

Package: SimuRg (via r-universe)

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Title Building, Fitting and Evaluating PK/PD Modeles

Version 0.2.0

Description Provides a unified workflow for building, fitting using external engines, and evaluating ordinary differential equation (ODE)-based pharmacokinetic/pharmacodynamic (PK/PD) models. Supports generation of estimation scenarios and control files for external engines (e.g., 'Monolix'), simulation of models using 'rxode2', and creation of goodness-of-fit diagnostics. Includes tools for covariate modeling, virtual population design, and local and global sensitivity analyses.

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Contents

data_pbc	2
ds_covval	3
gfo4cov	4
parest	5
sg_converter	6
sg_covsens_sim	7
sg_covsens_vis	12
sg_dummy	17
sg_fit	26
sg_globalsens_sim	32
sg_globalsens_vis	35
sg_gof_obpr	37
sg_gof_par_cov	39
sg_gof_par_dist	40
sg_gof_res	42
sg_gof_res_dist	44
sg_gof_tp	46
sg_localsens_sim	47
sg_localsens_vis	50
sg_modbuild	53
sg_modcomp	57
sg_multistart	57
sg_parsum	61
sg_predist_sim	62
sg_sim	63
sg_sim_tp	69
sg_translator	71
sg_vpc_sim	73
sg_vpc_vis	75
sg_vpop_est	78
warfarin	81
Index	83

data_pbc

Primary Biliary Cirrhosis (PBC) dataset

Description

Clinical trial dataset with Primary Biliary Cirrhosis patient data including survival outcomes and covariates.

Usage

data_pbc

Format

data_pbc:

A data frame with 280 rows and 15 columns:

id identifier of the individual
years time in years (survival time)
status2 survival status (0 = censored, 1 = event)
drug treatment group ("D-penicil" or "placebo")
age age of the individual in years
sex sex of the individual ("male" or "female")
ascites presence of ascites ("Yes" or "No")
hepatomegaly presence of hepatomegaly ("Yes" or "No")
spiders presence of spider angiomas ("Yes" or "No")
edema edema status ("No edema", "edema no diuretics", or "edema despite diuretics")
serBilir serum bilirubin level (mg/dl)
serChol serum cholesterol level (mg/dl)
albumin albumin level (g/dl)
platelets platelet count (per cubic mm/1000)

ds_covval

Warfarin covariate values dataset

Description

Individual covariate values for 100 patients used in the warfarin covariate sensitivity simulation, including demographic and pharmacogenomic variables.

Usage

ds_covval

Format

ds_covval:

A data frame with 100 rows and 9 columns:

ID individual patient identifier
AGE age in years
WEIGHT body weight in kg
BMI body mass index (kg/m²)
LG_WEIGHT log-transformed body weight (log10 scale), used as continuous covariate in the PK model
LG_AGE log-transformed age (log10 scale), used as continuous covariate in the PK model
SEX sex of the individual (0 = female, 1 = male)
CYP2C9 CYP2C9 genotype encoded as integer: 0 = **1/*1* (reference), 1 = **1/*2*, 2 = **1/*3*, 3 = **2/*2*, 4 = **2/*3*, 5 = **3/*3*
VKORC1 VKORC1 genotype ("GG", "AG", or "AA")

Source

Derived from `data_fin_i.csv` — individual covariate dataset for the warfarin PopPK covariate sensitivity analysis.

gfo4cov

Warfarin PopPK model fitting output (4-covariate model)

Description

A list containing the full output of a warfarin population pharmacokinetic model fit with four covariates (SEX, LG_WEIGHT, CYP2C9, VKORC1), including parameter estimates, model diagnostics, individual predictions, and covariate data.

Usage

gfo4cov

Format

gfo4cov:

A named list with 12 elements:

PROJNAME character string identifying the project run ("run_4cov")

SUMTAB data frame with 15 rows and 11 columns containing the parameter summary table.

Columns: PAR (parameter name), VALUE (estimate), TYPE (category: "Typical values", "Covariate effects", "Random effects", or "Residual error model"), EST (estimation status), SE (standard error), RSE (relative SE, \ LCI and UCI (95\ ETAshrinkage_var, ETAshrinkage_sd, EPSshrinkage_sd (shrinkage metrics).

SDTAB data frame with 1600 rows and 11 columns — the standard table of observations and model predictions. Columns: ID, TIME, DV (observed), DVID, PRED (population prediction), IPRED (individual prediction), RES (residual), IRES (individual residual), WRES (weighted residual), IWRES (individual weighted residual), MDV.

PATAB data frame with 100 rows and 7 columns — individual parameter table. Columns: ID, eta_ka, eta_Vd, eta_CL (individual ETA random effects), ka, Vd, CL (individual post-hoc PK parameter estimates).

COVMAT 15×15 numeric matrix — covariance matrix of the population parameter estimates.

CORRMAT 15×15 numeric matrix — correlation matrix of the population parameter estimates.

OFV data frame with 1 row and 4 columns — model fit criteria: LL (log-likelihood), AIC, BIC, BICc.

OMEGAMAT 3×3 numeric matrix — OMEGA variance-covariance matrix of the inter-individual random effects (eta_ka, eta_Vd, eta_CL).

SIGMAMAT 1×1 numeric matrix — SIGMA residual error variance matrix.

EVTAB data frame with 100 rows and 6 columns — dosing/event table. Columns: ID, TIME, EVID, AMT, ADM, CMT.

COTAB data frame with 100 rows and 6 columns — continuous covariate table. Columns: ID, AGE, WEIGHT, BMI, LG_WEIGHT (log10-transformed weight), LG_AGE (log10-transformed age).

CATAB data frame with 100 rows and 4 columns — categorical covariate table. Columns: ID, SEX (0 = female, 1 = male), CYP2C9 (genotype integer code, 0–5), VKORC1 (genotype: "GG", "AG", or "AA").

Source

Generated from the warfarin PopPK model fitting run "run_4cov" used in the covariate sensitivity analysis workflow.

parest	<i>Warfarin population PK parameter estimates</i>
--------	---

Description

Final parameter estimates from a warfarin population pharmacokinetic model, including typical values, covariate effects, random effects, and residual error parameters with associated uncertainty measures.

Usage

```
parest
```

Format

parest:

A data frame with 15 rows and 12 columns:

parameter parameter name as used in the model

value final parameter estimate

TYPE parameter category: "Typical values", "Covariate effects", "Random effects", or "Residual error model"

EST estimation status ("ESTIMATED")

SE standard error of the estimate

RSE relative standard error (%)

LCI lower bound of the 95 % confidence interval

UCI upper bound of the 95 % confidence interval

ETAshrinkage_var ETA shrinkage on the variance scale (%); NA for non-random-effect parameters

ETAshrinkage_sd ETA shrinkage on the SD scale (%); NA for non-random-effect parameters

EPSshrinkage_sd EPS shrinkage on the SD scale (%); NA for non-residual-error parameters

Source

Derived from par_fin_i.csv — warfarin PopPK model estimation output.

 sg_converter

 Convert Monolix project output to R objects

Description

Reads and parses Monolix project output files (.mlxtran and associated data files) into a structured list of SimuRg objects including parameter estimates, individual predictions, residuals, and diagnostic information.

Usage

```
sg_converter(folder_path, proj_name, save_file = FALSE)
```

Arguments

folder_path	Character string. Path to the directory containing Monolix project files.
proj_name	Character string. Name of the Monolix project (without file extension).
save_file	Logical. If TRUE, saves GCO and GFO JSON files to folder_path with names <proj_name>_GCO.json and <proj_name>_GFO.json, and also saves two RData files: <proj_name>_GCO.RData (object gco) and <proj_name>_GFO.RData (object gfo).

Details

This function serves as a bridge between Monolix output and R by parsing the .mlxtran file and associated data files to create a comprehensive R object containing all relevant model outputs. This facilitates further analysis, visualization, and reporting in R.

The function automatically detects and imports various components of Monolix output including population parameters, individual parameters, covariates, and diagnostic metrics.

If save_file = TRUE, the function additionally writes GCO and GFO JSON files and .RData files to folder_path.

Value

Returns a list with the following components:

- GFO: SimuRg generalized fit object output object with:
 - SDTAB: A tibble containing data used for fitting
 - SUMTAB: A tibble with parameter summary statistics containing
 - SIGMAT: Residual variability matrix
 - OMEGAMAT: Inter-individual variability matrix
 - OCCMAT: Inter-occasion variability matrix (NA if not present)
 - EVTAB: A tibble with event information
 - PATAB: A tibble with individual parameter estimates
 - COTAB: A tibble with continuous covariates

- CATAB: A tibble with categorical covariates
- REGTAB: Regression parameters (empty data.frame if not present)
- OFV: A tibble with objective function values
- COVMAT: Variance-covariance matrix of parameter estimates
- CORRMAT: Correlation matrix of parameter estimates
- OPTIONS: Model options (NULL if not present)
- PROJNAME: Project name
- GCO: SimuRg generalized control object parsed from the mlxtran project with:
 - headers: List of dataset column descriptors (name, use, type)
 - data: Path to source data file
 - model: Path to model file
 - task_opt: Task options placeholder (empty object)
 - covs: Covariate names detected
 - project_name: Project name
 - theta: List of population parameter definitions (NAME, INIT, EST, TRANS)
 - ruv: Residual error model definition (YNAME, DVID, TRANS, PRED, ERR, INIT, EST)
 - re: Between-subject variability matrices (init, est)
 - occ: Between-occasion variability matrices (init, est)
 - modelText: Text content of the model file

Examples

```
library(stringr)
# Convert Monolix project results
test_folder <- system.file("extdata", "Monolix_objects", package = "SimuRg")
if (substr(test_folder, nchar(test_folder), nchar(test_folder)) != "/")
  test_folder <- str_c(test_folder, "/")
proj_name <- "proj-solo"
result <- sg_converter(folder_path = test_folder, proj_name = proj_name)
# save(results, file = "./models/simurg_object/Warfarin_PK.RData")
# Access individual predictions
head(result$GFO$SDTAB)

# View parameter estimates
print(result$GFO$SUMTAB)

# Check objective function value
print(result$GFO$OFV)
```

Description

Evaluate the impact of continuous and categorical covariates on model parameters and exposure metrics. For each covariate the function perturbs its value to selected quantiles (continuous) or observed categories (categorical) while holding the remaining covariates at their reference values, simulates from the pharmacometric model under parameter uncertainty, and summarises the resulting percent change relative to the reference.

Usage

```
sg_covsens_sim(
  fpath_i = NULL,
  ds_parest = NULL,
  ds_covs = NULL,
  model,
  stimes,
  et,
  est_covmat,
  npop = 5,
  cont_cov_l,
  cat_cov_l,
  quantiles = c(0.1, 0.9),
  outputs = "Cc",
  aggr = c("min", "max", "mean")
)
```

Arguments

fpath_i	String or sg-fit object. If the string is given, the path to .Rdata or .json file with sg-fit object is expected.
ds_parest	Data.frame. Parameter estimates table with columns parameter and value. Required when fpath_i is NULL; must be provided together with ds_covs. Default is NULL.
ds_covs	data.frame. Subject-level covariate dataset (one row per subject) containing both continuous and categorical covariate columns. Required when fpath_i is NULL; must be provided together with ds_parest. Default is NULL.
model	A model object passed to sg_sim().
stimes	numeric vector. Sampling time points for steady-state simulation (e.g. generated by a cycling function over dosing intervals).
et	data.frame. Event (dosing) table. Must contain at least columns id, time, amt, evid and Regimen. Additional columns such as ii, addl, dur and Dose are used when present.
est_covmat	Data.frame. Parameter estimation covariance matrix. The first column (X1) must list parameter names; remaining columns (named identically) form the symmetric variance-covariance matrix.
npop	integer. Number of population-level simulation replicates drawn from the parameter uncertainty distribution. Default is 5.

cont_cov_1	<p>A named list. Each element defines one continuous covariate and must itself be a list with components:</p> <p>NAME Character. Column name of the (transformed) covariate in the dataset (e.g. "LG_AGE").</p> <p>UTNAME Character or NULL. Column name of the untransformed (back-transformed) covariate (e.g. "AGE"). If NULL or NA, defaults to NAME.</p> <p>REF Character or numeric. Reference value for the covariate. Use "median" to derive from data, or a numeric value.</p> <p>NICENAME Character or NULL. Display label for plots and tables (e.g. "Age, years").</p> <p>par_vec Character vector. Model parameter(s) affected by this covariate (e.g. c("CL")).</p>
cat_cov_1	<p>A named list. Each element defines one categorical covariate and must itself be a list with components:</p> <p>NAME Character. Column name of the covariate (e.g. "SEX").</p> <p>NICENAME Character or NULL. Display label.</p> <p>REF Character or NULL. Reference category value. If NULL, the first factor level (alphabetically) is used.</p> <p>par_vec Character vector. Model parameter(s) affected by this covariate (e.g. c("ka")).</p>
quantiles	A numeric vector of length 2. Lower and upper quantiles of the continuous covariate distribution to test. Default is $c(0.1, 0.9)$.
outputs	character vector. Name(s) of the model output variable(s) to evaluate (e.g. "Cc" or c("Cc", "Effect")). Default is "Cc".
aggr	character vector. Exposure aggregation metric(s) applied over the simulation time grid. Allowed values: "min" (Cmin), "max" (Cmax), "mean" (Cavg). Default is c("min", "max", "mean").

Details

Two data-source modes are supported (mutually exclusive):

- **File mode** — supply `fpath_i` (path to a SimuRg JSON / RData object that contains SUMTAB, CATAB and COTAB).
- **Table mode** — supply both `ds_parest` (parameter estimates) and `ds_covs` (covariate dataset).

The analysis proceeds in two stages for each covariate type (continuous and categorical):

1. **Parameter sensitivity** — the covariate is set to its quantile or category value and the model is evaluated at time zero (no ODE integration) to quantify the direct effect on each parameter.
2. **Exposure sensitivity** — the full time-course is simulated over `stimes` and exposure metrics (`aggr`) are computed.

Uncertainty is propagated by sampling `npop` parameter vectors from a multivariate normal distribution parameterised by the population estimates and the estimation covariance matrix (`est_covmat`).

Value

A named list of three elements:

`$PARSENS` data.frame. Full sensitivity results for model parameters: percent change statistics (mean, median, percentiles) for each covariate–parameter combination, with columns NICEN, VAR, KEY, LAB, Type and summary statistics (mean through P975).

`$SUMPARSENS` data.frame. Compact summary table with columns Parameter, Covariate, Cov. percentile, Cov. value, Mean and 90%CI.

`$EXPSSENS` data.frame. Sensitivity of exposure metrics (Cmin, Cmax, Cavg) to covariates, structured identically to `$PARSENS`.

