# Package 'GeoTox'

November 15, 2024

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Version 0.2.0
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```

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calculate\_response Calculate response

# Description

Calculate mixture response for GeoTox population data

# Usage

```
calculate_response(x, ...)
```

#### **Arguments**

- x GeoTox object
- ... additional arguments passed to other functions. See details.

#### **Details**

Additional parameters include time, BW, and scaling for calc\_internal\_dose, and max\_mult for calc\_concentration\_response.

#### Value

The same object with additional fields added or updated

#### See Also

calc\_internal\_dose, calc\_invitro\_concentration, calc\_concentration\_response

#### **Examples**

```
# Use a subset of the package data for demonstration purposes
set.seed(2357)
n <- 10 # Population size
m <- 5 # Number of regions
idx <- if (m < 100) sample(1:100, m) else 1:100
# Create GeoTox object and populate required fields
geoTox <- GeoTox() |>
  # Simulate populations for each region
  simulate_population(age = split(geo_tox_data$age, ~FIPS)[idx],
                      obesity = geo_tox_data$obesity[idx, ],
                      exposure = split(geo_tox_data$exposure, ~FIPS)[idx],
                      simulated_css = geo_tox_data$simulated_css,
                      n = n) >
  # Estimated Hill parameters
  set_hill_params(geo_tox_data$dose_response |>
                    fit_hill(assay = "endp", chem = "casn") |>
                    dplyr::filter(!tp.sd.imputed, !logAC50.sd.imputed))
# Response computations can now be done
geoTox <- geoTox |> calculate_response()
```

calc\_concentration\_response

Calculate the mixture response from one of three different approaches: IA, GCA, or Hazard Quotient

### **Description**

Calculate the combined response of multiple chemicals. It calculates the generalized concentration addition response, the independent action response, and a hazard quotient

# Usage

```
calc_concentration_response(
   C_invitro,
   hill_params,
   max_mult = 1.5,
   fixed = FALSE
)
```

# **Arguments**

#### Value

list of data frames

# **Examples**

```
calc_independent_action
```

**Independent Action** 

#### **Description**

Calculate independent action response for a set of chemicals with Hill concentration-response curves.

#### Usage

```
calc_independent_action(conc, max, AC50, Emax, n = 1)
```

#### **Arguments**

conc	concentrations in regular space
max	maximal (asymptotic) responses
AC50	concentrations of half-maximal response
Emax	maximum mixture response
n	Hill coefficients (slopes)

#### **Details**

The concentration is computed as:

$$IA = E_{max} \times \left(1 - \prod_i \left(1 - \frac{x_i}{E_{max}}\right)\right),$$

where  $x_i = hill\_val(conc_i, max_i, AC50_i, n_i)$  is the Hill model response function for each chemical.

# Value

response value

# See Also

hill\_val

```
n_chem <- 5
conc <- 10^sample(-1:4, n_chem, replace = TRUE)
max <- 80 * runif(n_chem)
AC50 <- 10^(5 * runif(n_chem) - 1)
Emax <- 100
calc_independent_action(conc, max, AC50, Emax)</pre>
```

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calc\_internal\_dose

Calculate internal chemical dose

# **Description**

Estimate the internal dose from inhalation of a chemical given inhalation rate, time, and body weight

# Usage

```
calc_internal_dose(C_ext, IR, time = 1, BW = 1, scaling = 1)
```

# **Arguments**

C_ext	ambient chemical concentration in $\frac{mg}{m^3}$
IR	inhalation rate in $\frac{m^3}{day}$
time	total time in days
BW	body weight in $kg$
scaling	scaling factor encompassing any required unit adjustments

#### **Details**

Input C\_ext must be a matrix or list of matrices. Input IR must be an atomic vector or list of atomic vectors. The time, BW and scaling arguments are scalars.

