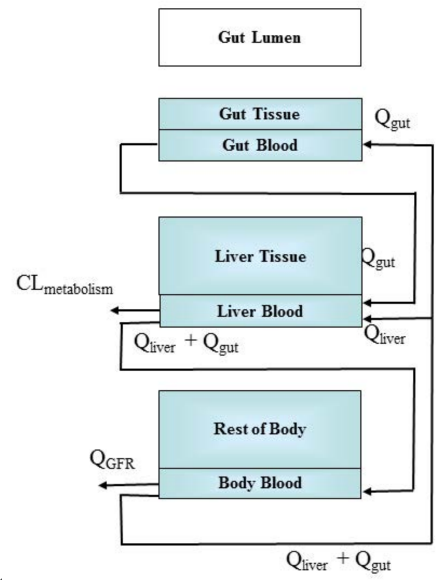
Note that the model parameters have units of hours while the model output is in days.

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 113



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

114 solve_pbtk

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,dose=.5,days=1,tsteps=2)
params <-parameterize_3comp(chem.cas="80-05-7")
solve_3comp(parameters=params)</pre>
```

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, times = NULL,
   parameters = NULL, days = 10, tsteps = 4, daily.dose = 1,
   dose = NULL, doses.per.day = NULL, initial.values = NULL,
   plots = F, suppress.messages = F, species = "Human", iv.dose = F,
   output.units = "uM", method = "lsoda", rtol = 1e-08,
   atol = 1e-12, default.to.human = F, recalc.blood2plasma = F,
   recalc.clearance = F, dosing.matrix = NULL,
   adjusted.Funbound.plasma = T, regression = T,
   restrictive.clearance = T, minimum.Funbound.plasma = 1e-04, ...)
```

Arguments

plots

Either the chemical name, CAS number, or the parameters must be specified. chem.name chem.cas Either the chemical name, CAS number, or the parameters must be specified. Optional time sequence for specified number of days. Dosing sequence begins times at the beginning of times. Chemical parameters from parameterize_pbtk function, overrides chem.name parameters and chem.cas. days Length of the simulation. The number of time steps per hour. tsteps Total daily dose, mg/kg BW. daily.dose Amount of a single dose, mg/kg BW. Overwrites daily.dose. dose 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 all outputs if true.

solve_pbtk 115

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.

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

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).

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.

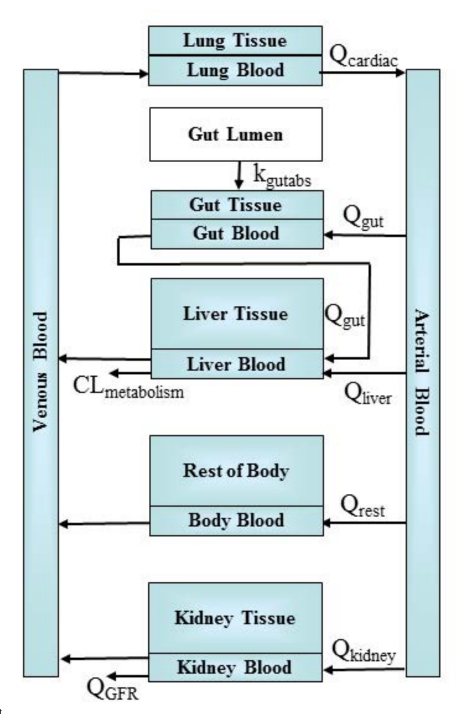
The compartments used in this model are the gutlumen, gut, liver, kidneys, veins, arteries, lungs, and the rest of the body.

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

116 solve_pbtk



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.

spleen_mass_children 117

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

```
solve_pbtk(chem.name='Bisphenol-A',dose=.5,days=1,doses.per.day=2,tsteps=2)
out <- solve_pbtk(chem.name='bisphenola',dose=0,output.units='mg',</pre>
                   plots=TRUE,initial.values=c(Agut=200))
params <- parameterize_pbtk(chem.cas="80-05-7")</pre>
solve_pbtk(parameters=params)
## Not run:
parameters <- parameterize_pbtk(chem.name = "triclosan", species = "rat")</pre>
parameters["Funbound.plasma"] <- 0.1</pre>
out <- solve_pbtk(parameters=parameters)</pre>
library("ggplot2")
out <- solve_pbtk(chem.name = "Bisphenol A", days = 50, doses.per.day = 3)
plot.data <- as.data.frame(out)</pre>
css <- calc_analytic_css(chem.name = "Bisphenol A")</pre>
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)
## End(Not run)
```

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.