See Also

[sg_sim](#), [read_smrg_obj](#)

Examples

```
library(dplyr)
library(rxode2)

# --- Covariate definitions ---
cont_cov_l <- list(
  LG_AGE = list(NAME = "LG_AGE", UTNAME = "AGE",
               REF = "median", NICENAME = "Age, years",
               par_vec = c("CL")),
  LG_WEIGHT = list(NAME = "LG_WEIGHT", UTNAME = "WEIGHT",
                  REF = "median", NICENAME = "Weight, kg",
                  par_vec = c("Vd"))
)

cat_cov_l <- list(
  SEX = list(NAME = "SEX", NICENAME = "Sex",
            REF = "0", par_vec = c("ka")),
  CYP2C9 = list(NAME = "CYP2C9", NICENAME = "CYP2C9 genotype",
               REF = NULL, par_vec = c("CL"))
)

# --- Dosing ---
ev_t_input <- tribble(
  ~id, ~time, ~ii, ~amt, ~addl, ~dur, ~evid, ~Regimen, ~Dose,
  1, 0, 336, 10, 21, 0.5, 1, "0.3 mg/kg Q2W", 0.3
)

# --- Model ---
model <- RxODE({
  # Doses in mg
  # Time in hours

  ### Parameter values
  # Typical values
  ka_pop = 0.073;
  Vd_pop = 14.8;
```

```

CL_pop = 0.347;

# Random effects
omega_ka = 0;
omega_Vd = 0;
omega_CL = 0;

# Covariate effect
# Continuous
beta_CL_LG_AGE = 0.49990114;
beta_Vd_LG_WEIGHT = 0.60529433;

# Categorical
beta_CL_CYP2C9_1_2 = -0.339;
beta_CL_CYP2C9_1_3 = -0.574;
beta_CL_CYP2C9_2_2 = -1.079;
beta_CL_CYP2C9_2_3 = -0.745;
beta_CL_CYP2C9_3_3 = -2.13;

beta_ka_SEX_1 = -0.12198035;

# Residual error
Cc_b = 0;

# Transformations
ka_tv = exp(ka_pop);
Vd_tv = exp(Vd_pop);
CL_tv = exp(CL_pop);

CL_multiplier = 1.0; # Default/reference
ka_multiplier = 1.0;

if (SEX == "1") {ka_multiplier = exp(beta_ka_SEX_1)}

if (CYP2C9 == "1") {
  CL_multiplier = exp(beta_CL_CYP2C9_1_2);
} else if (CYP2C9 == "2") {
  CL_multiplier = exp(beta_CL_CYP2C9_1_3);
} else if (CYP2C9 == "3") {
  CL_multiplier = exp(beta_CL_CYP2C9_2_2);
} else if (CYP2C9 == "4") {
  CL_multiplier = exp(beta_CL_CYP2C9_2_3);
} else if (CYP2C9 == "5") {
  CL_multiplier = exp(beta_CL_CYP2C9_3_3);
}

ka = ka_tv*ka_multiplier*exp(omega_ka);
Vd = Vd_tv*exp(beta_Vd_LG_WEIGHT * LG_WEIGHT + omega_Vd); #Vd_tv*exp(omega_Vd);
CL = CL_tv*CL_multiplier*exp(beta_CL_LG_AGE * LG_AGE + omega_CL);

### Explicit functions
Cc = Ac/Vd;

```

```

### Initial conditions
Ad(0) = 0;
Ac(0) = 0;

### Differential equations
d/dt(Ad) = - ka*Ad;
d/dt(Ac) = ka*Ad - CL*Cc;

Cc_ResErr = Cc*(1 + Cc_b);
})

# --- Estimation covariance (mock) ---
pnames      <- parest$parameter
npar        <- length(pnames)
set.seed(1)
m_cov       <- matrix(0.02, npar, npar)
diag(m_cov) <- 0.05 + runif(npar, 0, 0.05)
m_cov       <- (m_cov + t(m_cov)) / 2
est_covmat  <- as_tibble(cbind(X1 = pnames, as.data.frame(m_cov)))
names(est_covmat)[-1] <- pnames

# --- Simulation times (steady-state cycle 10) ---
ss_cycle <- 10
stimes_ss <- c(
  ss_cycle * 4 * 7 * 24 + c(seq(0, 23.5, 0.5), seq(24, 335, 1)),
  ss_cycle * 4 * 7 * 24 + 2 * 7 * 24 + c(seq(0, 23.5, 0.5), seq(24, 335, 1))
)

# --- Run ---
result <- sg_covsens_sim(
  fpath_i=NULL, ds_parest = parest, ds_covs = ds_covval,
  model = model, stimes = stimes_ss, et = ev_t_input,
  est_covmat = est_covmat, npop = 10,
  cont_cov_l = cont_cov_l, cat_cov_l = cat_cov_l,
  quantiles = c(0.2, 0.8), aggr = c("max"),
  outputs = "Cc"
)
print(result[["PARSENS"]])
print(result[["SUMPARSENS"]])
print(result[["EXPSENS"]])

```

Description

Draw a forest-style graphic (points with uncertainty intervals) from the sensitivity tables produced by `sg_covsens_sim`. Each facet row corresponds to a model output or exposure metric (VAR); each point is a covariate scenario (LAB), coloured by covariate type (Type).

Usage

```
sg_covsens_vis(
  covsens_res,
  plot_type = c("PARSENS", "EXPESENS"),
  exclude_vars = NULL,
  ci_quantiles = c("P025", "P975"),
  ci_limits = c(0.8, 1.25),
  ci_band_alpha = 0.2,
  ci_band_col = "firebrick",
  ref_line_col = "grey25",
  palette = MSDcol[c(1, 3, 4, 5, 6, 7)],
  point_size = 2.5,
  errorbar_width = 0.2,
  lab_y = "Mean (95% CI)\nchange from reference",
  cap = NULL
)
```

Arguments

<code>covsens_res</code>	Named list as returned by <code>sg_covsens_sim()</code> . Must contain the element selected by type (PARSENS and/or EXPESENS data.frames with columns LAB, VAR, mean, Type, and the interval columns named by <code>ci_quantiles</code>).
<code>plot_type</code>	Character scalar: "PARSENS" (default) or "EXPESENS".
<code>exclude_vars</code>	Character vector. Contains VAR levels to omit (e.g. "Cc_Cmin"). NULL keeps all rows.
<code>ci_quantiles</code>	Character vector of length 2: names of the lower and upper uncertainty columns in the sensitivity table, in that order. Defaults <code>c("P025", "P975")</code> to match the default percentiles in <code>sg_covsens_sim</code> . Use other names (e.g. <code>c("P05", "P95")</code>) if you changed quantiles in the simulation and the columns exist.
<code>ci_limits</code>	Numeric vector of length 2: lower and upper bounds of the shaded acceptance band and of the dotted horizontal guides. Default <code>c(0.8, 1.25)</code> is a common bioequivalence-style window on the ratio scale.
<code>ci_band_alpha</code>	Numeric in $[0, 1]$: transparency of the shaded band. Default 0.2.
<code>ci_band_col</code>	Color for the band fill and dotted limit lines. Default "firebrick".
<code>ref_line_col</code>	Color for the dashed horizontal line at $y = 1$ (no change from reference). Default "grey25".
<code>palette</code>	Colors for the Type scale (continuous vs categorical covariates, etc.). Recycled if there are more levels than colors. Default <code>MSDcol[c(1, 3, 4, 5, 6, 7)]</code> .
<code>point_size</code>	Point size for <code>geom_point</code> . Default 2.5.

errorbar_width	Numeric. Width argument for geom_errorbar. Default 0.2.
lab_y	Axis label for the numeric scale (this becomes the horizontal axis after coord_flip()). Default mentions a 95\ if you use different ci_quantiles.
cap	Optional figure caption, passed to ggplot2::labs(caption = ...) (e.g. text describing reference covariate values). Default NULL.

Details

Two views are available, matching the named elements of the simulation output:

- PARSENS — sensitivity of individual parameters (simulation at time zero, no ODE time course).
- EXPSENS — sensitivity of exposure summaries (e.g. Cmin, Cmax, Cavg) after full simulation over stimes in sg_covsens_sim.

Values on the y-axis are ratios relative to the reference scenario: 1 means no change. In sg_covsens_sim, percent change relative to reference is transformed to this scale before tabulation. The shaded region between ci_limits highlights a target interval; points whose intervals lie largely inside can be read as scenarios consistent with that criterion, subject to study-specific rules.

The plot layers are drawn in order: reference band, error bars, points, reference line at 1, dotted lines at ci_limits, then faceting by VAR and flipped coordinates so labels read along the vertical axis.

Value

A ggplot2 object (inactive until printed or saved). You can add further layers or themes with the usual **ggplot2** API.

See Also

[sg_covsens_sim](#)

Examples

```
library(tibble)
library(dplyr)
library(rxode2)
# Typical workflow: run the simulation (see examples in ?sg_covsens_sim),
# then visualise parameter and exposure sensitivity.

cont_cov_l <- list(
  LG_AGE = list(NAME = "LG_AGE", UTNAME = "AGE",
               REF = "median", NICENAME = "Age, years",
               par_vec = c("CL")),
  LG_WEIGHT = list(NAME = "LG_WEIGHT", UTNAME = "WEIGHT",
                  REF = "median", NICENAME = "Weight, kg",
                  par_vec = c("Vd"))
)

cat_cov_l <- list(
  SEX = list(NAME = "SEX", NICENAME = "Sex",
```

```

        REF = "0", par_vec = c("ka")),
    CYP2C9 = list(NAME = "CYP2C9", NICENAME = "CYP2C9 genotype",
        REF = NULL, par_vec = c("CL"))
)

# --- Dosing ---
ev_t_input <- tribble(
  ~id, ~time, ~ii, ~amt, ~addl, ~dur, ~evid, ~Regimen, ~Dose,
  1, 0, 336, 10, 21, 0.5, 1, "0.3 mg/kg Q2W", 0.3
)

# --- Model ---
model <- RxODE({
  # Doses in mg
  # Time in hours

  ### Parameter values
  # Typical values
  ka_pop = 0.073;
  Vd_pop = 14.8;
  CL_pop = 0.347;

  # Random effects
  omega_ka = 0;
  omega_Vd = 0;
  omega_CL = 0;

  # Covariate effect
  # Continuous
  beta_CL_LG_AGE = 0.49990114;
  beta_Vd_LG_WEIGHT = 0.60529433;

  # Categorical
  beta_CL_CYP2C9_1_2 = -0.339;
  beta_CL_CYP2C9_1_3 = -0.574;
  beta_CL_CYP2C9_2_2 = -1.079;
  beta_CL_CYP2C9_2_3 = -0.745;
  beta_CL_CYP2C9_3_3 = -2.13;

  beta_ka_SEX_1 = -0.12198035;

  # Residual error
  Cc_b = 0;

  # Transformations
  ka_tv = exp(ka_pop);
  Vd_tv = exp(Vd_pop);
  CL_tv = exp(CL_pop);

  CL_multiplier = 1.0; # Default/reference
  ka_multiplier = 1.0;

  if (SEX == "1") {ka_multiplier = exp(beta_ka_SEX_1)}

```

```

if (CYP2C9 == "1") {
  CL_multiplier = exp(beta_CL_CYP2C9_1_2);
} else if (CYP2C9 == "2") {
  CL_multiplier = exp(beta_CL_CYP2C9_1_3);
} else if (CYP2C9 == "3") {
  CL_multiplier = exp(beta_CL_CYP2C9_2_2);
} else if (CYP2C9 == "4") {
  CL_multiplier = exp(beta_CL_CYP2C9_2_3);
} else if (CYP2C9 == "5") {
  CL_multiplier = exp(beta_CL_CYP2C9_3_3);
}

ka = ka_tv*ka_multiplier*exp(omega_ka);
Vd = Vd_tv*exp(beta_Vd_LG_WEIGHT * LG_WEIGHT + omega_Vd); #Vd_tv*exp(omega_Vd);
CL = CL_tv*CL_multiplier*exp(beta_CL_LG_AGE * LG_AGE + omega_CL);

### Explicit functions
Cc = Ac/Vd;

### Initial conditions
Ad(0) = 0;
Ac(0) = 0;

### Differential equations
d/dt(Ad) = - ka*Ad;
d/dt(Ac) = ka*Ad - CL*Cc;

Cc_ResErr = Cc*(1 + Cc_b);
})

# --- Estimation covariance (mock) ---
pnames <- parest$parameter
npar <- length(pnames)
set.seed(1)
m_cov <- matrix(0.02, npar, npar)
diag(m_cov) <- 0.05 + runif(npar, 0, 0.05)
m_cov <- (m_cov + t(m_cov)) / 2
est_covmat <- as_tibble(cbind(X1 = pnames, as.data.frame(m_cov)))
names(est_covmat)[-1] <- pnames

# --- Simulation times (steady-state cycle 10) ---
ss_cycle <- 10
stimes_ss <- c(
  ss_cycle * 4 * 7 * 24 + c(seq(0, 23.5, 0.5), seq(24, 335, 1)),
  ss_cycle * 4 * 7 * 24 + 2 * 7 * 24 + c(seq(0, 23.5, 0.5), seq(24, 335, 1))
)

result <- sg_covsens_sim(
  fpath_i = NULL, ds_parest = parest, ds_covs = ds_covval,
  model = model, stimes = stimes_ss, et = ev_t_input,
  est_covmat = est_covmat, npop = 10,
  cont_cov_l = cont_cov_l, cat_cov_l = cat_cov_l,
  quantiles = c(0.1, 0.9), aggr = c("min", "max", "mean"),

```

```

    outputs = "Cc"
  )

  p_par <- sg_covsens_vis(result, plot_type = "PARSENS")
  p_exp <- sg_covsens_vis(result, plot_type = "EXPESENS")
  print(p_par)
  print(p_exp)

  # Alternate interval columns (must exist in the sensitivity tables)
  sg_covsens_vis(result, ci_quantiles = c("P05", "P95"))

  # Drop selected metrics from the exposure panel
  sg_covsens_vis(result, plot_type = "EXPESENS", exclude_vars = c("Cc_Cmin"))

```

sg_dummy

The function to store common parameters description

Description

The function to store common parameters description

Usage

```

sg_dummy(
  ...,
  abreaks = scales::pretty_breaks(7),
  addcov = TRUE,
  addOFV = TRUE,
  addline = TRUE,
  aggr = NULL,
  alpha_i = 0.5,
  atol = 1e-08,
  cap,
  cat_cov_l,
  ci_band_alpha = 0.2,
  ci_band_col = "firebrick",
  ci_limits = c(0.8, 1.25),
  ci_quantiles = c("P025", "P975"),
  cont_cov_l,
  cov_cols,
  cov,
  covint = "locf",
  covsens_res,
  ciLow = 0.025,
  ciUp = 0.975,
  col_i,

```

```
col_lab,  
covs,  
data,  
dens,  
ds_covs,  
ds_i,  
ds_parest = NULL,  
dt_obs_fl = FALSE,  
DVID = 1,  
dv_col = "DV",  
emp_perc = TRUE,  
errorbar_width = 0.2,  
est_covmat,  
et,  
eta_seq = NULL,  
excl_col = NULL,  
exclude_vars = NULL,  
f_scales = "fixed",  
facet_i,  
fill_i = NULL,  
filt = "T",  
fit = FALSE,  
fpath_i,  
free_stat = "free",  
group_i = "VAR",  
headers,  
ncores = 1,  
id_col = NULL,  
indiv = TRUE,  
inits = NULL,  
keep = NULL,  
lab_x,  
lab_y,  
legend_fl = FALSE,  
levels_discrete = 10,  
log_x = FALSE,  
log_y = FALSE,  
log_axes = FALSE,  
lty_i = NULL,  
max_x = NULL,  
max_y = NULL,  
method = "lsoda",  
min_x = NULL,  
min_y = NULL,  
model,  
n_bins = 30,  
n_quantiles = 3,  
n_sim,
```

```
no_leg = FALSE,
npop = 1,
nsub = 1,
occ,
opt_name = "Monolix",
omega,
outputs = NULL,
output = NULL,
path_to_fitter = NULL,
path_to_save_output = NULL,
par_bounds,
par_seq = NULL,
par_type = "Ind",
params = NULL,
piLow = 0.1,
piUp = 0.9,
plot_type = "DIST",
point_size = 2.5,
pred.corr = FALSE,
project_name,
quantiles = c(0.1, 0.9),
re,
ref_line_col = "grey25",
rtol = 1e-06,
run_id = 1,
ruv,
sc_factor = 1,
scale = NULL,
seed = 123,
sigma = NULL,
shp_i = NULL,
smooth = TRUE,
stat_comp = NULL,
stimes = NULL,
task_opt = NULL,
tdist = TRUE,
time_col = "TIME",
theor_perc = TRUE,
theor_percCI = TRUE,
theta = NULL,
thetamat = NULL,
tsld = FALSE,
val_col = "VALUE",
wrap_i = NULL,
wrap_ncol = NULL,
wrap_nrow = NULL
)
```

Arguments

...	Other arguments that will be passed to rxSolve function.
abreaks	A function that generates axis breaks. Default is <code>scales::pretty_breaks(7)</code> .
addcov	Logical. If TRUE, columns with covariate values will be added to the resulting dataset. Default is TRUE.
addOFV	Logical. If TRUE, information about OFV and AIC of the model will be added. Default is TRUE.
addline	Logical. If TRUE, lines connecting observations of individual subjects will be added. Default is TRUE.
aggr	A string that specifies the aggregation type. Set to NULL for no aggregation, ID for aggregation by time, population and ID, NPOP for aggregation by population and time, TIME for aggregation by time. Default is NULL.
alpha_i	Numeric. Transparency level (from 0 to 1) for points/lines. Default is 0.5.
atol	A numeric absolute tolerance used by the ODE solver to determine if a good solution has been achieved. This is also used in the solved linear model to check if prior doses do not add anything to the solution. Default is 1e-8.
cap	A string that specifies the plot caption.
cat_cov_l	A named list. Each element defines one categorical covariate and must itself be a list with components: NAME Character. Column name of the covariate (e.g. "SEX"). NICENAME Character or NULL. Display label. REF Character or NULL. Reference category value. If NULL, the first factor level (alphabetically) is used. par_vec Character vector. Model parameter(s) affected by this covariate (e.g. <code>c("ka")</code>).
ci_band_alpha	Numeric in $[0, 1]$: transparency of the shaded band. Default 0.2.
ci_band_col	Color for the band fill and dotted limit lines. Default "firebrick".
ci_limits	Numeric vector of length 2: lower and upper bounds of the shaded acceptance band and of the dotted horizontal guides. Default <code>c(0.8, 1.25)</code> is a common bioequivalence-style window on the ratio scale.
ci_quantiles	Character vector of length 2: names of the lower and upper uncertainty columns in the sensitivity table, in that order. Defaults <code>c("P025", "P975")</code> to match the default percentiles in <code>sg_covsens_sim</code> . Use other names (e.g. <code>c("P05", "P95")</code>) if you changed quantiles in the simulation and the columns exist.
cont_cov_l	A named list. Each element defines one continuous covariate and must itself be a list with components: NAME Character. Column name of the (transformed) covariate in the dataset (e.g. "LG_AGE"). UTNAME Character or NULL. Column name of the untransformed (back-transformed) covariate (e.g. "AGE"). If NULL or NA, defaults to NAME. REF Character or numeric. Reference value for the covariate. Use "median" to derive from data, or a numeric value.