The internal dose is calculated as:

$$D_{int} = \frac{C_{ext} \times IR \times time}{BW} \times scaling$$

### Value

list of matrices containing internal chemical doses in  $\frac{mg}{kg}$ 

```
# Single population
C_ext <- matrix(1:15, ncol = 3)
IR <- 1:5
calc_internal_dose(C_ext, IR)

# Multiple populations
C_ext <- list(
    "a" = matrix(1:15 / 10, ncol = 3),
    "b" = matrix(1:8, ncol = 2)
)
IR <- list(1:5, 1:4 / 2)
calc_internal_dose(C_ext, IR)</pre>
```

calc\_invitro\_concentration

Calculate in vitro concentration

# **Description**

Estimate the *in vitro* equivalent plasma concentration given internal chemical dose and steady-state plasma concentration.

# Usage

```
calc_invitro_concentration(D_int, C_ss = NULL)
```

# **Arguments**

D_int	internal chemical dose in $\frac{mg}{kg}$
C_ss	steady-state plasma concentration in $\frac{\mu M}{mg/kg}$

#### **Details**

Input D\_int must be a matrix or list of matrices. Input C\_ss must be a numeric atomic vector or matrix, or a list of those types.

The in vitro equivalent plasma concentration is calculated as:

$$C_{plasma} = C_{ss} \times D_{int}$$

#### Value

list of matrices containing concentrations in  $\mu M$ 

```
# Single population
D_int <- matrix(1:15, ncol = 3)
C_ss <- 1:5
calc_invitro_concentration(D_int, C_ss)

# Multiple populations
D_int <- list(
    "a" = matrix(1:15 / 10, ncol = 3),
    "b" = matrix(1:8, ncol = 2)
)
C_ss <- list(1:5, 1:4 / 2)
calc_invitro_concentration(D_int, C_ss)</pre>
```

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compute\_sensitivity Compute response sensitivity to parameter variation.

#### **Description**

Compute response sensitivity to parameter variation.

### Usage

```
compute_sensitivity(
    x,
    vary = c("age", "obesity", "css_params", "fit_params", "C_ext"),
    max_mult = NULL
)
```

### **Arguments**

```
x GeoTox object.
vary which parameter to vary.
max_mult input for calc_concentration_response step.
```

#### Value

output from calc\_concentration\_response

```
# Use a subset of the package data for demonstration purposes
set.seed(2357)
n <- 10 # Population size
m <- 5 # Number of regions
idx <- if (m < 100) sample(1:100, m) else 1:100
# Create GeoTox object and populate required fields
geoTox <- GeoTox() |>
  # Simulate populations for each region
  simulate_population(age = split(geo_tox_data$age, ~FIPS)[idx],
                      obesity = geo_tox_data$obesity[idx, ],
                      exposure = split(geo_tox_data$exposure, ~FIPS)[idx],
                      simulated_css = geo_tox_data$simulated_css,
                      n = n) >
  # Estimated Hill parameters
  set_hill_params(geo_tox_data$dose_response |>
                    fit_hill(assay = "endp", chem = "casn") |>
                    dplyr::filter(!tp.sd.imputed, !logAC50.sd.imputed))
# Sensitivity computations can now be done
age_resp <- geoTox |> compute_sensitivity()
obesity_resp <- geoTox |> compute_sensitivity(vary = "obesity")
```

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fit\_hill

Fit 2- or 3-parameter Hill model

# **Description**

Fit 2- or 3-parameter Hill model

# Usage

```
fit_hill(
    x,
    conc = "logc",
    resp = "resp",
    fixed_slope = TRUE,
    chem = NULL,
    assay = NULL
)
```

# **Arguments**

x data frame of dose response data.

conc column name of base-10 log scaled concentration.

resp column name of response. fixed\_slope if TRUE, slope is fixed at 1.

chem (optional) column name of chemical identifiers. assay (optional) column name of assay identifiers.

#### **Details**

Optional chem and assay identifiers can be used to fit multiple chemicals and/or assays. Returned columns tp is the top asymptote and logAC50 is the 50% response concentration. If the computation of the standard deviations of these two parameters fails, then the standard deviation is set equal to the parameter estimate and is indicated by the respective imputed flag being TRUE.