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').
```

118 spline_heightweight

Value

A vector of spleen masses in kg.

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 119

Description

Smoothing splines and KDE residuals pre-calculated from NHANES hematocrit and age data by race/ethnicity and gender.

Usage

spline_hematocrit

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	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

120 tissue.data

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

Tables.Rdata.stamp

A timestamp of table creation

Description

A timestamp of table creation

Usage

Tables.Rdata.stamp

Format

An object of class character of length 1.

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

tissue_masses_flows 121

Format

A data.frame containing 13 rows and 20 columns.

Author(s)

John Wambaugh, Robert Pearce, and Nisha Sipes

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.

122 tissue_scale

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

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

ToxCast2015subset 123

ToxCast2015subset

ToxCast and 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.

Usage

ToxCast2015subset

Format

A data.table with 62412 rows and 5 columns

Author(s)

Caroline Ring

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.

124 wambaugh2019

wambaugh2019

in vitro Toxicokinetic Data from Wambaugh et al. (submitted)

Description

These data are the new HTTK in vitro data for chemicals reported in Wambaugh et al. (submitted) 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 125

Source

Wambaugh et al. (submitted)

References

Wambaugh et al. "Assessing Toxicokinetic Uncertainty and Variability in Risk Prioritization", submitted.

wambaugh2019.nhanes

NHANES Chemical Intake Rates for chemicals in Wambaugh et al. (submitted))

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. (submitted)

References

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

126 wambaugh2019.raw

wambaugh2019.raw Raw Bayesian in vitro Toxicokinetic Data Analysis from Wambaugh et al. (submitted)

Description

These data are the new HTTK in vitro data for chemicals reported in Wambaugh et al. (submitted) 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 127

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. (submitted)

References

Wambaugh et al. "Assessing Toxicokinetic Uncertainty and Variability in Risk Prioritization", submitted

128 well_param

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. (submitted)

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.

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

Wetmore.data 129

Format

A data frame with 53940 rows and 10 variables:

area_bottom cell_yield diam

sysID

v_total

v_working

well_desc

well_number

Author(s)

Greg Honda

Source

http://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.

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.

130 wfl

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.

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

wfl 131

Format