	NICENAME	Character or NULL. Display label for plots and tables (e.g. "Age, years").
	par_vec	Character vector. Model parameter(s) affected by this covariate (e.g. c("CL")).
cov_cols		A character vector specifying the names of the columns with covariates.
cov		Covariates passed to <code>sg_sim()</code> . Default is NULL.
covint		String. Specifies the interpolation method for time-varying covariates. When solving ODEs it often samples times outside the sampling time specified in event table. When this happens, the time varying covariates are interpolated. Currently this can be: <ul style="list-style-type: none"> • "linear" interpolation, which interpolates the covariate by solving the line between the observed covariates and extrapolating the new covariate value • "locf" last observation carried forward • "NOCB" next observation carried backward. This is the same method that NONMEM uses • "midpoint" last observation carried forward to midpoint; next observation carried backward to midpoint Default is "locf"
covsens_res		Named list as returned by <code>sg_covsens_sim()</code> . Must contain the element selected by <code>type</code> (PARSENS and/or EXPSENS data.frames with columns LAB, VAR, mean, Type, and the interval columns named by <code>ci_quantiles</code>).
ciLow		Numeric. Lower confidence interval bound. Default is 0.025
ciUp		Numeric. Upper confidence interval bound. Default is 0.975
col_i		String. Column name for color
col_lab		String. Label for color legend
covs		A list of covariate structures. Each element should be a list containing covariate relationship definitions with the following structure: <ul style="list-style-type: none"> • PAR - string, name of the parameter to which covariate effect is applied • COVNAME - string, name of the covariate column in the dataset • FUNC - string, functional form of the covariate effect • TRANS - string, transformation applied to covariate • REF - numeric, reference value for categorical covariates • INIT - numeric, initial value for the covariate effect parameter • EST - logical, whether to estimate the covariate effect Can be NULL, if no covariates are used. Default is NULL
data		String. Path to the dataset used to fit a model. Should be a CSV file containing the pharmacokinetic/pharmacodynamic data with appropriate column structure matching the headers specification
dens		Logical. If TRUE, plot histogram/density of residuals instead of scatter
ds_covs		Data.frame. The dataframe with covariates
ds_i		Data.frame. The data frame with source data.

ds_parest	Data.frame. Parameter estimates table with columns parameter and value. Required when fpath_i is NULL; must be provided together with ds_covs. Default is NULL.
dt_obs_fl	Logical. Show observed data points. Default is FALSE
DVID	Restrict SDTAB to one observation type. Numeric values select the DVID column (default 1). If SDTAB has DVNAME, a character or factor is matched to DVNAME first; otherwise a digit-only string is coerced to numeric DVID.
dv_col	Character. Name of DV column in data_i. Default is DV
emp_perc	Logical. Show empirical percentiles. Default is TRUE
errorbar_width	Numeric. Width argument for geom_errorbar. Default 0.2.
est_covmat	Data.frame. Parameter estimation covariance matrix. The first column (X1) must list parameter names; remaining columns (named identically) form the symmetric variance-covariance matrix.
et	An event table passed to sg_sim().
eta_seq	Vector of strings. Character vector of parameter names to be plotted. If NULL, all parameters be included. Default is NULL
excl_col	Character vector. Contains column names to exclude from synthesis. Default: NULL
exclude_vars	Character vector. Contains VAR levels to omit (e.g. "Cc_Cmin"). NULL keeps all rows.
f_scales	String, one of "fixed", "free", "free_x", "free_y". User can specify whether the scales (x and y axes) should be fixed across all panels ("fixed"), free for each panel ("free"), or free only in one dimension ("free_x" or "free_y"). Default is "fixed"
facet_i	String. Column name for facet
fill_i	String. Column name for fill aesthetic. Default is NULL
filt	String. Provide a filter to apply. Default is "T"
fit	Logical. If TRUE, the model fitting will be executed immediately using the specified fitter. If FALSE, only the fit configuration file will be generated without running the fit. Set to FALSE for file preparation only, or TRUE to run the complete fitting process. Default is FALSE.
fpath_i	String or sg-fit object. If the string is given, the path to .Rdata or .json file with sg-fit object is expected.
free_stat	String. Facet scaling option. One of "free", "free_x", "free_y", or "fixed". Default is 'free'.
group_i	String. Primary grouping variable for lines. Default is 'VAR'.
headers	List. A list specifying the data frame column headers. Each element should be a list containing column information with the following structure: <ul style="list-style-type: none"> • name - string, column name in the dataset • use - string, column usage type. Valid values include: <ul style="list-style-type: none"> – "identifier" for subject ID columns – "time" for time columns

	<ul style="list-style-type: none"> – "observation" for dependent variable columns – "observationtype" for observation type identifier – "administration" for administration route – "amount" for dose amount – "eventidentifier" for event ID – "missingdependentvariable" for missing DV flag – "covariate" for covariate columns <ul style="list-style-type: none"> • type - string or NULL, data type specification. For observations use "continuous", "count/categorical" or "event", depending on the nature of observations. For covariates use "continuous" or "categorical". Can be NULL for non-covariate columns.
ncores	Integer. Number of cores used for calculations. Default is 1.
id_col	Character string. Specify the name of the identifier column to exclude from synthesis. Default: NULL.
indiv	Logical. If TRUE, use individual predictions ("IPRED"); otherwise use population predictions ("PRED"). Default is TRUE.
inits	A named vector specifying initial conditions for model variables; Default is NULL.
keep	Vector of strings. Columns of event table to keep in the output data frame. Default is NULL.
lab_x	String. X-axis label.
lab_y	String. Y-axis label.
legend_fl	Logical. Show legend. Default is FALSE.
levels_discrete	Integer. Maximum unique values to consider a variable discrete. Default is 10.
log_x	Logical. If TRUE, a logarithmic scale is applied to x-axis. Default is FALSE.
log_y	Logical. If TRUE, a logarithmic scale is applied to y-axis. Default is FALSE.
log_axes	Logical. If TRUE, a logarithmic scale is applied to all axes. Default is FALSE.
lty_i	String. Column name for linetype aesthetic. Default is NULL.
max_x	Numeric. X-axis maximum limit. Default is NA.
max_y	Numeric. Y-axis maximum limit. Default is NA.
method	Character string. "PRCC" or "eFAST".
min_x	Numeric. X-axis minimum limit. Default is NA.
min_y	Numeric. Y-axis minimum limit. Default is NA.
model	A model object passed to sg_sim().
n_bins	Integer. Number of bins to use in the histogram. Default is 30.
n_quantiles	Integer. Number of quantile groups for continuous variables in col_i. Default is 3.
n_sim	Integer. Number of samples (LHS size for PRCC, base frequency size for eFAST).

no_leg	Logical. If TRUE, the legend is not shown. Default is FALSE
npop	Integer. Number of population replicates. Default is 1.
nsub	Integer. Number of subjects sampled per population (omega/sigma matrices per ID). Default is 1.
occ	A list specifying interoccasion variability properties, containing: <ul style="list-style-type: none"> • <code>init</code> - matrix, initial values for the interoccasion variance-covariance matrix. Same structure as <code>re\$init</code> but for occasion-to-occasion variability. Use 0 for no interoccasion variability • <code>est</code> - matrix, logical matrix specifying which interoccasion variance-covariance elements to estimate. Use TRUE to estimate, FALSE to fix, NA for elements not applicable. Typically all elements are NA when no interoccasion variability is modeled.
opt_name	String. Specify the optimizer/fitter to use for model fitting. Currently supported options: <ul style="list-style-type: none"> • "Monolix" - uses Monolix Suite for population pharmacokinetic modeling (generates .mlxtran files) • "Simurg" - uses SimuRg internal fitter (generates .R files with JSON control structure) Default is "Monolix".
omega	A named matrix or vector.
outputs	Vector of strings. Names of the model variables to output. If NULL, all variables returned. Default is NULL.
output	A character vector of outputs to keep. Passed to <code>sg_sim()</code> .
path_to_fitter	String. The path to the program fitter executable. For Monolix, this should point to the monolix.bat file. If NULL, "C:/ProgramData/Lixoft/MonolixSuite2023R1/bin/monolix.bat" will be used as default. Default is NULL.
path_to_save_output	String. Path to save fit output files. Should be a valid directory path where the fit results and project files will be saved. If NULL, current working directory will be used. Default is NULL.
par_bounds	A tibble or data frame with columns PAR, LB, UB.
par_seq	Vector of strings. Character vector of parameter names to be plotted. If NULL, all parameters be included. Default is NULL
par_type	String. A character string specifying the type of parameters used for theoretical distribution overlay (only relevant when <code>plot_type = 'DIST'</code> and <code>tdist = TRUE</code>). If 'Ind' - Individual (default), distributions are shown on the natural parameter scale assuming log-normal variability. If 'RE' - Random Effect, distributions are shown on the ETA scale, assuming a normal distribution with mean zero and covariance defined by \$OMEGAMAT, without transformation.
params	Character vector of parameter names to vary.
piLow	Numeric. Lower prediction interval bound. Default is 0.10.
piUp	Numeric. Upper prediction interval bound Default is 0.90.
plot_type	Character. Type of plot to produce: <ul style="list-style-type: none"> • "DIST" (default) - histogram of individual parameters,

	<ul style="list-style-type: none"> • "QQ" - QQ-plot of individual parameters.
point_size	Point size for geom_point. Default 2.5.
pred.corr	Logical. Apply prediction correction. Default is FALSE.
project_name	String. The name of the Monolix project without file extension. This will be used as the base name for output files and directories.
quantiles	A numeric vector of length 2. Lower and upper quantiles of the continuous covariate distribution to test. Default is $c(0.1, 0.9)$.
re	List. A list specifying options for random effects used in model fit, containing: <ul style="list-style-type: none"> • <code>init</code> - matrix, initial values for the variance-covariance matrix of random effects. Rows and columns should correspond to parameters defined in <code>theta</code>. Diagonal elements represent variances, off-diagonal elements represent covariances. Use 0 for no variability • <code>est</code> - matrix, logical matrix of same dimensions as <code>init</code> specifying which variance-covariance elements to estimate. Use TRUE to estimate, FALSE to fix, NA to not use this random effect.
ref_line_col	Color for the dashed horizontal line at $y = 1$ (no change from reference). Default "grey25".
rtol	Numeric. A numeric relative tolerance used by the ODE solver to determine if a good solution has been achieved. This is also used in the solved linear model to check if prior doses do not add anything to the solution. Default is 1e-6.
run_id	Integer. Tested model ID. Default is 1.
ruv	A list specifying options for residual error used in model fit, containing: <ul style="list-style-type: none"> • <code>YNAME</code> - string, output name. Typically "y1", "y2",... • <code>DVID</code> - numeric, observation type identifier corresponding to <code>DVID</code> column values • <code>TRANS</code> - string, residual error distribution. Can be: "normal", "logNormal", "logitNormal" • <code>PRED</code> - string, prediction variable name from the model • <code>ERR</code> - string, error model type. Options include: <ul style="list-style-type: none"> - "constant" for additive error - "proportional" for proportional error - "combined1" for combined additive and proportional error • <code>INIT</code> - numeric vector, initial values for error parameters (length depends on error model) • <code>EST</code> - logical vector, whether to estimate each error parameter (same length as <code>INIT</code>) • <code>BLQM</code> - below limit of quantification method (can be NULL) For several observation types list of lists should be provided: one residual error (<code>ruv</code>) object for each observations.
sc_factor	Numeric. Scaling factor for DV/PRED/IPRED values. Default is 1 (no scaling).
scale	A numeric named vector. Scaling for ODE parameters of the system. The names must correspond to the parameter identifiers in the ODE specification. Each of the ODE variables will be divided by the scaling factor. Default is NULL.

seed	Integer. Random seed for synthetic data generation reproducibility. Default is 123.
sigma	A named matrix representing a sigma covariance matrix or its Cholesky decomposition; Default is NULL.
shp_i	String. Column name for shape aesthetic. Default is NULL.
smooth	Logical. Add LOESS smooth line. Default is TRUE.
stat_comp	A character vector of summary statistics to compute. Supported internally: "mean", "median", "min", "max", "sd", "cmax", "SS".
stimes	A numeric vector of simulation times.
task_opt	String. Additional task options to be passed to the fitting software. For Monolix, this can include specific task configurations or optimization settings. When NULL, default tasks (populationParameters, individualParameters, fim, logLikelihood) will be used. Default is NULL.
tdist	Logical. If TRUE, overlay theoretical parameter distributions based on population mean and OMEGA matrix. Default is TRUE.
time_col	String. The column to use as a time column. Currently, can be only TIME. Default is TIME.
theor_perc	Logical. Show theoretical percentiles. Default is TRUE.
theor_percCI	Logical. Show CI around theoretical percentiles. Default is TRUE.
theta	A named numeric vector of baseline parameters. Default is NULL. Parameters listed in params are replaced by sampled values.
thetamat	A named variance-covariance matrix (for parameters brought to normal distribution). Default is NULL.
tsld	Logical. If TRUE, uses time since last dose instead of time from first dose. Default is FALSE.
val_col	String. Name of value column. Default is VALUE.
wrap_i	String. Faceting formula for facet_wrap. Default is NULL.
wrap_ncol	Integer. Number of columns for facet_wrap. Default is NULL.
wrap_nrow	Integer. Number of rows for facet_wrap. Default is NULL.

sg_fit

Run fit with monolix/simurg/nonmem fitter

Description

Run fit with monolix/simurg/nonmem fitter

Usage

```
sg_fit(
  model,
  data,
  headers,
  theta,
  ruv,
  re,
  occ,
  covs,
  project_name,
  task_opt = NULL,
  opt_name = "Monolix",
  fit = FALSE,
  path_to_save_output = NULL,
  path_to_fitter = NULL,
  max_wait_time = 3600
)
```

Arguments

model	string. Path to a txt file file with model structure in Monolix syntax. This should be a valid file path pointing to a model file that defines the pharmacokinetic/pharmacodynamic model structure
data	String. Path to the dataset used to fit a model. Should be a CSV file containing the pharmacokinetic/pharmacodynamic data with appropriate column structure matching the headers specification
headers	List. A list specifying the data frame column headers. Each element should be a list containing column information with the following structure: <ul style="list-style-type: none"> • name - string, column name in the dataset • use - string, column usage type. Valid values include: <ul style="list-style-type: none"> – "identifier" for subject ID columns – "time" for time columns – "observation" for dependent variable columns – "observationtype" for observation type identifier – "administration" for administration route – "amount" for dose amount – "eventidentifier" for event ID – "missingdependentvariable" for missing DV flag – "covariate" for covariate columns • type - string or NULL, data type specification. For observations use "continuous", "count/categorical" or "event", depending on the nature of observations. For covariates use "continuous" or "categorical". Can be NULL for non-covariate columns.
theta	tibble or data.frame. Should contain the following columns:

- NAME - string, parameter name. Should match names specified in the model file
 - TRANS - string, parameter transformation type. Can be: "normal", "logNormal", "logitNormal"
 - INIT - numeric, initial value for the parameter
 - LB - numeric, lower bound for logit transformation, NA for other transformations
 - UB - numeric, upper bound for logit transformation, NA for other transformation
 - EST - logical, whether to estimate this parameter.
- ruv A list specifying options for residual error used in model fit, containing:
- YNAME - string, output name. Typically "y1", "y2",...
 - DVID - numeric, observation type identifier corresponding to DVID column values
 - TRANS - string, residual error distribution. Can be: "normal", "logNormal", "logitNormal"
 - PRED - string, prediction variable name from the model
 - ERR - string, error model type. Options include:
 - "constant" for additive error
 - "proportional" for proportional error
 - "combined1" for combined additive and proportional error
 - INIT - numeric vector, initial values for error parameters (length depends on error model)
 - EST - logical vector, whether to estimate each error parameter (same length as INIT)
 - BLQM - below limit of quantification method (can be NULL) For several observation types list of lists should be provided: one residual error (ruv) object for each observations.
- re List. A list specifying options for random effects used in model fit, containing:
- `init` - matrix, initial values for the variance-covariance matrix of random effects. Rows and columns should correspond to parameters defined in `theta`. Diagonal elements represent variances, off-diagonal elements represent covariances. Use 0 for no variability
 - `est` - matrix, logical matrix of same dimensions as `init` specifying which variance-covariance elements to estimate. Use TRUE to estimate, FALSE to fix, NA to not use this random effect.
- occ A list specifying interoccasion variability properties, containing:
- `init` - matrix, initial values for the interoccasion variance-covariance matrix. Same structure as `re$init` but for occasion-to-occasion variability. Use 0 for no interoccasion variability
 - `est` - matrix, logical matrix specifying which interoccasion variance-covariance elements to estimate. Use TRUE to estimate, FALSE to fix, NA for elements not applicable. Typically all elements are NA when no interoccasion variability is modeled.

covs	<p>A list of covariate structures. Each element should be a list containing covariate relationship definitions with the following structure:</p> <ul style="list-style-type: none"> • PAR - string, name of the parameter to which covariate effect is applied • COVNAME - string, name of the covariate column in the dataset • FUNC - string, functional form of the covariate effect • TRANS - string, transformation applied to covariate • REF - numeric, reference value for categorical covariates • INIT - numeric, initial value for the covariate effect parameter • EST - logical, whether to estimate the covariate effect Can be NULL, if no covariates are used. Default is NULL
project_name	String. The name of the Monolix project without file extension. This will be used as the base name for output files and directories.
task_opt	String. Additional task options to be passed to the fitting software. For Monolix, this can include specific task configurations or optimization settings. When NULL, default tasks (populationParameters, individualParameters, fim, logLikelihood) will be used. Default is NULL.
opt_name	String. Specify the optimizer/fitter to use for model fitting. Currently supported options: <ul style="list-style-type: none"> • "Monolix" - uses Monolix Suite for population pharmacokinetic modeling (generates .mlxtran files) • "Simurg" - uses SimuRg internal fitter (generates .R files with JSON control structure) Default is "Monolix".
fit	Logical. If TRUE, the model fitting will be executed immediately using the specified fitter. If FALSE, only the fit configuration file will be generated without running the fit. Set to FALSE for file preparation only, or TRUE to run the complete fitting process. Default is FALSE.
path_to_save_output	String. Path to save fit output files. Should be a valid directory path where the fit results and project files will be saved. If NULL, current working directory will be used. Default is NULL.
path_to_fitter	String. The path to the program fitter executable. For Monolix, this should point to the monolix.bat file. If NULL, "C:/ProgramData/Lixoft/MonolixSuite2023R1/bin/monolix.bat" will be used as default. Default is NULL.
max_wait_time	numeric. Maximum time in seconds to wait for fit results to complete. Default is 3600 seconds (1 hour). Set to Inf for no timeout.

Value

if option `fit = TRUE`, generalized simurg output object is returned. Otherwise, the file for fit is written and no output is returned

Examples

```
library(tibble)
library(dplyr)
```

```

library(stringr)
# Specify structural model path. For fit, should be in Monolix syntax
model <- system.file("extdata", "models", "model_PK_1c.txt", package = "SimuRg")
# Specify data path. Should be ADPPK-like format
data <- system.file("extdata", "datasets", "dspk-warf.csv", package = "SimuRg")

# Specify headers. Should be a list of lists with the following elements:
# name - string, name of the column
# use - string, use of the column from Monolix documentation
# type - string, type of the column. for use = "covariate", should be
# "continuous" or "categorical", for other uses should be NULL
headers <- list(list(name = "ID", use = "identifier", type = NULL),
               list(name = "TIME", use = "time", type = NULL),
               list(name = "DV", use = "observation", type = "continuous"),
               list(name = "DVID", use = "observationtype", type = NULL),
               list(name = "ADM", use = "administration", type = NULL),
               list(name = "AMT", use = "amount", type = NULL),
               list(name = "EVID", use = "eventidentifier", type = NULL),
               list(name = "MDV", use = "missingdependentvariable", type = NULL),
               list(name = "AGE", use = "covariate", type = "continuous"),
               list(name = "AGE_centered", use = "covariate", type = "continuous"),
               list(name = "SEX", use = "covariate", type = "categorical"),
               list(name = "WEIGHT", use = "covariate", type = "continuous"),
               list(name = "BMI", use = "covariate", type = "continuous"))

# Dataset with the parameters properties. Should be a tibble with the following columns:
# NAME - string, name of the parameter
# TRANS - string, distribution of the parameter. Should be one of the following:
# "normal", "logNormal", "logitNormal"
# INIT - numeric, initial value of the parameter or its fixed value
# LB - numeric, lower bound of the parameter for logit transformation
# UB - numeric, upper bound of the parameter for logit transformation
# EST - logical, estimation status of the parameter. For estimation, should
# be TRUE, for fixed, should be FALSE
theta <- tribble(~NAME, ~TRANS, ~INIT, ~LB, ~UB, ~EST,
                "Cl", "logNormal", 0.2, NA, NA, TRUE,
                "V", "logNormal", 20, NA, NA, TRUE,
                "ka", "logNormal", 0.2, NA, NA, TRUE
                )

# Examples of random effect model specification
# Examples of random effect model specification for fit
# Single observation (legacy format):
# YNAME - string, name of the observation, ususally y1, y2, ...
# DVID - numeric, observation type identifier corresponding to DVID column values
# TRANS - string, residual error distribution. Can be: "normal", "logNormal",
# "logitNormal"
# PRED - string, prediction variable name from the model
# ERR - string, error model type. Options include: "constant" for additive
# error, "proportional" for proportional error, "combined1" for combined
# additive and proportional error
# INIT - numeric vector, initial values for error parameters
# (length depends on error model)
# EST - logical vector, whether to estimate each error parameter

```

```

# (same length as INIT)
# BLQM - below limit of quantification method (can be NULL)
# Single observation (legacy format):
ruv <- list(YNAME = "y1", DVID = 1, TRANS = "normal", PRED = "Cc",
           ERR = "combined1", INIT = c(1, 1), EST = c(TRUE, TRUE), BLQM = NULL)

# Multiple observations (recommended format):
ruv <- list(
  list(YNAME = "y1", DVID = 1, TRANS = "normal", PRED = "Cc",
       ERR = "combined1", INIT = c(1, 1), EST = c(TRUE, TRUE), BLQM = NULL),
  list(YNAME = "y2", DVID = 2, TRANS = "normal", PRED = "EFF",
       ERR = "proportional", INIT = c(0.1), EST = c(TRUE), BLQM = NULL)
)

# Example of random effects (RE) specification.
#
# init matrix: provides the initial values for the random effects.
# est matrix: controls how each random effect is handled:
# TRUE - the random effect will be estimated,
# FALSE - the random effect will be fixed at its initial value,
# NA - no random effect will be applied.
#
# To fit a model without random effects, set all entries in the
# est matrix to NA.
#
# The same logic applies to the between-occasion variability
# matrix (occ).

re <- list(init = tribble(~Cl, ~V, ~ka,
                        1, 0, 0,
                        0, 0, 0,
                        0, 0, 1) %>% as.matrix(),
          est = tribble(~Cl, ~V, ~ka,
                       TRUE, NA, NA,
                       NA, NA, NA,
                       NA, NA, TRUE) %>% as.matrix())

# Example of between-occasion variability (BOV) specification. The structure
# is the same as for RE, but for BOV

occ <- list(init = tribble(~Cl, ~V, ~ka,
                          0, 0, 0,
                          0, 0, 0,
                          0, 0, 0) %>% as.matrix(),
          est = tribble(~Cl, ~V, ~ka,
                       NA, NA, NA,
                       NA, NA, NA,
                       NA, NA, NA) %>% as.matrix())

# Example of covariate specification. Should be a list of lists with the following elements:
# PAR - string, name of the parameter to which the covariate is applied
# COVNAME - string, name of the covariate
# FUNC - string, function to apply to the covariate. Should be "linear" for
# continuous covariates, "categorical" for categorical covariates
# TRANS - string, transformation of the covariate. Should be "median" for

```

```

# continuous covariates, "reference" for categorical covariates
# INIT - numeric, initial value of the covariate
# EST - logical, estimation status of the covariate. For estimation, should
#be TRUE, for fixed, should be FALSE
covs <- list(list(PAR = "V", COVNAME = "AGE", FUNC = "linear",
                 TRANS = "median", INIT = 1, EST = TRUE),
             list(PAR = "ka", COVNAME = "SEX", REF = 0, INIT = 1, EST = TRUE))
output_path <- str_c(tempdir(), "/")
fitter_path <- "C:/ProgramData/Lixoft/MonolixSuite2023R1/bin/monolix.bat"
# Examples of task_opt parameter

task_opt <- paste("populationParameters()", "individualParameters()",
                 "logLikelihood()", sep = "\n")
task_opt_lin <- paste("populationParameters()", "individualParameters()",
                    "fim(method = Linearization)",
                    "logLikelihood(method = Linearization)", sep = "\n")

# Generalized control object
# gco <-list(headers = headers,
#           data = data,
#           model = model,
#           task_opt = task_opt
#           covs = covs,
#           project_name = "test-proj",
#           theta = theta,
#           ruv = ruv,
#           re = re,
#           occ = occ,
#           modelText = "")
result <- sg_fit(model, data, headers, theta, ruv, re, occ, covs,
                project_name = "my_project", fit = FALSE, # set fit = TRUE for fit
                path_to_save_output = output_path,
                path_to_fitter = fitter_path)

```

sg_globalsens_sim

Global sensitivity analysis via PRCC or eFAST

Description

Performs global sensitivity analysis using either PRCC (Partial Rank Correlation Coefficients with LHS sampling) or eFAST (extended Fourier Amplitude Sensitivity Test). For each sampled parameter set, the function runs simulations via `sg_sim()`, reduces trajectories to scalar summary statistics, and computes sensitivity indices for each output–statistic pair.

Usage

```

sg_globalsens_sim(
  method = c("PRCC", "eFAST"),
  model,
  params,

```

```

    par_bounds,
    n_sim,
    stimes,
    output,
    stat_comp = c("mean", "min", "max"),
    et,
    theta = NULL,
    cov = NULL
  )

```

Arguments

method	Character string. "PRCC" or "eFAST".
model	A model object passed to sg_sim().
params	Character vector of parameter names to vary.
par_bounds	A tibble or data frame with columns PAR, LB, UB.
n_sim	Integer. Number of samples (LHS size for PRCC, base frequency size for eFAST).
stimes	A numeric vector of simulation times.
output	A character vector of outputs to keep. Passed to sg_sim().
stat_comp	A character vector of summary statistics to compute. Supported internally: "mean", "median", "min", "max", "sd", "cmax", "SS".
et	An event table passed to sg_sim().
theta	A named numeric vector of baseline parameters. Default is NULL. Parameters listed in params are replaced by sampled values.
cov	Covariates passed to sg_sim(). Default is NULL.

Value

List with:

result	Tibble containing sensitivity analysis results. For PRCC method columns include: PAR, VAR, STAT, estimate, p.value. For eFAST method columns include: PAR, VAR, STAT, TYPE, VALUE, METHOD.
summary	Tibble of scalar summaries used for sensitivity computation: sim.id, VAR, STAT, value.
design	Data.frame of sampled parameter values (real scale) with sim.id. For PRCC this corresponds to LHS samples. For eFAST this corresponds to the FAST design matrix.
bounds	Parameter bounds used in the analysis.

Examples

```

# Example model
library(rxode2)
library(tibble)
source(system.file("extdata", "RxODE_model", "example_rxode_model.R", package = "SimuRg")) # mod_ex

# Set up event table
et_base <- tribble(
  ~id, ~time, ~evid, ~cmt, ~amt, ~addl, ~ii, ~IGFR, ~POPn,
  1, 0, 1, 1, 10, 2, 24, 112, 1
)

# Define parameter bounds
inits <- rxInits(mod_ex)
par_bounds <- tibble::tibble(
  PAR = c("POPCL", "POPVC"),
  LB = inits[c("POPCL", "POPVC")] * (1 - 0.9),
  UB = inits[c("POPCL", "POPVC")] * (1 + 0.9)
)

# Run eFAST sensitivity analysis
res_efast <- sg_globalsens_sim(
  method = c("eFAST"),
  model = mod_ex,
  params = c("POPCL"),
  par_bounds = par_bounds,
  n_sim = 100,
  stimes = seq(0, 168, 10),
  output = "Cc",
  cov = c("IGFR", "POPn"),
  et = et_base,
  stat_comp = c("mean")
)

# Run PRCC sensitivity analysis
res_prcc <- sg_globalsens_sim(
  method = c("PRCC"),
  model = mod_ex,
  params = c("POPCL", "POPVC"),
  par_bounds = par_bounds,
  n_sim = 100,
  stimes = seq(0, 168, 10),
  output = "Cc",
  cov = c("IGFR", "POPn"),
  et = et_base,
  stat_comp = c("mean")
)

```

sg_globalsens_vis *Visualize global sensitivity analysis results*

Description

Generates visualization plots for results produced by `sg_globalsens_sim()`. Supported visualizations:

- PRCC: heatmap or barplot
- eFAST: barplot of first and total order indices

Usage

```
sg_globalsens_vis(  
  x,  
  type = c("heatmap", "bar"),  
  params = NULL,  
  vars = NULL,  
  stats = NULL  
)
```

Arguments

<code>x</code>	Result object returned by <code>sg_globalsens_sim()</code>
<code>type</code>	Plot type for PRCC: "heatmap" or "bar"
<code>params</code>	Optional vector of parameters to display
<code>vars</code>	Optional vector of output variables
<code>stats</code>	Optional vector of statistics to display

Value

ggplot object

Examples

```
# Example model  
library(rxode2)  
library(tibble)  
source(system.file("extdata", "RxODE_model", "example_rxode_model.R", package = "SimuRg")) # mod_ex  
  
# Set up event table  
et_base <- tribble(  
  ~id, ~time, ~evid, ~cmt, ~amt, ~addl, ~ii, ~IGFR, ~POPNN,  
  1, 0, 1, 1, 10, 2, 24, 112, 1  
)
```

```

# Define parameter bounds
inits <- rxInits(mod_ex)
par_bounds <- tibble::tibble(
  PAR = c("POPCL", "POPVC"),
  LB = inits[c("POPCL", "POPVC")] * (1 - 0.9),
  UB = inits[c("POPCL", "POPVC")] * (1 + 0.9)
)

# Run eFAST sensitivity analysis
res_efast <- sg_globalsens_sim(
  method = c("eFAST"),
  model = mod_ex,
  params = c("POPCL"),
  par_bounds = par_bounds,
  n_sim = 100,
  stimes = seq(0, 168, 10),
  output = "Cc",
  cov = c("IGFR", "POPN"),
  et = et_base,
  stat_comp = c("mean")
)

efast_p <- sg_globalsens_vis(
  res_efast,
  params = c("POPCL"),
  vars = "Cc",
  stats = c("mean")
)

efast_p
# Run PRCC sensitivity analysis
res_prcc <- sg_globalsens_sim(
  method = c("PRCC"),
  model = mod_ex,
  params = c("POPCL", "POPVC"),
  par_bounds = par_bounds,
  n_sim = 100,
  stimes = seq(0, 168, 10),
  output = "Cc",
  cov = c("IGFR", "POPN"),
  et = et_base,
  stat_comp = c("mean")
)

prcc_p <- sg_globalsens_vis(
  res_prcc,
  type = c("heatmap"),
  params = c("POPCL", "POPVC"),
  vars = "Cc",
  stats = c("mean")
)

prcc_p

```

sg_gof_obpr

*Observed vs predicted plot***Description**

Function generates observed versus predicted (OBS vs PRED/IPRED) scatter plots, a fundamental goodness-of-fit diagnostic tool in pharmacometric modeling. This visualization assesses model adequacy by comparing observed clinical measurements against model-predicted values, enabling identification of systematic bias, heteroscedasticity, and model misspecification patterns. Function contains options for faceting, coloring by covariates, and trend lines.