#### Value

data frame of fit parameters.

```
# Multiple assays, multiple chemicals
df <- geo_tox_data$dose_response
fit_hill(df, assay = "endp", chem = "casn")

# Single assay, multiple chemicals
df <- geo_tox_data$dose_response |>
    dplyr::filter(endp == "TOX21_H2AX_HTRF_CHO_Agonist_ratio")
```

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GeoTox

GeoTox S3 object

# **Description**

An S3 object that can be used to help organize the data and results of a GeoTox analysis.

# Usage

```
GeoTox()
## S3 method for class 'GeoTox'
plot(x, type = c("resp", "hill", "exposure", "sensitivity"), ...)
```

# **Arguments**

```
x GeoTox object.type type of plot.... arguments passed to subsequent methods.
```

#### Value

```
a GeoTox S3 object
```

#### See Also

```
plot_resp, plot_hill, plot_exposure, plot_sensitivity
```

```
# Use a subset of the package data for demonstration purposes
set.seed(2357)
n <- 10 # Population size
m <- 5 # Number of regions
idx <- if (m < 100) sample(1:100, m) else 1:100

geoTox <- GeoTox() |>
    # Set region and group boundaries (for plotting)
    set_boundaries(region = geo_tox_data$boundaries$county,
```

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```
group = geo_tox_data$boundaries$state) |>
  # Simulate populations for each region
  simulate_population(age
                                   = split(geo_tox_data$age, ~FIPS)[idx],
                                   = geo_tox_data$obesity[idx, ],
                     obesity
                     exposure = split(geo_tox_data$exposure, ~FIPS)[idx],
                      simulated_css = geo_tox_data$simulated_css,
                                   = n) |>
  # Estimated Hill parameters
  set_hill_params(geo_tox_data$dose_response |>
                    fit_hill(assay = "endp", chem = "casn") |>
                    dplyr::filter(!tp.sd.imputed, !logAC50.sd.imputed)) |>
  # Calculate response
  calculate_response() |>
  # Perform sensitivity analysis
  sensitivity_analysis()
# Print GeoTox object
geoTox
# Plot hill fits
plot(geoTox, type = "hill")
# Plot exposure data
plot(geoTox, type = "exposure", ncol = 5)
# Plot response data
plot(geoTox)
plot(geoTox, assays = "TOX21_H2AX_HTRF_CH0_Agonist_ratio")
# Plot sensitivity data
plot(geoTox, type = "sensitivity")
plot(geoTox, type = "sensitivity", assay = "TOX21_H2AX_HTRF_CHO_Agonist_ratio")
```

geo\_tox\_data

GeoTox Data

#### **Description**

Sample data for use in vignettes and function examples. See the Package Data vignette, vignette("package\_data", package = "GeoTox"), for details on how this data was gathered.

#### Usage

```
geo_tox_data
```

#### **Format**

A list with items:

**exposure** 2019 AirToxScreen exposure concentrations for a subset of chemicals in North Carolina counties.

dose\_response Subset of chemicals curated by ICE cHTS as active within a set of assays.

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age County population estimates for 7/1/2019 in North Carolina.

**obesity** CDC PLACES obesity data for North Carolina counties in 2020.

simulated\_css Simulated steady-state plasma concentrations for various age groups and obesity status combinations.

boundaries County and state boundaries for North Carolina in 2019.

get\_fixed\_age

Get C\_ss Data for Fixed Age

# Description

```
Get C_ss Data for Fixed Age
```

# Usage

```
get_fixed_age(simulated_css, age)
```

# Arguments

age list of atomic vectors containing ages.

# Value

list of matrices containing median C\_ss values.

# **Examples**

get\_fixed\_css

Get Fixed C\_ss Data

# **Description**

Get C\_ss values for use in sensitivity\_analysis and compute\_sensitivity.

# Usage

```
get_fixed_css(simulated_css, age, obesity, C_ss)
```

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# Arguments

```
simulated_css list of pre-generated C_ss data, for details see: vignette("package_data", package = "GeoTox").

age list of atomic vectors containing ages.

obesity list of atomic vectors containing obesity status.