```
A data.table object with variables
```

Sex 'Male' or 'Female' Length length in cm

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

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

http://www.cdc.gov/growthcharts/who/girls_weight_head_circumference.htm and http://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

Index

*Topic Export	calc_mc_css, 27
export_pbtk_jarnac, 54	calc_mc_oral_equiv, 31
export_pbtk_sbml, 55	*Topic cheminformatics
*Topic Literature	get_chem_id, 59
get_lit_cheminfo, 62	*Topic datasets
get_lit_css, 63	Tables.Rdata.stamp, 120
get_lit_oral_equiv, 65	*Topic data
*Topic Monte-Carlo	age_dist_smooth, 6
calc_mc_css, 27	armitage_input, 10
<pre>calc_mc_oral_equiv, 31</pre>	bmiage, 13
get_lit_css, 63	chem.invivo.PK.aggregate.data,40
<pre>get_lit_oral_equiv, 65</pre>	chem.invivo.PK.data,40
monte_carlo, 86	chem.invivo.PK.summary.data,43
*Topic Parameter	chem.lists,46
available_rblood2plasma, 11	<pre>chem.physical_and_invitro.data, 47</pre>
<pre>calc_elimination_rate, 23</pre>	howgate, 70
<pre>calc_hepatic_clearance, 25</pre>	johnson, 81
calc_ionization, 26	mcnally_dt, 85
calc_rblood2plasma,34	nhanes_mec_svy, 88
calc_total_clearance, 37	Obach2008, 89
<pre>calc_vdist, 38</pre>	onlyp, 90
get_rblood2plasma,67	pc.data,99
<pre>lump_tissues, 83</pre>	pharma, 100
parameterize_1comp,91	physiology.data, 101
parameterize_3comp,92	sipes2017, 104
parameterize_pbtk,94	sipes2017.table, 105
parameterize_schmitt,96	spline_heightweight, 118
parameterize_steadystate,98	spline_hematocrit, 119
<pre>predict_partitioning_schmitt, 102</pre>	spline_serumcreat, 119
*Topic Retrieval	tissue.data, 120
<pre>get_cheminfo, 58</pre>	ToxCast2015subset, 123
get_lit_cheminfo, 62	wambaugh2019, <u>124</u>
*Topic Solve	wambaugh2019.nhanes, 125
calc_analytic_css, 15	wambaugh2019.raw, 126
calc_stats, 36	wambaugh2019.seem3, 128
honda.ivive, 69	well_param, 128
solve_1comp, 108	Wetmore.data, 129
solve_3comp, 111	Wetmore2012, 130
solve_pbtk, 114	wf1, 130
*Topic Statistics	*Topic httk-pop
calc_stats, 36	age_dist_smooth, 6
*Topic Steady-State	age_draw_smooth, 7
calc_css, 21	blood_weight, 12

INDEX 133

bmiage, 13	calc_ionization, 26
ckd_epi_eq,49	calc_mc_css, 27
estimate_gfr, 52	<pre>calc_mc_oral_equiv, 31</pre>
estimate_gfr_ped, 52	calc_rblood2plasma, 34
estimate_hematocrit, 53	calc_stats, 36
<pre>gen_age_height_weight, 56</pre>	calc_total_clearance, 37
gen_height_weight, 57	calc_vdist, 38
get_gfr_category, 60	chem.invivo.PK.aggregate.data,40
<pre>get_weight_class, 67</pre>	chem.invivo.PK.data, 40
hematocrit_infants, 68	chem.invivo.PK.summary.data,43
httkpop, 70	chem.lists, 46
httkpop_bio, 71	<pre>chem.physical_and_invitro.data, 47</pre>
httkpop_direct_resample, 72	ckd_epi_eq, 49
httkpop_direct_resample_inner, 73	convert_httk, 50
httkpop_generate, 74	
httkpop_virtual_indiv, 76	draw_fup_clint, 50 , 51
is_in_inclusive, 80	
mcnally_dt, 85	estimate_gfr, 52
nhanes_mec_svy, 88	estimate_gfr_ped,52
rfun, 103	estimate_hematocrit, 53
spline_heightweight, 118	export_pbtk_jarnac,54
spline_hematocrit, 119	<pre>export_pbtk_sbml, 55</pre>
spline_serumcreat, 119	
tissue_masses_flows, 121	<pre>gen_age_height_weight, 56</pre>
tissue_scale, 122	<pre>gen_height_weight, 57</pre>
well_param, 128	get_chem_id, 59
wfl, 130	get_cheminfo, 50, 51, 58, 61
*Topic package	get_gfr_category, 60
httk-package, 4	get_httk_params, 61, 71
	<pre>get_lit_cheminfo, 62</pre>
add_chemtable, 5	get_lit_css, 63
age_dist_smooth, 6	<pre>get_lit_oral_equiv, 65</pre>
age_draw_smooth, 7	get_physchem_param, 66
armitage_estimate_sarea,8	get_rblood2plasma,67
armitage_eval,8	<pre>get_weight_class, 67</pre>
armitage_input, 10	
available_rblood2plasma, 11	hematocrit_infants, 68
. 10	honda.ivive, 69
blood_mass_correct, 12	howgate, 70
blood_weight, 12, 12	httk (httk-package), 4
bmiage, 13	httk-package, 4
body_surface_area, 14	httkpop, 70
bone_mass_age, 14	httkpop-package (httkpop), 70
brain_mass, 15	httkpop_bio, 50, 71
calc analytic acc 15	httkpop_direct_resample, 72
calc_analytic_css,15 calc_analytic_css_1comp,17	httkpop_direct_resample_inner, 73
calc_analytic_css_1comp, 17 calc_analytic_css_3comp, 18	httkpop_generate, 70, 71, 74
• • • • • • • • • • • • • • • • • • • •	httkpop_virtual_indiv,76
calc_analytic_css_3compss, 19	in list 77 80
calc_analytic_css_pbtk,20 calc_css,21	in.list, 77, 80
calc_css, 21 calc_elimination_rate, 23	is.expocast(in.list),77 is.httk,78,78
calc_errmination_rate, 25	is.nhanes (in.list), 77
care_nepatic_creat affec, 23	13.1111diles (111.113t), //

134 INDEX

is.pharma(in.list),77 is.tox21(in.list),77	tissue_scale, 122 ToxCast2015subset, 123
is.toxcast(in.list),77	
is_in_inclusive, 80	wambaugh2019, 124 wambaugh2019.nhanes, 125
johnson, 81	wambaugh2019.raw, 126 wambaugh2019.seem3, 128
kidney_mass_children, 81	well_param, 128 Wetmore.data, 129
liver_mass_children, 82	Wetmore2012, 130
load_sipes2017, <u>82</u>	wfl, 130
<pre>lump_tissues, 83</pre>	,
lung_mass_children, 84	
mcnally_dt, 85	
monte_carlo, 86	
nhanes_mec_svy, 88	
Obach2008, 89	
onlyp, 90	
pancreas_mass_children,90	
parameterize_1comp, 91	
parameterize_3comp, 92	
parameterize_pbtk, 94	
parameterize_schmitt,96	
parameterize_stimitt, 90 parameterize_steadystate, 98	
pc.data, 99	
pharma, 100	
physiology.data, 101	
predict_partitioning_schmitt, 102	
r_left_censored_norm, 104	
rfun, 103	
Sipes2017 (sipes2017), 104	
sipes2017, 104	
sipes2017.table, 105	
skeletal_muscle_mass, 107	
skeletal_muscle_mass_children, 107, 107	
skin_mass_bosgra, 108	
solve_1comp, 108	
solve_3comp, 111	
solve_pbtk, 114	
spleen_mass_children, 117	
spline_heightweight, 118	
spline_hematocrit, 119	
spline_serumcreat, 119	
Tables.Rdata.stamp, 120	
tc.dt.sub (ToxCast2015subset), 123	
tissue.data, 120	
tissue masses flows 121	