Usage

```
sg_gof_obpr(
  fpath_i,
  DVID = 1,
  cov_cols = NULL,
  indiv = TRUE,
  addline = TRUE,
  alpha_i = 0.5,
  smooth = TRUE,
  log_axes = FALSE,
  sc_factor = 1,
  abreaks = scales::pretty_breaks(7),
  lab_x = "Model-predicted values",
  lab_y = "Observed values",
  col_i = NULL,
  col_lab = NULL,
  facet_i = NULL,
  f_scales = "fixed",
  no_leg = FALSE,
  n_quantiles = 3,
  levels_discrete = 10
)
```

Arguments

fpath_i	String or sg-fit object. If the string is given, the path to .Rdata or .json file with sg-fit object is expected.
DVID	Restrict SDTAB to one observation type. Numeric values select the DVID column (default 1). If SDTAB has DVNAME, a character or factor is matched to DVNAME first; otherwise a digit-only string is coerced to numeric DVID.
cov_cols	A character vector specifying the names of the columns with covariates.

indiv	Logical. If TRUE, use individual predictions ("IPRED"); otherwise use population predictions ("PRED"). Default is TRUE.
addline	Logical. If TRUE, lines connecting observations of individual subjects will be added. Default is TRUE.
alpha_i	Numeric. Transparency level (from 0 to 1) for points/lines. Default is 0.5.
smooth	Logical. Add LOESS smooth line. Default is TRUE.
log_axes	Logical. If TRUE, a logarithmic scale is applied to all axes. Default is FALSE.
sc_factor	Numeric. Scaling factor for DV/PRED/IPRED values. Default is 1 (no scaling).
abreaks	A function that generates axis breaks. Default is <code>scales::pretty_breaks(7)</code> .
lab_x	X-axis label. Default "Model-predicted values"
lab_y	Y-axis label. Default "Observed values"
col_i	String. Column name for color
col_lab	String. Label for color legend
facet_i	String. Column name for facet
f_scales	String, one of "fixed", "free", "free_x", "free_y". User can specify whether the scales (x and y axes) should be fixed across all panels ("fixed"), free for each panel ("free"), or free only in one dimension ("free_x" or "free_y"). Default is "fixed"
no_leg	Logical. If TRUE, the legend is not shown. Default is FALSE
n_quantiles	Integer. Number of quantile groups for continuous variables in col_i. Default is 3.
levels_discrete	Integer. Maximum unique values to consider a variable discrete. Default is 10.

Value

A `ggplot2` object

Examples

```
# Basic example with mock data
set.seed(123) # For reproducibility
mock_obj <- list(
  SDTAB = data.frame(
    ID = rep(1:3, each = 5),
    TIME = rep(c(0, 1, 2, 4, 8), 3),
    DV = rnorm(15, mean = 10, sd = 2),
    PRED = rnorm(15, mean = 10, sd = 1.5),
    IPRED = rnorm(15, mean = 10, sd = 1.2),
    MDV = rep(0, 15)
  ),
  COTAB = data.frame(ID = 1:3, AGE = c(30, 45, 60)),
  CATAB = data.frame(ID = 1:3, RACE = c("Hispanic", "Hispanic", "Asian"))
)

# Basic plot
```

```

p <- sg_gof_obpr(mock_obj)

# With covariates and faceting
p <- sg_gof_obpr(
  mock_obj,
  cov_cols = "RACE",
  col_i = "RACE",
  facet_i = "RACE"
)

```

sg_gof_par_cov

Plot random effects or individual parameters vs covariates

Description

Generates ggplot2 visualizations for either random effects (RE) or individual parameters (IndPar) versus continuous or categorical covariates from a Simurg object.

Usage

```
sg_gof_par_cov(fpath_i, ptype = "REvsCov", cat_cov = NULL, cont_cov = NULL)
```

Arguments

fpath_i	String or sg-fit object. If the string is given, the path to .Rdata or .json file with sg-fit object is expected.
ptype	Character. Type of plot: "REvsCov" for random effects or "IndParvsCov" for individual parameters.
cat_cov	Optional tibble with categorical covariates. Must have columns COV and optionally COVNAME for labels.
cont_cov	Optional tibble with continuous covariates. Must have columns COV and optionally COVNAME for labels.

Value

A list of ggplot objects. Returns versus continuous covariates vs_contcov and versus categorical covariates vs_catcov plots.

Examples

```

library(tibble)
cont_cov <- tibble(
  COV = c("AGE", "WEIGHT"),
  COVNAME = c("Age, years", "Body weight, kg")
)
cat_cov <- tibble(

```

```

COV = c("SEX", "VKORC1_gentyp"),
COVNAME = c("Sex, M/F", "VKORC1 genotype")
)
fpath_i <- system.file("extdata", "simurg_object", "Warfarin_PK.RData", package = "SimuRg")
p <- sg_gof_par_cov(
  fpath_i = fpath_i,
  ptype = "IndParvsCov",
  cont_cov = cont_cov,
  cat_cov = cat_cov
)
p$vs_contcov
p$vs_catcov

```

sg_gof_par_dist	<i>Plot distributions of individual parameters with optional theoretical density</i>
-----------------	--

Description

This function creates histograms of individual parameter estimates, with optional overlay of the theoretical distribution based on the population parameters and OMEGA matrix.

Usage

```

sg_gof_par_dist(
  fpath_i,
  par_seq = NULL,
  par_type = "Ind",
  n_bins = 30,
  tdist = TRUE,
  plot_type = "DIST"
)

```

Arguments

fpath_i	String or sg-fit object. If the string is given, the path to .Rdata or .json file with sg-fit object is expected.
par_seq	Vector of strings. Character vector of parameter names to be plotted. If NULL, all parameters be included. Default is NULL
par_type	String. A character string specifying the type of parameters used for theoretical distribution overlay (only relevant when plot_type = 'DIST' and tdist = TRUE). If 'Ind' - Individual (default), distributions are shown on the natural parameter scale assuming log-normal variability. If 'RE' - Random Effect, distributions are shown on the ETA scale, assuming a normal distribution with mean zero and covariance defined by \$OMEGAMAT, without transformation.
n_bins	Integer. Number of bins to use in the histogram. Default is 30.

<code>tdist</code>	Logical. If TRUE, overlay theoretical parameter distributions based on population mean and OMEGA matrix. Default is TRUE.
<code>plot_type</code>	Character string specifying the type of plot to generate. Options: 'DIST' for parameter distributions (default), 'QQ' for Q-Q plots, 'correlations' for correlation matrix plot.

Details

The function visualizes the distribution of post hoc individual parameter estimates (from \$PATAB) and optionally compares them with the expected population-level variability using \$SUMTAB\$DISTRIBUTION (see Details). Shrinkage values are calculated based on ETAsrinkage_var from \$SUMTAB and included in facet labels.

Theoretical distributions are generated by sampling from a multivariate normal distribution (mean zero, covariance from \$OMEGAMAT) and mapping to the individual-parameter scale using typical values and the DISTRIBUTION column: logNormal uses $TV * \exp(ETA)$, normal uses $TV + ETA$, logitNormal uses the inverse logit of $\logit(TV) + ETA$. Parameters with missing or empty DISTRIBUTION do not get a theoretical curve.

Value

A ggplot object containing histogram(s) of individual parameter distributions, optionally overlaid with theoretical densities.

Examples

```
# Basic usage: distribution plots for all parameters
fpath_i <- system.file("extdata", "simurg_object", "Warfarin_PK.RData", package = "SimuRg")
sg_gof_par_dist(fpath_i = fpath_i)

# Distribution plots for selected parameters (natural scale)
sg_gof_par_dist(fpath_i = fpath_i, par_seq = c("ka", "Cl"))

# Distribution plots with theoretical densities disabled
sg_gof_par_dist(fpath_i = fpath_i, tdist = FALSE)

# Distribution plots for ETA
sg_gof_par_dist(fpath_i = fpath_i, par_seq = c("eta_ka", "eta_Cl"), par_type = "RE")

# Q-Q plots for all ETA parameters
sg_gof_par_dist(fpath_i = fpath_i, plot_type = "QQ")

# Q-Q plot for a specific ETA parameter
sg_gof_par_dist(fpath_i = fpath_i, plot_type = "QQ", par_seq = "eta_ka")

# Correlation matrix for selected parameters
sg_gof_par_dist(fpath_i = fpath_i, plot_type = "correlations")
```

sg_gof_res

*Create residual diagnostic plots***Description**

Generates residual diagnostic plots versus time/predictions. Supports faceting, covariate coloring, quantile binning for continuous covariates, and optional smoothing.

Usage

```
sg_gof_res(
  fpath_i,
  DVID = 1,
  cov_cols = NULL,
  indiv = TRUE,
  vs_time = TRUE,
  weighted = TRUE,
  addline = TRUE,
  alpha_i = 0.5,
  smooth = TRUE,
  log_x = FALSE,
  abreaks = scales::pretty_breaks(7),
  lab_y = NULL,
  lab_x = NULL,
  col_i = NULL,
  col_lab = NULL,
  facet_i = NULL,
  f_scales = "fixed",
  n_bins = 50,
  min_y = NA,
  max_y = NA,
  min_x = NA,
  max_x = NA,
  legend_fl = FALSE,
  n_quantiles = 3,
  levels_discrete = 10
)
```

Arguments

fpath_i	String or sg-fit object. If the string is given, the path to .Rdata or .json file with sg-fit object is expected.
DVID	Restrict SDTAB to one observation type. Numeric values select the DVID column (default 1). If SDTAB has DVNAME, a character or factor is matched to DVNAME first; otherwise a digit-only string is coerced to numeric DVID.
cov_cols	A character vector specifying the names of the columns with covariates.

indiv	Logical. If TRUE, use individual predictions ("IPRED"); otherwise use population predictions ("PRED"). Default is TRUE.
vs_time	Logical. If TRUE, plot residuals vs TIME; otherwise, plot residuals vs predictions (IPRED/PRED).
weighted	Logical. If TRUE, use weighted residuals; otherwise, use RES/IRES
addline	Logical. If TRUE, lines connecting observations of individual subjects will be added. Default is TRUE.
alpha_i	Numeric. Transparency level (from 0 to 1) for points/lines. Default is 0.5.
smooth	Logical. Add LOESS smooth line. Default is TRUE.
log_x	Logical. If TRUE, a logarithmic scale is applied to x-axis. Default is FALSE.
abreaks	A function that generates axis breaks. Default is scales::pretty_breaks(7).
lab_y	String. Y-axis label.
lab_x	String. X-axis label.
col_i	String. Column name for color
col_lab	String. Label for color legend
facet_i	String. Column name for facet
f_scales	String, one of "fixed", "free", "free_x", "free_y". User can specify whether the scales (x and y axes) should be fixed across all panels ("fixed"), free for each panel ("free"), or free only in one dimension ("free_x" or "free_y"). Default is "fixed"
n_bins	Integer. Number of bins to use in the histogram. Default is 30.
min_y	Numeric. Y-axis minimum limit. Default is NA.
max_y	Numeric. Y-axis maximum limit. Default is NA.
min_x	Numeric. X-axis minimum limit. Default is NA.
max_x	Numeric. X-axis maximum limit. Default is NA.
legend_fl	Logical. Show legend. Default is FALSE.
n_quantiles	Integer. Number of quantile groups for continuous variables in col_i. Default is 3.
levels_discrete	Integer. Maximum unique values to consider a variable discrete. Default is 10.

Value

A ggplot2 object

Examples

```
# Basic example with mock data
set.seed(123) # For reproducibility
n_subjects <- 50
mock_obj <- list(
  SDTAB = do.call(rbind, lapply(1:n_subjects, function(id) {
    n_obs <- 6
```

```

times <- sort(runif(n_obs, min = 0, max = 24)) # random times between 0 and 24h
data.frame(
  ID    = id,
  TIME  = times,
  DV    = rnorm(n_obs, mean = 10, sd = 2),
  PRED  = rnorm(n_obs, mean = 10, sd = 1.5),
  IPRED = rnorm(n_obs, mean = 10, sd = 1.2),
  IWRES = rnorm(n_obs, mean = 0, sd = 0.8),
  IRES  = rnorm(n_obs, mean = 0, sd = 1.2),
  MDV   = 0
)
)),
COTAB = data.frame(
  ID = 1:n_subjects,
  AGE = sample(20:80, n_subjects, replace = TRUE)
),
CATAB = data.frame(
  ID = 1:n_subjects,
  RACE = sample(c("Hispanic", "Asian", "Caucasian"), n_subjects, replace = TRUE)
)
)

# Basic plot: individual weighted residuals vs TIME (weighted = TRUE, vs_time = TRUE)
p <- sg_gof_res(mock_obj, smooth = FALSE)

# With covariates and faceting (use RACE as facet and AGE as color)
p <- sg_gof_res(
  mock_obj,
  smooth = TRUE,
  cov_cols = c("RACE", "AGE"),
  col_i = "RACE",
  facet_i = "AGE",
  indiv = TRUE,
  weighted = TRUE,
  vs_time = FALSE,
  legend_fl = TRUE
)
p

```

sg_gof_res_dist

Plot distribution or QQ-plot of residuals from NONMEM/Simurg run

Description

This function visualizes residuals from estimation results stored in a Simurg output object. The residuals can be plotted either as histograms with overlaid normal density curves, or as QQ-plots to assess normality.

Usage

```
sg_gof_res_dist(
  fpath_i,
  res_type = "RES",
  DVID = 1,
  n_bins = 30,
  ndist = TRUE,
  plot_type = "DIST"
)
```

Arguments

fpath_i	String or sg-fit object. If the string is given, the path to .Rdata or .json file with sg-fit object is expected.
res_type	Character vector. One or several types of residuals to plot. Values must correspond to column name(s) in smrg_obj\$SDTAB, e.g. "RES", "IWRES", "IRES". If multiple residual types are provided, plots will be generated for each of them.
DVID	Restrict SDTAB to one observation type. Numeric values select the DVID column (default 1). If SDTAB has DVNAME, a character or factor is matched to DVNAME first; otherwise a digit-only string is coerced to numeric DVID.
n_bins	Integer. Number of bins to use in the histogram. Default is 30.
ndist	Logical. If TRUE, overlays the corresponding normal density curve on the histogram (default: TRUE).
plot_type	Character. Type of plot to produce: <ul style="list-style-type: none"> • "DIST" (default) - histogram of individual parameters, • "QQ" - QQ-plot of individual parameters.

Value

A ggplot object.

Examples

```
# Plot the distribution of individual weighted residuals (IWRES)
# (default \code{DVID = 1}; pass \code{DVID} or a \code{DVNAME} string to select another endpoint)
fpath_i <- system.file("extdata", "simurg_object", "Warfarin_PK.RData",
  package = "SimuRg")
sg_gof_res_dist(fpath_i = fpath_i, res_type = "IWRES")

# QQ-plots for the same default endpoint
sg_gof_res_dist(fpath_i = fpath_i, res_type = "RES", plot_type = "QQ")

# Several residual columns at once (still one \code{DVID} unless you change it)
sg_gof_res_dist(fpath_i = fpath_i, res_type = c("IWRES", "IRES"))
```

sg_gof_tp

*Plot time profiles of the fitted data***Description**

Plot time profiles of the fitted data

Usage

```
sg_gof_tp(
  fpath_i,
  pop = TRUE,
  filt = "T",
  cov_cols = NULL,
  col_i = NULL,
  DVID = 1,
  tsld = FALSE,
  f_scales = "free",
  sort_by = NULL,
  desc = FALSE,
  log_y = FALSE,
  lab_x = "Time since first dose, h",
  lab_y = "Plasma concentration, mmol/L",
  cap = str_c("empty circles - observed data\n", "solid lines with point - ",
    "individual predictions\n", "dashed grey lines with ",
    "point - population predictions"),
  n_quantiles = 3,
  levels_discrete = 10
)
```

Arguments

fpath_i	String or sg-fit object. If the string is given, the path to .Rdata or .json file with sg-fit object is expected.
pop	Logical. If TRUE (default), adds population predictions (PRED) as dashed lines and points.
filt	String. Provide a filter to apply. Default is "T"
cov_cols	A character vector specifying the names of the columns with covariates.
col_i	String. Column name for color
DVID	Restrict SDTAB to one observation type. Numeric values select the DVID column (default 1). If SDTAB has DVNAME, a character or factor is matched to DVNAME first; otherwise a digit-only string is coerced to numeric DVID.
tsld	Logical. If TRUE, uses time since last dose instead of time from first dose. Default is FALSE.

f_scales	String, one of "fixed", "free", "free_x", "free_y". User can specify whether the scales (x and y axes) should be fixed across all panels ("fixed"), free for each panel ("free"), or free only in one dimension ("free_x" or "free_y"). Default is "fixed"
sort_by	Character vector of column names to sort ID facets by. Use "DOSE" to sort by last dose. Other names (e.g. covariates) must exist in SimuRg object.
desc	Logical. If TRUE, sort in descending order for all columns in sort_by.
log_y	Logical. If TRUE, a logarithmic scale is applied to y-axis. Default is FALSE.
lab_x	String. X-ax label. Default is "Time since first dose, h"
lab_y	String. Y-ax label. Default is "Plasma concentration, mmol/L"
cap	String. Plot caption. Default is "empty circles - observed data solid lines with point - individual predictions dashed grey lines with point - population predictions"
n_quantiles	Integer. Number of quantile groups for continuous variables in col_i. Default is 3.
levels_discrete	Integer. Maximum unique values to consider a variable discrete. Default is 10.