C_ss list of matrices containing C_ss values.
```

#### Value

list of matrices or atomic vectors containing C\_ss values.

# **Examples**

get\_fixed\_obesity

Get C\_ss Data for Fixed Obesity Status

# Description

Get C\_ss Data for Fixed Obesity Status

#### Usage

```
get_fixed_obesity(simulated_css, obesity)
```

# **Arguments**

### Value

list of matrices containing median C\_ss values.

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# **Examples**

 $get\_fixed\_other$ 

Get median C\_ss Values

# **Description**

Get median C\_ss Values

# Usage

```
get_fixed_other(C_ss)
```

# **Arguments**

 $C_s$ 

list of matrices containing C\_ss data

# Value

list of atomic vectors containing median C\_ss values.

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Get C\_ss Data for Fixed C\_ss Generation Parameters

# **Description**

Get C\_ss Data for Fixed C\_ss Generation Parameters

# Usage

```
get_fixed_params(simulated_css, age)
```

# Arguments

```
simulated_css list of pre-generated C_ss data, for details see: vignette("package_data",
```

package = "GeoTox").

age list of atomic vectors containing ages.

#### Value

list of matrices containing C\_ss values.

# **Examples**

```
\label{eq:get_fixed_params} $$ \gcd_{css} = \gcd_{tox_{data}} simulated_{css}, $$ age = list(c(25, 35, 55), c(15, 60))) $$
```

hill\_conc

Hill model concentration

# Description

Calculate the concentration in regular space for a given response value.

# Usage

```
hill_conc(resp, max, AC50, n)
```

#### **Arguments**

resp	response value
1 656	response varae

max maximal (asymptotic) response

AC50 concentration of half-maximal response

n Hill coefficient (slope)

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# **Details**

This is a regular space version of tcpl::tcplHillConc().

The concentration is computed as:

$$conc = AC50*(\frac{max}{resp} - 1)^{-1/n}$$

#### Value

concentration in regular space

# See Also

hill\_val

# **Examples**

```
\label{eq:hill_conc} \begin{array}{lllll} hill\_conc(c(0.2,\ 0.5,\ 0.75),\ 1,\ 0.01,\ 1) \\ hill\_conc(c(0.2,\ 0.5,\ 0.9),\ 1,\ c(0.1,\ 0.01,\ 0.001),\ 2) \end{array}
```

hill\_val

Hill model response

# **Description**

Calculate the response for a given concentration in regular space.

# Usage

```
hill_val(conc, max, AC50, n)
```

# **Arguments**

conc concentration in regular space max maximal (asymptotic) response

AC50 concentration of half-maximal response

n Hill coefficient (slope)

#### **Details**

This is a regular space version of tcpl::tcplHillVal().

The Hill model is defined as:

$$resp = \frac{max}{1 + (\frac{AC50}{conc})^n}$$

# Value

response value

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#### See Also

```
hill_conc
```

#### **Examples**

```
hill_val(c(0.0025, 0.01, 0.03), 1, 0.01, 1)
hill_val(c(0.05, 0.01, 0.003), 1, c(0.1, 0.01, 0.001), 2)
```

plot\_exposure

Plot exposure data.

#### **Description**

Plot exposure data.

### Usage

```
plot_exposure(
  exposure,
  region_boundary,
  group_boundary = NULL,
  chem_label = "chnm",
  ncol = 2
)
```

# Arguments

```
exposure list of exposure data named by region label.

region_boundary

"sf" data.frame mapping features to a "geometry" column. Used to color regions.

group_boundary (optional) "sf" data.frame containing a "geometry" column. Used to draw outlines.

chem_label label for facet_wrap.

ncol number of columns to wrap.
```

# Value

ggplot2 object.