Value

A list of plots with predicted time profiles, faceted by id

Examples

```
fpath_i <- system.file("extdata", "simurg_object", "Warfarin_PK.RData",
                      package = "SimuRg")
plot_list <- sg_gof_tp(fpath_i)
print(plot_list[[1]])
```

sg_localsens_sim *Perform local sensitivity analysis simulations*

Description

Generates simulation datasets for a model varying each specified parameter within provided bounds or relative percentage. Supports multiple outputs and covariates.

Usage

```
sg_localsens_sim(
  model,
  params,
  stimes,
  output = NULL,
  perc = 0.2,
```

```

  lb = NULL,
  ub = NULL,
  cov = NULL,
  et,
  theta = NULL,
  n_sim = 10
)

```

Arguments

model	An RxODE model object.
params	Character vector of parameter names to vary in the analysis.
stimes	Numeric vector of time points for the simulation.
output	Character vector of model outputs (variables) to keep. Default is NULL (all outputs).
perc	Numeric. If lb/ub not provided, defines \pm percentage range around parameter base value. Default 0.2.
lb	Numeric vector of lower bounds for each parameter. Length must match params. Default NULL.
ub	Numeric vector of upper bounds for each parameter. Length must match params. Default NULL.
cov	Character vector of covariate names to include in the simulation. Default NULL.
et	Event table for the simulation.
theta	Named numeric vector of baseline parameters. Default NULL.
n_sim	Integer. Number of points to simulate between LB and UB for each parameter. Default 10.

Value

A tibble containing the simulated results with the following columns:

NPOP Optional. Population identification number. From simulation.

ID Optional. Individual subject ID. From simulation.

TIME Time points of simulation.

VAR Name of the output variable (model compartment or biomarker).

VALUE Simulated value of the variable.

mean, median, min, max, sd, etc. Optional. Aggregated statistics, if aggregation applied.

COV1,...COVn Optional. Covariate values, if cov provided.

PARNAME Name of the parameter being varied in this simulation.

PARVAL Value of the parameter used in this simulation.

PARVAL_NORM Normalized parameter value between 0 and 1 relative to LB and UB. Useful for plotting color gradients.

Examples

```

library(rxode2)
library(tibble)
mod_ex <- RxODE({
  # Doses in mg
  # Time in hours
  ### Parameter values
  # Typical
  POPCL = 5;
  POPVC = 180;
  POPQ = 7;
  POPVP = 52;
  POPKTR = 6;
  FBIOPAR = 1;
  # Covariate coefficients
  POPIGFRCOV = 0.8;
  POPPATCOVCLGR1 = 0.85;
  POPPATCOVCLGR2 = 0.85;
  POPPATCOVCLGR3 = 0.85;
  POPPATCOVCLGR4 = 0.85;
  # Random effects
  PPVCL = 0;
  PPVVC = 0;
  PPVKTR = 0;
  # Residual error
  RUV = 0;
  ### Covariates
  PATCOVCL = 1
  if(POP1 == 1){PATCOVCL = POPPATCOVCLGR1}
  if(POP1 == 2){PATCOVCL = POPPATCOVCLGR2}
  if(POP1 == 3){PATCOVCL = POPPATCOVCLGR3}
  if(POP1 == 4){PATCOVCL = POPPATCOVCLGR4}
  IGFRCOV = (IGFR/112)^POPIGFRCOV
  ## Parameters
  CL = POPCL * IGFRCOV * PATCOVCL * exp(PPVCL);
  VC = POPVC * exp(PPVVC);
  Q = POPQ;
  VP = POPVP;
  KTR = POPKTR * exp(PPVKTR);
  ### Explicit functions
  Cc = Ac/VC;          # nmol/L
  Cp = Ap/VP;         # nmol/L
  ### Initial conditions
  At1(0) = 0;        # mg
  At2(0) = 0;        # mg
  At3(0) = 0;        # mg
  At4(0) = 0;        # mg
  At5(0) = 0;        # mg
  At6(0) = 0;        # mg
  Ad(0) = 0;         # mg
  Ac(0) = 0;         # mg
  Ap(0) = 0;         # mg

```

```

### ODEs
d/dt(At1) = - KTR*At1;
d/dt(At2) = KTR*At1 - KTR*At2;
d/dt(At3) = KTR*At2 - KTR*At3;
d/dt(At4) = KTR*At3 - KTR*At4;
d/dt(At5) = KTR*At4 - KTR*At5;
d/dt(At6) = KTR*At5 - KTR*At6;
d/dt(Ad) = KTR*At6 - KTR*Ad;
d/dt(Ac) = KTR*Ad - CL*Cc - Q*(Cc - Cp);
d/dt(Ap) = Q*(Cc - Cp);
FBI0 = FBIOPAR
f(At1) = FBI0*1000000/505;
CHECKRUV = RUV;
Cc_ResErr = Cc + RUV*Cc;
})
et_base <- tribble(
  ~id, ~time, ~evid, ~cmt, ~amt, ~addl, ~ii, ~IGFR, ~POPN,
  1, 0, 1, 1, 10, 2, 24, 112, 1
)
sens_loc <- sg_localsens_sim(
  model = mod_ex,
  params = c("POPCL", "POPVC"),
  stimes = seq(0, 168, 0.1),
  output = "Cc",
  perc = 0.9,
  cov = c("IGFR", "POPN"),
  et = et_base
)

```

sg_localsens_vis

Visualize Local Sensitivity Analysis Results

Description

This function creates plots to visualize the results of a local sensitivity analysis: (1) a family of curves plot showing how model outputs vary over time for different parameter perturbations, and (2) a tornado plot summarizing the relative influence of each parameter on selected summary metrics (e.g., output value at a chosen time or Cmax).

Usage

```

sg_localsens_vis(
  sens_data,
  tornado_time = NULL,
  metrics = "value",
  ref_data = NULL,
  log_scale = FALSE,

```

```

    color_low = "#1f77b4",
    color_high = "#ff7f0e",
    facet_scales = "free"
  )

```

Arguments

sens_data	A data frame containing sensitivity analysis results from <code>sg_localsens_sim()</code>
tornado_time	Numeric value specifying the time point (in the same units as TIME) at which to evaluate the model output for the tornado plot. Defaults to the maximum time in <code>sens_data</code> if not specified.
metrics	Character vector specifying which metrics to include in the tornado plot. Supported options are "value" (value at <code>tornado_time</code>) and "cmax" (maximum output). Defaults to "value".
ref_data	Optional data frame providing reference (baseline) output values, formatted similarly to <code>sens_data</code> . If NULL, the function uses the mid-range parameter value as the baseline.
log_scale	Logical. If TRUE, the y-axis of the family-of-curves plot is displayed on a log scale.
color_low	Character. Color used for the lower parameter bound (default: "#1f77b4").
color_high	Character. Color used for the upper parameter bound (default: "#ff7f0e").
facet_scales	Character string passed to <code>facet_grid()</code> , defining how y-axis scales behave across facets (default = "free").

Value

A named list containing:

`family_of_curves` A `ggplot2` object showing parameter sensitivity curves over time.

`tornado` A `ggplot2` object showing the tornado plot of relative sensitivity.

Examples

```

library(rxode2)
library(tibble)
mod_ex <- RxODE({
  # Doses in mg
  # Time in hours
  ### Parameter values
  # Typical
  POPCL = 5;
  POPVC = 180;
  POPQ = 7;
  POPVP = 52;
  POPKTR = 6;
  FBIOPAR = 1;
  # Covariate coefficients
  POPIGFRCOV = 0.8;

```

```

POPPATCOVCLGR1 = 0.85;
POPPATCOVCLGR2 = 0.85;
POPPATCOVCLGR3 = 0.85;
POPPATCOVCLGR4 = 0.85;
# Random effects
PPVCL = 0;
PPVVC = 0;
PPVKTR = 0;
# Residual error
RUV = 0;
### Covariates
PATCOVCL = 1
if(POPn == 1){PATCOVCL = POPPATCOVCLGR1}
if(POPn == 2){PATCOVCL = POPPATCOVCLGR2}
if(POPn == 3){PATCOVCL = POPPATCOVCLGR3}
if(POPn == 4){PATCOVCL = POPPATCOVCLGR4}
IGFRCOV = (IGFR/112)^POPIGFCOV
## Parameters
CL = POPCL * IGFRCOV * PATCOVCL * exp(PPVCL);
VC = POPVC * exp(PPVVC);
Q = POPQ;
VP = POPVP;
KTR = POPKTR * exp(PPVKTR);
### Explicit functions
Cc = Ac/VC;          # nmol/L
Cp = Ap/VP;          # nmol/L
### Initial conditions
At1(0) = 0;          # mg
At2(0) = 0;          # mg
At3(0) = 0;          # mg
At4(0) = 0;          # mg
At5(0) = 0;          # mg
At6(0) = 0;          # mg
Ad(0) = 0;           # mg
Ac(0) = 0;           # mg
Ap(0) = 0;           # mg
### ODEs
d/dt(At1) = - KTR*At1;
d/dt(At2) = KTR*At1 - KTR*At2;
d/dt(At3) = KTR*At2 - KTR*At3;
d/dt(At4) = KTR*At3 - KTR*At4;
d/dt(At5) = KTR*At4 - KTR*At5;
d/dt(At6) = KTR*At5 - KTR*At6;
d/dt(Ad) = KTR*At6 - KTR*Ad;
d/dt(Ac) = KTR*Ad - CL*Cc - Q*(Cc - Cp);
d/dt(Ap) = Q*(Cc - Cp);
FBI0 = FBIOPAR
f(At1) = FBI0*1000000/505;
CHECKRUV = RUV;
Cc_ResErr = Cc + RUV*Cc;
})
et_base <- tribble(
  ~id, ~time, ~evid, ~cmt, ~amt, ~addl, ~ii, ~IGFR, ~POPn,

```

```

    1, 0, 1, 1, 10, 2, 24, 112, 1
  )
sens_loc <- sg_localsens_sim(
  model = mod_ex,
  params = c("POPCL", "POPVC"),
  stimes = seq(0, 168, 0.1),
  output = "Cc",
  perc = 0.9,
  cov = c("IGFR", "POPN"),
  et = et_base
)
plots <- sg_localsens_vis(
  sens_data = sens_loc,
  tornado_time = 168,
  metrics = c("value", "cmax"),
  log_scale = TRUE,
  color_low = "#4575b4",
  color_high = "#d73027",
  facet_scales = "free"
)

# Display plots
print(plots$family_of_curves)
print(plots$tornado)

```

sg_modbuild

Build estimation model scenarios

Description

Constructs and exports multiple model configuration scenarios by combining population parameters, random effects, residual variability, occasion effects, and covariate assignments. Each scenario is defined by unique combinations of these model components and exported as `.mlxtran` files. Additionally, a summary CSV file describing all scenarios is generated.

Usage

```

sg_modbuild(
  mod_lst,
  data,
  headers,
  ruv_lst,
  theta_lst,
  re_lst,
  path,
  occ_lst,
  covs_lst = NULL,
  task_lst = NULL,

```

```

    opt_name = "Simurg",
    project_name = "my_project"
  )

```

Arguments

mod_lst	List of model file paths or model identifiers used for scenario generation.
data	Character string. Path to the dataset used in model fitting or simulation.
headers	Predefined list of dataset column names (e.g., ID, TIME, DV, etc.) used by the modeling framework.
ruv_lst	List specifying residual unexplained variability (RUV) structures. Each element contains RUV properties (e.g., YNAME, ERR) and mapping to data.
theta_lst	List of tibbles describing population parameter properties. Each tibble should contain columns such as NAME, INIT, and EST.
re_lst	List of random effects (RE) specifications. Each element includes matrices <code>init</code> and <code>est</code> defining initial and estimated covariance structures.
path	Character. Directory path where output files (CSV summary and model files) will be written. Default is current working directory.
occ_lst	List of occasion effect matrices. Each element includes <code>init</code> and <code>est</code> matrices defining inter-occasion variability.
covs_lst	Optional list describing covariate-parameter relationships. Each element should include fields such as PAR, COVNAME, and EST indicating inclusion in the model.
task_lst	Optional list defining additional tasks or configuration options for each scenario. Default is NULL.
opt_name	Character. Optimization engine name (e.g., "Monolix", "Simurg"). Default "Simurg".
project_name	Character. Base name for the exported project files. Default "my_project".

Details

This function automates the construction of multiple model configurations for model evaluation, sensitivity analysis, or scenario testing. It expands parameter and variability definitions, constructs all possible combinations, and exports model specifications for external fitting tools.

Value

The function writes two types of outputs to the specified path:

- A CSV file (`scenarios_info.csv`) summarizing all generated scenarios with columns:
 - scenario_number** Unique index of the scenario.
 - model** Model file or path used in the scenario.
 - data** Path to the dataset used.
 - theta** Active population parameters and initial values used.
 - RUV** Residual error structure(s) used.
 - RE** Random effect and correlation terms included.

OCC Active occasion effects.

COVS Included covariate-parameter relationships.

- Individual .mlxtran model files for each generated scenario, named according to the pattern <project_name>_<i>.mlxtran.