```
# Load package data
exposure <- split(geo_tox_data$exposure, ~FIPS)
region_boundary <- geo_tox_data$boundaries$county
group_boundary <- geo_tox_data$boundaries$state
# Plot county exposure data</pre>
```

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plot\_hill

Plot Hill equation fits.

#### **Description**

Plot Hill equation fits.

# Usage

```
plot_hill(hill_params, xlim = c(-1, 4))
```

#### **Arguments**

```
hill_params output from fit_hill.

xlim log-10 scaled concentration limits.
```

#### Value

ggplot2 object.

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plot\_resp

Plot response data

#### **Description**

Plot response data

# Usage

```
plot_resp(
   df,
   region_boundary,
   group_boundary = NULL,
   assay_quantiles = c(Median = 0.5),
   summary_quantiles = c(`10th percentile` = 0.1)
)
```

### **Arguments**

```
df output from resp_quantiles.

region_boundary

"sf" data.frame mapping features to a "geometry" column. Used to color map regions.

group_boundary

"sf" data.frame containing a "geometry" column. Used to draw outlines around groups of regions.

assay_quantiles

named numeric vector of assay quantile labels.

summary_quantiles

named numeric vector of summary quantile labels.
```

# Value

ggplot2 object.

```
# Use example boundary data from package
region_boundary <- geo_tox_data$boundaries$county
group_boundary <- geo_tox_data$boundaries$state
n <- nrow(region_boundary)</pre>
```

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```
# Single assay quantile
df <- data.frame(id = region_boundary$FIPS,</pre>
                 metric = "GCA.Eff",
                 assay_quantile = 0.5,
                 value = runif(n)^3
# Default plot
plot_resp(df, region_boundary)
# Add group boundary, a state border in this case
plot_resp(df, region_boundary, group_boundary)
# Change quantile label
plot_resp(df, region_boundary, group_boundary,
          assay_quantiles = c("Q50" = 0.5))
# Multiple assay quantiles
df <- data.frame(id = rep(region_boundary$FIPS, 2),</pre>
                 metric = "GCA.Eff",
                 assay_quantile = rep(c(0.25, 0.75), each = n),
                 value = c(runif(n)^3, runif(n)^3 + 0.15)
plot_resp(df, region_boundary, group_boundary,
          assay_quantiles = c("Q25" = 0.25, "Q75" = 0.75))
# Summary quantiles
df <- data.frame(id = rep(region_boundary$FIPS, 4),</pre>
                 assay_quantile = rep(rep(c(0.25, 0.75), each = n), 2),
                 summary_quantile = rep(c(0.05, 0.95), each = n * 2),
                 metric = "GCA.Eff",
                 value = c(runif(n)^3, runif(n)^3 + 0.15,
                           runif(n)^3 + 0.7, runif(n)^3 + 0.85)
plot_resp(df, region_boundary, group_boundary,
          assay_quantiles = c("A_Q25" = 0.25, "A_Q75" = 0.75),
          summary_quantiles = c("S_Q05" = 0.05, "S_Q95" = 0.95))
```

plot\_sensitivity

Plot results of sensitivity analysis.

# Description

Plot results of sensitivity analysis.

# Usage

```
plot_sensitivity(
    x,
    metric = "GCA.Eff",
    assay = NULL,
    y = "",
    xlab = metric,
    ylab = ""
)
```

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# Arguments

```
x GeoTox object.

metric metric to plot. Valid choices are "GCA.Eff", "IA.Eff", "GCA.HQ.10", and "IA.HQ.10".

assay which assay to plot, if multiple exist.

y y value or text for bottom of ridge plot.

xlab x-axis label.

ylab y-axis label.
```

#### Value

ggplot2 object.

# Examples

resp\_quantiles

Get response quantiles

# **Description**

Get response quantiles

# Usage

```
resp_quantiles(
  resp,
  metric = c("GCA.Eff", "IA.Eff", "GCA.HQ.10", "IA.HQ.10"),
  assays = NULL,
  assay_summary = FALSE,
```

resp\_quantiles

```
assay_quantiles = c(Median = 0.5),
summary_quantiles = c(`10th percentile` = 0.1)
)
```

#### **Arguments**

```
resp calculated mixture response output from calc_concentration_response.

metric response metric, one of "GCA.Eff", "IA.Eff", "GCA.HQ.10" or "IA.HQ.10".

assays assays to summarize. If NULL and multiple assays exist, then the first assay is used.

assay_summary boolean indicating whether to summarize across assays.

assay_quantiles numeric vector of assay quantiles.

summary_quantiles numeric vector of quantiles to compute across all assay quantiles.
```

#### **Details**

The columns of the returned data frame will vary based on the inputs. If assays is specified and assay\_summary is FALSE, then the resulting data frame will have an assay column. If assay\_summary is TRUE, then the data frame will have an summary\_quantile column.

#### Value

data frame with computed response quantiles.

```
# Dummy response data
resp <- list(</pre>
 "r1" = data.frame(assay = c("a1", "a1", "a2", "a2"),
                    sample = c(1, 2, 1, 2),
                    GCA.Eff = c(1, 2, 3, 4),
                    IA.Eff = c(5, 6, 7, 8),
                    "GCA.HQ.10" = c(9, 10, 11, 12),
                    "IA.HQ.10" = c(13, 14, 15, 16))
# Summarize single assay
resp_quantiles(resp)
# Specify assay
resp_quantiles(resp, assays = "a1")
# Specify quantiles
resp_quantiles(resp, assays = "a1", assay_quantiles = c(0.25, 0.75))
# Specify metric
resp_quantiles(resp, assays = "a1", metric = "IA.HQ.10")
# Summarize across assays
resp_quantiles(resp, assay_summary = TRUE)
# Specify quantiles
suppressWarnings(
```

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sample\_Css

Sample from pre-generated C\_ss data

# Description

Sample from pre-generated C\_ss data

# Usage

```
sample_Css(simulated_css, age, obesity)
```

#### **Arguments**

# Value

list of matrices containing C\_ss values. Columns are sorted to have consistent order across functions.

24 sensitivity\_analysis

#### **Description**

Perform sensitivity analysis

### Usage

```
sensitivity_analysis(x, max_mult = list(NULL, NULL, NULL, 1.2, NULL))
```

#### **Arguments**

```
x GeoTox object.
```

max\_mult numeric list of length 5 for each step of the sensitivity analysis.

# **Details**

This wrapper function will sequentially call the compute\_sensitivity function with inputs age, obesity, css\_params, fit\_params, and C\_ext. The results will be returned as a named list and stored in the sensitivity field of the input GeoTox object.

Values of NULL in the max\_mult input will use the default value stored in the GeoTox object (x\*par\*resp\*max\_mult). When a GeoTox object is created this is initialized at 1.5, but can be changed via the calculate response function or directly in the object.

#### Value

The same GeoTox object with added sensitivity field.

# See Also

```
compute_sensitivity
```

set\_boundaries 25

set\_boundaries

Set GeoTox boundaries

# **Description**

Set GeoTox boundaries

# Usage

```
set_boundaries(x, region = NULL, group = NULL)
```

# Arguments

x GeoTox object.

region "sf" data.frame mapping features to a "geometry" column. Used when coloring

map regions.

group "sf" data.frame containing a "geometry" column. Used to draw outlines around

groups of regions.

#### Value

same GeoTox object with boundaries set.

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set\_hill\_params

Set Hill parameters for a GeoTox object.

# **Description**

Set Hill parameters for a GeoTox object.

# Usage

```
set_hill_params(x, hill_params)
```

# **Arguments**

```
x GeoTox object.
hill_params output of fit_hill.
```

#### Value

same GeoTox object with Hill parameters set.