Examples

```
library(dplyr)
folder_path <- system.file("extdata", package = "SimuRg")
mod_lst <- list(paste(folder_path, "/models/model_PK_1c.txt", sep = "/"),
               paste(folder_path, "/models/model_PK_2c.txt", sep = "/"))

### path to the dataset
data <- paste(folder_path, "datasets", "dspk-warf.csv", sep = "/")
re_lst_1 <- list(
  list(init = tribble(~Cl, ~Vd, ~ka, ~Vp, ~Q,
                    1, 0, 0, 0, 0,
                    0, 1, 0, 0, 0,
                    0, 0, 1, 0, 0,
                    0, 0, 0, 1, 0,
                    0, 0, 0, 0, 1) %>% as.matrix(),
    est = tribble(~Cl, ~Vd, ~ka, ~Vp, ~Q,
                 TRUE, NA, NA, NA, NA,
                 NA, TRUE, NA, NA, NA,
                 NA, NA, TRUE, NA, NA,
                 NA, NA, NA, TRUE, NA,
                 NA, NA, NA, NA, TRUE) %>% as.matrix(),
    block = tribble(~Cl, ~Vd, ~ka, ~Vp, ~Q,
                   FALSE, NA, NA, NA, NA,
                   NA, FALSE, NA, NA, NA,
                   NA, NA, TRUE, NA, NA,
                   NA, NA, NA, FALSE, NA,
                   NA, NA, NA, NA, FALSE) %>% as.matrix())
)
headers <- list(list(name = "ID", use = "identifier", type = NULL),
                list(name = "TIME", use = "time", type = NULL),
                list(name = "DV", use = "observation", type = "continuous"),
                list(name = "DVID", use = "observationtype", type = NULL),
                list(name = "ADM", use = "administration", type = NULL),
                list(name = "AMT", use = "amount", type = NULL),
                list(name = "EVID", use = "eventidentifier", type = NULL),
                list(name = "MDV", use = "missingdependentvariable", type = NULL),
                list(name = "AGE", use = "covariate", type = "continuous"),
                list(name = "AGE_centered", use = "covariate", type = "continuous"),
                list(name = "SEX", use = "covariate", type = "categorical"),
                list(name = "WEIGHT", use = "covariate", type = "continuous"),
                list(name = "BMI", use = "covariate", type = "continuous"))

ruv_lst_2 <- list(
  # structural model 1
  list(
    list(YNAME = "y1",
```

```

    DVID = 1,
    TRANS = "normal",
    PRED = "Cc",
    ERR = list("constant", "proportional", "combined1"),
    # options to test (can be length = 1, i.e., "constant" or "combined1")
    INIT = list(1, 1, c(1, 1)),
    # options to test (can be length = 1, i.e., 1 or c(1, 1))
    EST = list(TRUE, TRUE, c(TRUE, TRUE))
    # options to test (can be length = 1, i.e., T or c(T, T))BLQM = NULL))
  ),
  # structural model 2
  list(
    list(YNAME = "y1",
      DVID = 1,
      TRANS = "normal",
      PRED = "Cc",
      ERR = list("constant", "proportional"),
      # options to test (can be length = 1, i.e., "constant" or "combined1")
      INIT = list(1, 1), # options to test (can be length = 1, i.e., 1 or c(1, 1))
      EST = list(TRUE, TRUE)
      # options to test (can be length = 1, i.e., T or c(T, T))BLQM = NULL))
    )
  )
)

theta_lst_2 <- list(
  # structural model 1
  tribble(~NAME, ~TRANS, ~INIT, ~LB, ~UB, ~EST,
    "Cl", "logNormal", 0.2, NA, NA, TRUE,
    "Vd", "logNormal", 20, NA, NA, TRUE,
    "ka", "logNormal", 0.2, NA, NA, TRUE),
  # structural model 2
  tribble(~NAME, ~TRANS, ~INIT, ~LB, ~UB, ~EST,
    "Cl", "logNormal", 0.2, NA, NA, TRUE,
    "Vd", c("Normal", "logNormal"), c(10, 20, 30), NA, NA, TRUE,
    "ka", "logNormal", c(0.2, 0.1, 0.5), NA, NA, TRUE,
    # INIT could be length > 1
    "Vp", c("Normal", "logNormal"), 10, NA, NA, TRUE,
    "Q", "logNormal", 5, NA, NA, TRUE))

path <- tempdir()
sg_modbuild(
  mod_lst = mod_lst[1],
  data = data,
  headers = headers,
  ruv_lst = ruv_lst_2,
  theta_lst = theta_lst_2,
  re_lst = re_lst_1,
  occ_lst = re_lst_1,
  covs_lst = NULL,
  path = path,
  project_name = "tests_test_project"
)
clr_files <- list.files(path, full.names = TRUE)

```

```
unlink(cldr_files, recursive = TRUE, force = TRUE)
```

sg_modcomp	<i>Returns dataframe with summary statistics for model comparison</i>
------------	---

Description

Returns dataframe with summary statistics for model comparison

Usage

```
sg_modcomp(fpath_i, run_id = 1)
```

Arguments

fpath_i	String or sg-fit object. If the string is given, the path to .Rdata or .json file with sg-fit object is expected.
run_id	Integer. Tested model ID. Default is 1.

Value

A data.frame with model summary metrics for further comparison

Examples

```
fpath_i <- system.file("extdata", "simurg_object", "Warfarin_PK.RData", package = "SimuRg")
sum_tab <- sg_modcomp(fpath_i, 3)
print(sum_tab)
```

sg_multistart	<i>Build multistart scenarios with varying initial values</i>
---------------	---

Description

Generates multiple scenarios with identical model structure (model, RUV, RE, OCC, COVS) and different sampled initial values for population parameters.

For each start, initial values of estimated parameters are sampled uniformly within specified intervals, and a corresponding .mlxtran file is generated. A summary table describing all multistart scenarios is written to disk.

Usage

```

sg_multistart(
  mod,
  data,
  headers,
  ruv,
  theta,
  re,
  occ,
  path,
  covs_lst = NULL,
  opt_name = "Simurg",
  project_name = "multistart_project",
  n_starts = 10,
  theta_intervals = NULL,
  interval_factor = c(0.2, 5),
  vary_params = NULL,
  seed = NULL
)

```

Arguments

mod	Model file path or model identifier used for scenario generation.
data	Character string. Path to the dataset used in model fitting or simulation.
headers	Predefined list of dataset column names (e.g., ID, TIME, DV, etc.) used by the modeling framework.
ruv	Residual unexplained variability (RUV) structure. Each element contains RUV properties (e.g., YNAME, ERR) and mapping to data.
theta	Tibble describing population parameter properties. Tibble should contain columns such as NAME, INIT, and EST.
re	Random effects (RE) specification. Includes matrices <code>init</code> and <code>est</code> defining initial and estimated covariance structures.
occ	Occasion effect matrices. Includes <code>init</code> and <code>est</code> matrices defining inter-occasion variability.
path	Character. Directory path where output files (CSV summary and model files) will be written. Default is current working directory.
covs_lst	Optional list describing covariate-parameter relationships. Each element should include fields such as PAR, COVNAME, and EST indicating inclusion in the model.
opt_name	Character. Optimization engine name (e.g., "Monolix", "Simurg"). Default "Simurg".
project_name	Character. Base name for the exported project files. Default "my_project".
n_starts	Number of multistart runs (number of initial value scenarios).
theta_intervals	Optional data frame specifying sampling intervals for initial values. Must contain columns NAME, lower, and upper. If NULL, intervals are derived from <code>interval_factor</code> .

interval_factor	Numeric vector of length 2 specifying multiplicative lower and upper bounds factors relative to original INIT values (default $c(0.2, 5)$).
vary_params	List of parameters which initial values are varied. Default value implies all parameters are varied. If theta_interval is specified, list of parameters is taken from theta_interval
seed	Optional random seed for reproducible sampling of initial values. If NULL, the global RNG state is not modified.

Details

For each start:

- Population parameter initial values INIT for estimated parameters are sampled uniformly.
- A scenario summary is stored.
- The corresponding .mlxtran project files are written.

If seed is provided, the function temporarily sets the random number generator state and restores it upon exit.

Value

The function writes two types of outputs to the specified path:

- A CSV file (scenarios_info.csv) summarizing all generated scenarios with columns:
 - scenario_number** Unique index of the scenario.
 - model** Model file or path used in the scenario.
 - data** Path to the dataset used.
 - theta** Active population parameters and initial values used.
 - RUV** Residual error structure(s) used.
 - RE** Random effect and correlation terms included.
 - OCC** Active occasion effects.
 - COVS** Included covariate-parameter relationships.
- Individual .mlxtran model files for each generated scenario, named according to the pattern <project_name>_<i>.mlxtran.

Examples

```
library(dplyr)
folder_path <- system.file("extdata", package = "SimuRg")

mod <- paste(folder_path, "/models/model_PK_1c.txt", sep = "/")

data <- paste(folder_path, "datasets", "dspk-warf.csv", sep = "/")
re <- list(init = tribble(~Cl, ~Vd, ~ka, ~Vp, ~Q,
                        1, 0, 0, 0, 0,
                        0, 1, 0, 0, 0,
                        0, 0, 1, 0, 0,
```

```

      0, 0, 0, 1, 0,
      0, 0, 0, 0, 1) %>% as.matrix(),
est = tribble(~Cl, ~Vd, ~ka, ~Vp, ~Q,
              TRUE, NA, NA, NA, NA,
              NA, TRUE, NA, NA, NA,
              NA, NA, TRUE, NA, NA,
              NA, NA, NA, TRUE, NA,
              NA, NA, NA, NA, TRUE) %>% as.matrix(),
block = tribble(~Cl, ~Vd, ~ka, ~Vp, ~Q,
               FALSE, NA, NA, NA, NA,
               NA, FALSE, NA, NA, NA,
               NA, NA, TRUE, NA, NA,
               NA, NA, NA, FALSE, NA,
               NA, NA, NA, NA, FALSE) %>% as.matrix())

headers <- list(list(name = "ID", use = "identifier", type = NULL),
               list(name = "TIME", use = "time", type = NULL),
               list(name = "DV", use = "observation", type = "continuous"),
               list(name = "DVID", use = "observationtype", type = NULL),
               list(name = "ADM", use = "administration", type = NULL),
               list(name = "AMT", use = "amount", type = NULL),
               list(name = "EVID", use = "eventidentifier", type = NULL),
               list(name = "MDV", use = "missingdependentvariable", type = NULL),
               list(name = "AGE", use = "covariate", type = "continuous"),
               list(name = "AGE_centered", use = "covariate", type = "continuous"),
               list(name = "SEX", use = "covariate", type = "categorical"),
               list(name = "WEIGHT", use = "covariate", type = "continuous"),
               list(name = "BMI", use = "covariate", type = "continuous"))

theta <- tribble(~NAME, ~TRANS, ~INIT, ~LB, ~UB, ~EST,
                "Cl", "logNormal", 0.2, NA, NA, TRUE,
                "Vd", "logNormal", 20, NA, NA, TRUE,
                "ka", "logNormal", 0.2, NA, NA, TRUE)

theta_intervals <- list(
  Cl = c(0.25*theta$INIT[theta$NAME == "Cl"], 4*theta$INIT[theta$NAME == "Cl"]),
  #Vd = c(0.5*theta$INIT[theta$NAME == "Vd"], 2*theta$INIT[theta$NAME == "Vd"]),
  ka = c(1.25*theta$INIT[theta$NAME == "ka"], 1.5*theta$INIT[theta$NAME == "ka"])
)

ruv <- list(YNAME = "y1",
            DVID = 1,
            TRANS = "normal",
            PRED = "Cc",
            ERR = "proportional",
            INIT = 1,
            EST = TRUE)

n_starts <- 20

path <- tempdir()
sg_multistart(
  mod = mod,

```

```

    data = data,
    headers = headers,
    ruv = ruv,
    theta = theta,
    re = re,
    occ = re,
    n_starts = n_starts,
    theta_intervals = theta_intervals,
    path = path,
    project_name = "multistart_test_project"
  )
  clr_files <- list.files(path, full.names = TRUE)
  unlink(clr_files, recursive = TRUE, force = TRUE)

```

sg_parsum

Extract parameter summary (theta, eta, SE, RSE, CI, CV, shrinkage)

Description

Extract parameter summary (theta, eta, SE, RSE, CI, CV, shrinkage)

Usage

```
sg_parsum(fpath_i, addOFV = TRUE)
```

Arguments

fpath_i	String or sg-fit object. If the string is given, the path to .Rdata or .json file with sg-fit object is expected.
addOFV	Logical. If TRUE, information about OFV and AIC of the model will be added. Default is TRUE.

Value

A table with parameter summary

Examples

```

fpath_i <- system.file("extdata", "simurg_object", "Warfarin_PK.RData", package = "SimuRg")
sum_tab <- sg_parsum(fpath_i)
print(sum_tab)

```

sg_predist_sim *Perform simulations for prediction distribution plots*

Description

Perform simulations for prediction distribution plots

Usage

```
sg_predist_sim(fpath_i, model, time_col = "TIME", outputs = NULL, npop = 500)
```

Arguments

fpath_i	String or sg-fit object. If the string is given, the path to .Rdata or .json file with sg-fit object is expected.
model	A model object passed to sg_sim().
time_col	String. The column to use as a time column. Currently, can be only TIME. Default is TIME.
outputs	Vector of strings. Names of the model variables to output. If NULL, all variables returned. Default is NULL.
npop	Integer. Number of population replicates. Default is 1.

Value

A dataset with simulation results

Examples

```
library(rxode2)
fpath_i <- system.file("extdata", "simurg_object", "Warfarin_PK.RData", package = "SimuRg")
mod_fin <- rxode2({
  # Differential equations
  d/dt(Ad) = -ka * Ad
  d/dt(Ac) = ka * Ad - Cl/V * Ac
  # Concentration calculations
  Cc = Ac / V
})
sg_predist_sim(fpath_i, mod_fin, outputs = "Cc")
```

sg_sim

Perform simulations from an rxode2 model

Description

Runs simulations using an rxode2 model object with explicit R objects for parameters (theta), variance-covariance (thetamat), omega, and sigma.

Usage

```
sg_sim(
  model,
  et,
  stimes = NULL,
  outputs = NULL,
  theta = NULL,
  omega = NULL,
  sigma = NULL,
  thetamat = NULL,
  covs = NULL,
  npop = 1,
  nsub = 1,
  aggr = NULL,
  addcov = TRUE,
  keep = NULL,
  scale = NULL,
  covint = "locf",
  inits = NULL,
  byID = NULL,
  byPOP = NULL,
  shared = NULL,
  ncores = 1,
  atol = 1e-08,
  rtol = 1e-06,
  ...
)
```

Arguments

model	A model object passed to <code>sg_sim()</code> .
et	An event table passed to <code>sg_sim()</code> .
stimes	A numeric vector of simulation times.
outputs	Vector of strings. Names of the model variables to output. If NULL, all variables returned. Default is NULL.
theta	A named numeric vector of baseline parameters. Default is NULL. Parameters listed in <code>params</code> are replaced by sampled values.

omega	A named matrix or vector.
sigma	A named matrix representing a sigma covariance matrix or its Cholesky decomposition; Default is NULL.
thetamat	A named variance-covariance matrix (for parameters brought to normal distribution). Default is NULL.
covs	A list of covariate structures. Each element should be a list containing covariate relationship definitions with the following structure: <ul style="list-style-type: none"> • PAR - string, name of the parameter to which covariate effect is applied • COVNAME - string, name of the covariate column in the dataset • FUNC - string, functional form of the covariate effect • TRANS - string, transformation applied to covariate • REF - numeric, reference value for categorical covariates • INIT - numeric, initial value for the covariate effect parameter • EST - logical, whether to estimate the covariate effect Can be NULL, if no covariates are used. Default is NULL
npop	Integer. Number of population replicates. Default is 1.
nsub	Integer. Number of subjects sampled per population (omega/sigma matrices per ID). Default is 1.
aggr	A string that specifies the aggregation type. Set to NULL for no aggregation, ID for aggregation by time, population and ID, NPOP for aggregation by population and time, TIME for aggregation by time. Default is NULL.
addcov	Logical. If TRUE, columns with covariate values will be added to the resulting dataset. Default is TRUE.
keep	Vector of strings. Columns of event table to keep in the output data frame. Default is NULL.
scale	A numeric named vector. Scaling for ODE parameters of the system. The names must correspond to the parameter identifiers in the ODE specification. Each of the ODE variables will be divided by the scaling factor. Default is NULL.
covint	String. Specifies the interpolation method for time-varying covariates. When solving ODEs it often samples times outside the sampling time specified in event table. When this happens, the time varying covariates are interpolated. Currently this can be: <ul style="list-style-type: none"> • "linear" interpolation, which interpolates the covariate by solving the line between the observed covariates and extrapolating the new covariate value • "locf" last observation carried forward • "NOCB" next observation carried backward. This is the same method that NONMEM uses • "midpoint" last observation carried forward to midpoint; next observation carried backward to midpoint Default is "locf"
inits	A named vector specifying initial conditions for model variables; Default is NULL.

byID	logical. If TRUE, replicate populations/subjects per event-table ID. Default NULL is treated as TRUE.
byPOP	logical. When both thetam and omega are used: if TRUE, replicate subjects by population. Default NULL is treated as TRUE.
shared	logical. If TRUE, one shared population draw for all IDs. Default NULL is treated as TRUE.
ncores	Integer. Number of cores used for calculations. Default is 1.
atol	A numeric absolute tolerance used by the ODE solver to determine if a good solution has been achieved. This is also used in the solved linear model to check if prior doses do not add anything to the solution. Default is 1e-8.
rtol	Numeric. A numeric relative tolerance used by the ODE solver to determine if a good solution has been achieved. This is also used in the solved linear model to check if prior doses do not add anything to the solution. Default is 1e-6.
...	optional arguments passed to rxode2::rxSolve (e.g. method, maxsteps).

Details

Index columns ID, POPN, and sim.id are included depending on what is used:

No uncertainty, no variability Only ID (and TIME, VAR, VALUE)

Only uncertainty (thetamat) POPN and ID

Only variability (omega) sim.id and ID

Uncertainty and variability POPN, sim.id, and ID

Other columns:

TIME Timepoint of the simulations

VAR Names of the output variables

VALUE Optional. Simulated value. Returned when no aggregation applied

mean, median, ... Optional. Aggregated statistic of simulated values. Returned when aggregation is applied

COV1, ...COVn Optional. Covariates value. Returned when addcov is TRUE

KEEP1, ...KEEPn Optional. Columns that were specified in the keep argument

Event table can use either id or ID. Parameters can be fixed per ID by including parameter names (e.g. *_pop, omega_*) as columns in the event table.

Value

A dataset with simulation results (long format).