# **Examples**

```
hill_params <- geo_tox_data$dose_response |>
  fit_hill(chem = "casn", assay = "endp") |>
  dplyr::filter(!tp.sd.imputed, !logAC50.sd.imputed)

geoTox <- GeoTox() |>
  set_hill_params(hill_params)
```

simulate\_age

Simulate ages

# **Description**

Simulate ages

# Usage

```
simulate_age(x, n = 1000)
```

#### Arguments

x data frame or list of data frames containing population data for age groups. Each

data frame must contain columns "AGEGRP" and "TOT\_POP".

n simulated sample size.

simulate\_exposure 27

#### **Details**

Each data frame must contain 19 rows. The first row represents the total population of all age groups while the next 18 rows represent age groups from 0 to 89 in increments of 5 years.

#### Value

List of arrays containing simulated ages.

#### **Examples**

```
# Single data frame
x <- data.frame(AGEGRP = 0:18, TOT_POP = 0)
# populate only age range 40-44, set population total of all ages
x$TOT_POP[c(1, 10)] <- 100
simulate_age(x, 5)

# List of 2 data frames
y <- data.frame(AGEGRP = 0:18, TOT_POP = 0)
# populate age ranges 5-9 and 50-54
y$TOT_POP[c(3, 12)] <- 10
# set population total for all age groups
y$TOT_POP[1] <- sum(y$TOT_POP)
simulate_age(list(x = x, y = y), 15)</pre>
```

simulate\_exposure

Simulate external exposure

#### **Description**

Simulate external exposure

# Usage

```
simulate_exposure(
   x,
   expos_mean = "mean",
   expos_sd = "sd",
   expos_label = "casn",
   n = 1000
)
```

#### **Arguments**

```
    x data frame or list of data frames containing exposure data.
    expos_mean column name of mean values.
    expos_sd column name of standard deviations.
    expos_label column name of labeling term, required if x has more than one row.
    n simulated sample size.
```

#### Value

list of matrices containing inhalation rates. Matrix columns are named using the values in the expos\_label column for more than one data frame row. Columns are sorted to have consistent order across functions.

#### **Examples**

simulate\_inhalation\_rate

Simulate inhalation rates

#### **Description**

Simulate inhalation rates

# Usage

```
simulate_inhalation_rate(x, IR_params = NULL)
```

### **Arguments**

x atomic vector or list of atomic vectors containing ages.

IR\_params (optional) data frame with columns "age", "mean" and "sd". See details for more information.

#### **Details**

The age column of the optional IR\_params data frame should be in ascending order and represent the lower value of age groups for the corresponding mean and sd values. When not provided, the default values will come from Table 6.7 of EPA's 2011 Exposure Factors Handbook using the mean of male and female values.

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# Value

List of atomic vectors containing inhalation rates.

#### **Examples**

simulate\_obesity

Simulate obesity status

### Description

Simulate obesity status

# Usage

```
simulate_obesity(
   x,
   obes_prev = "OBESITY_CrudePrev",
   obes_sd = "OBESITY_SD",
   obes_label = "FIPS",
   n = 1000
)
```

# **Arguments**

```
x data frame containing obesity data as a percentage from 0 to 100.

obes_prev column name of prevalence.

obes_sd column name of standard deviation.

obes_label column name of labeling term, required if x has more than one row.

n simulated sample size.
```

30 simulate\_population

### Value

List of arrays containing simulated obesity status.

#### **Examples**

simulate\_population

Simulate population data

### Description

Simulate population data for given input fields

### Usage

```
simulate_population(
    x,
    age = NULL,
    obesity = NULL,
    exposure = NULL,
    simulated_css = NULL,
    ...
)
```

#### **Arguments**

```
x GeoTox object.

age input x to function simulate_age. After simulating ages, the inhalation rate is subsequently calculated using simulate_inhalation_rate.

obesity input x to function simulate_obesity.

exposure input x to function simulate_exposure.

simulated_css input simulated_css to functions sample_Css and get_fixed_css.

... additional arguments passed to other functions. See details.
```

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# **Details**

Additional parameters include n for sample size, IR\_params for simulate\_inhalation\_rate, obes\_prev, obes\_sd, and obes\_label for simulate\_obesity, and expos\_mean, expos\_sd, and expos\_label for simulate\_exposure.

#### Value

The same object with simulated fields added.

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