Examples

```

library(rxode2)
library(dplyr)
library(tibble)
# Base 1-compartment PK model (no covariate)
mod <- rxode2::rxode2({
  ka_pop = 0.1;
  Vd_pop = 10;
  CL_pop = 0.5;

  omega_ka = 0;
  omega_Vd = 0;
  omega_CL = 0;

  Cc_b = 0;
  ka_tv = exp(ka_pop);
  Vd_tv = exp(Vd_pop);
  CL_tv = exp(CL_pop);

  ka = ka_tv * exp(omega_ka);
  Vd = Vd_tv * exp(omega_Vd);
  CL = CL_tv * exp(omega_CL);

  Cc = Ac / Vd;

  Ad(0) = 0;
  Ac(0) = 0;

  d/dt(Ad) = -ka * Ad;
  d/dt(Ac) = ka * Ad - CL * Cc;

  Cc_ResErr = Cc * (1 + Cc_b);
})

# Model with covariate: WTBL on Vd (Vd_tv = exp(Vd_pop) * (WTBL/70)^beta_WTBL_Vd_pop)
mod_cov <- rxode2::rxode2({
  ka_pop = 0.1;
  Vd_pop = 10;
  CL_pop = 0.5;

  beta_WTBL_Vd_pop = 0;

  omega_ka = 0;
  omega_Vd = 0;
  omega_CL = 0;

  Cc_b = 0;

  ka_tv = exp(ka_pop);
  Vd_tv = exp(Vd_pop) * (WTBL/70)^beta_WTBL_Vd_pop;
  CL_tv = exp(CL_pop);

```

```

ka = ka_tv * exp(omega_ka);
Vd = Vd_tv * exp(omega_Vd);
CL = CL_tv * exp(omega_CL);

Cc = Ac / Vd;

Ad(0) = 0;
Ac(0) = 0;

d/dt(Ad) = -ka * Ad;
d/dt(Ac) = ka * Ad - CL * Cc;

Cc_ResErr = Cc * (1 + Cc_b);
})

et_test <- tibble::tribble(
  ~ID, ~TIME, ~EVID, ~CMT, ~AMT,
  1, 0, 1, 1, 10,
  2, 0, 1, 1, 50
)

stimes_test <- seq(0, 24, 0.1)
output_test <- c("Ac", "Ad", "Cc", "Cc_ResErr")
theta_test <- c(ka_pop = log(0.1), Vd_pop = log(10), CL_pop = log(0.5))

thetamat_test <- matrix(c(0.05, 0.01, 0, 0.01, 0.05, 0, 0, 0, 0.05), nrow = 3, byrow = TRUE)
rownames(thetamat_test) <- colnames(thetamat_test) <- c("ka_pop", "Vd_pop", "CL_pop")

omega_test <- matrix(c(0.2, 0.05, 0, 0.05, 0.2, 0, 0, 0, 0.2), nrow = 3, byrow = TRUE)
rownames(omega_test) <- colnames(omega_test) <- c("omega_ka", "omega_Vd", "omega_CL")

sigma_test <- matrix(0.1); rownames(sigma_test) <- colnames(sigma_test) <- "Cc_b"

# No variability
sim1 <- sg_sim(model = mod, et = et_test, stimes = stimes_test, theta = theta_test,
              outputs = output_test)

# Population uncertainty, single scenario (ID)
sim2a <- sg_sim(model = mod, et = dplyr::filter(et_test, ID == 1),
               stimes = stimes_test, theta = theta_test, thetamat = thetamat_test,
               npop = 10)

# Population uncertainty, multiple IDs, byID = TRUE (populations are replicated
# for each scenario (ID)),
# shared = TRUE (the same populations are used for each scenario (ID))
sim2b <- sg_sim(model = mod, et = et_test, stimes = stimes_test, theta = theta_test,
               thetamat = thetamat_test, npop = 10, byID = TRUE, shared = TRUE)

# Population uncertainty, multiple IDs, byID = FALSE (one solve over full event table;
# npop must equal n(ID); ID - virtual subject)
sim2c <- sg_sim(model = mod, et = et_test, stimes = stimes_test, theta = theta_test,
               thetamat = thetamat_test, npop = nrow(et_test), byID = FALSE)

```

```

# Between-subject variability (BSV), multiple IDs, byID = TRUE (subjects are
# replicated for each scenario (ID)), shared = FALSE (separate subjects per ID)
sim3a <- sg_sim(model = mod, et = et_test, stimes = stimes_test, theta = theta_test,
               omega = omega_test, nsub = 10, byID = TRUE, shared = FALSE)

# Between-subject variability (BSV), byID = FALSE (one solve over full event table;
# nsub must equal n(ID); ID - virtual subject)
sim3b <- sg_sim(model = mod, et = et_test, stimes = stimes_test, theta = theta_test,
               omega = omega_test, nsub = nrow(et_test), byID = FALSE)

# BSV + uncertainty: byID = TRUE (populations/subjects are replicated for each
# scenario (ID)), byPOP = TRUE (subjects are replicated for each population),
# shared = FALSE (separate populations/subjects per ID)
sim4a <- sg_sim(model = mod, et = et_test, stimes = stimes_test, theta = theta_test,
               thetammat = thetammat_test, npop = 10, omega = omega_test, nsub = 5,
               byID = TRUE, byPOP = TRUE, shared = FALSE)

# BSV + uncertainty: byID = TRUE (populations/subjects are replicated for each
# scenario (ID)), byPOP = FALSE (subjects are not replicated between population;
# npop = nsub)
sim4b <- sg_sim(model = mod, et = et_test, stimes = stimes_test, theta = theta_test,
               thetammat = thetammat_test, npop = nrow(et_test), omega = omega_test,
               nsub = nrow(et_test), byID = TRUE, byPOP = FALSE)

# BSV + uncertainty: byID = FALSE (one solve over full event table; nsub and/or
# npop must equal n(ID); ID - virtual subject), byPOP = TRUE (subjects are
# replicated for each population), shared = FALSE (separate populations/subjects
# per ID)
sim4c <- sg_sim(model = mod, et = et_test, stimes = stimes_test, theta = theta_test,
               thetammat = thetammat_test, npop = 10, omega = omega_test, nsub = 5,
               byID = FALSE, byPOP = TRUE, shared = FALSE)

# BSV + uncertainty: byID = FALSE (one solve over full event table; nsub and/or
# npop must equal n(ID); ID - virtual subject), byPOP = FALSE (subjects are not
# replicated between population; npop = nsub)
sim4d <- sg_sim(model = mod, et = et_test, stimes = stimes_test, theta = theta_test,
               thetammat = thetammat_test, npop = nrow(et_test), omega = omega_test,
               nsub = nrow(et_test), byID = FALSE, byPOP = FALSE)

# Parameter override in event table
sim4 <- sg_sim(model = mod, et = dplyr::filter(et_test, ID == 1) %>%
               dplyr::mutate(Vd_pop = 2),
               stimes = stimes_test, theta = theta_test, thetammat = thetammat_test,
               npop = 1)

# Covariate (WTBL): pass covs and keep so WTBL is used and returned
theta_cov <- c(theta_test, beta_WTBL_Vd_pop = 0.5)
sim5 <- sg_sim(model = mod_cov, et = dplyr::filter(et_test, ID == 1) %>%

```

```

      dplyr::mutate(WTBL = 70),
    stimes = stimes_test, theta = theta_cov, covs = c("WTBL"),
    keep = c("WTBL"),
    thetamat = thetamat_test, npop = 5)

```

 sg_sim_tp

Time-profile plotting with optional summary bands

Description

Creates a ggplot time-profile plot from longitudinal data with optional summarization (mean/median/geometric mean), variance bands (SD/SE/IQR), percentile ribbons, faceting, and log scaling.

Usage

```

sg_sim_tp(
  ds_i,
  time_col = "TIME",
  val_col = "VALUE",
  group_i = "VAR",
  bands_i = NULL,
  cent_i = NULL,
  vrns_i = NULL,
  lperc_i = NULL,
  uperc_i = NULL,
  add_points = 0,
  col_i = NULL,
  fill_i = NULL,
  lty_i = NULL,
  shp_i = NULL,
  grid_i = NULL,
  wrap_i = NULL,
  free_stat = "free",
  wrap_ncol = NULL,
  wrap_nrow = NULL,
  min_x = NA,
  max_x = NA,
  min_y = NA,
  max_y = NA,
  log_y = FALSE,
  log_x = FALSE
)

```

Arguments

ds_i Data.frame. The data frame with source data.

time_col	String. The column to use as a time column. Currently, can be only TIME. Default is TIME.
val_col	String. Name of value column. Default is VALUE.
group_i	String. Primary grouping variable for lines. Default is 'VAR'.
bands_i	vector of characters. Character vector of length 2 with column names for custom ymin/ymax ribbons. Default is NULL
cent_i	string. Central tendency measure. One of 'mean', 'median', 'geom_mean', or NULL for raw values. Default is NULL
vrns_i	string. Variance band type. One of 'SD', 'SE', 'IQR', or NULL. Default is NULL
lperc_i	numeric. Lower percentile for percentile ribbons (must be provided together with uperc_i). Default is NULL
uperc_i	numeric. Upper percentile for percentile ribbons (must be provided together with lperc_i). Default is NULL
add_points	numeric. Size of points to add to the plot. If > 0, points will be added. Default is 0
col_i	String. Column name for color
fill_i	String. Column name for fill aesthetic. Default is NULL
lty_i	String. Column name for linetype aesthetic. Default is NULL.
shp_i	String. Column name for shape aesthetic. Default is NULL.
grid_i	string. Faceting formula for facet_grid (e.g., '~VAR' or 'A~B'). Default is NULL
wrap_i	String. Faceting formula for facet_wrap. Default is NULL.
free_stat	String. Facet scaling option. One of "free", "free_x", "free_y", or "fixed". Default is 'free'.
wrap_ncol	Integer. Number of columns for facet_wrap. Default is NULL.
wrap_nrow	Integer. Number of rows for facet_wrap. Default is NULL.
min_x	Numeric. X-axis minimum limit. Default is NA.
max_x	Numeric. X-axis maximum limit. Default is NA.
min_y	Numeric. Y-axis minimum limit. Default is NA.
max_y	Numeric. Y-axis maximum limit. Default is NA.
log_y	Logical. If TRUE, a logarithmic scale is applied to y-axis. Default is FALSE.
log_x	Logical. If TRUE, a logarithmic scale is applied to x-axis. Default is FALSE.

Value

A ggplot object with lines and optional ribbons/points/facets.

Examples

```

make_extended_mock_data <- function() {
  data.frame(
    TIME = rep(1:4, times = 6),
    VALUE = rnorm(24, mean = 10, sd = 2),
    VAR = rep(rep(c("A", "B"), each = 4), times = 3),
    Regimen = rep(c("R1", "R2", "R3"), each = 8),
    CAT1 = rep(c("Cat1", "Cat2"), each = 12)
  )
}
ds_sim <- make_extended_mock_data()
p <- sg_sim_tp(ds_i = ds_sim, group_i = 'VAR', col_i = 'VAR', fill_i = 'VAR',
              wrap_i = '~VAR', wrap_ncol = 2)

```

sg_translator

*Translate structural models between rxode2 and MLXTRAN syntax***Description**

Converts structural pharmacometric model files between rxode2 (R package) and MLXTRAN (Monolix) syntax. Supports ODE-based models and Monolix pkmodel macros.

Usage

```

sg_translator(
  input_path,
  to,
  output_path,
  dm_list = NULL,
  regressors = NULL,
  output_vars = NULL,
  macros = TRUE,
  stiff = TRUE
)

```

Arguments

input_path	Character string. Path to the input structural model file (.txt).
to	Character string. Target format: "mlxtran" or "rxode".
output_path	Character string. Path to write the translated model file (.txt).
dm_list	List of dosing macros with cmt and adm vectors. For example, list(cmt = 2, adm = 1) or list(cmt = c(2, 1), adm = c(1, 2)). Used to generate iv() dosing macros in MLXTRAN output.
regressors	Character vector of regressor variable names. For example, c("WTBL", "ADAN") generates WTBL = {use = regressor} in MLXTRAN.

output_vars	Character vector of output variable names for the MLXTRAN OUTPUT: section. For example, c("Cc_mgL", "CSF1_ngmL") generates output = {Cc_mgL, CSF1_ngmL}. Required when to = "mlxtran"; ignored when to = "rxode".
macros	Logical. If TRUE, attempt to convert rxode2 model to pkmodel macro format in MLXTRAN (only for simple 1- or 2-compartment oral PK models). Default TRUE.
stiff	Logical. If TRUE, add odeType = stiff to the MLXTRAN EQUATION section. Default TRUE.

Details

The function detects the source format automatically from file content.

rxode2 format uses # [INPUT], # [MODEL], # [OUTPUT] section markers, d/dt(X) ODE notation, f(X) for bioavailability, and a_{lag}(X) for lag time.

MLXTRAN format uses [LONGITUDINAL], PK:/EQUATION:/OUTPUT: sections, ddt_X ODE notation, compartment()/iv() dosing macros, and optionally pkmodel() macros.

When converting rxode2 to MLXTRAN with macros = TRUE, the function checks if the model structure matches a simple pkmodel() pattern (1- or 2-compartment with first-order oral absorption). If not, it falls back to full ODE format with compartment()/iv() macros.

MLXTRAN models using absorption()/elimination()/peripheral() macros (type 2) are expanded to explicit ODEs when converted to rxode2.

Value

Invisibly returns the translated model as a character string. The translated model is also written to output_path.

Examples

```
# Convert rxode2 model to MLXTRAN ODE format
rxode_model_hv_iv <- system.file("extdata", "models", "rxode",
                               "model_PK_hv_iv.txt", package = "SimuRg")
rxode_model_1c <- system.file("extdata", "models", "rxode",
                              "model_PK_1c.txt", package = "SimuRg")
monolix_model_hv_iv <- system.file("extdata", "models", "monolix",
                                  "model_PK_hv_iv.txt", package = "SimuRg")
monolix_model_1c <- system.file("extdata", "models", "monolix",
                                "model_PK_1c.txt", package = "SimuRg")

path_to_save <- tempdir()

sg_translator(rxode_model_hv_iv, to = "mlxtran",
              output_path = normalizePath(file.path(path_to_save,
                                                    "model_PK_hv_iv_mlx.txt")),
              mustWork = FALSE),
              output_vars = "Cc_mgL",
              dm_list = list(cmt = 1, adm = 1), stiff = TRUE)

# Convert rxode2 model to MLXTRAN with pkmodel macros
```

```

sg_translator(rxode_model_1c, to = "mlxtran",
              output_path = normalizePath(file.path(path_to_save,
                                                    "model_PK_1c_mlx.txt"),
                                          mustWork = FALSE),
              output_vars = "Cc_nM", macros = TRUE)

# Convert MLXTRAN model to rxode2
sg_translator(monolix_model_hv_iv, to = "rxode",
              output_path = normalizePath(file.path(path_to_save,
                                                    "model_PK_hv_iv_rx.txt"),
                                          mustWork = FALSE))

```

sg_vpc_sim

*Perform simulations for VPC plot***Description**

Perform simulations for VPC plot

Usage

```

sg_vpc_sim(
  fpath_i,
  gco = NULL,
  model = NULL,
  time_col = "TIME",
  outputs = NULL,
  npop = 100
)

```

Arguments

fpath_i	Either a character string specifying the file path to a saved <code>sg_fit</code> object (R data file), or a list object containing the <code>sg_fit</code> results directly. The object must contain: SDTAB, EVTAB, SUMTAB, OMEGAMAT, SIGMAMAT, and optionally COTAB and CATAB
gco	Generalized control object. Either an R list or path to Rdata/json file. Either <code>gco</code> or <code>model</code> file should be specified for simulations model specification. Defaults is NULL
model	A model object passed to <code>sg_sim()</code> .
time_col	String. The column to use as a time column. Currently, can be only TIME. Default is TIME.
outputs	Vector of strings. Names of the model variables to output. If NULL, all variables returned. Default is NULL.
npop	Integer specifying the number of virtual subjects to simulate per original individual. Higher values provide more robust percentile estimates but increase computation time. Default is 100

Value

A dataset with simulation results

Examples

```

library(rxode2)
library(tibble)
fpath_i <- system.file("extdata", "simurg_object", "Warfarin_PK.RData", package = "SimuRg")
# Simulate VPC from with generalized model object
mod <- rxode2::rxode2({
  ka_pop = 0.1;
  Vd_pop = 10;
  Cl_pop = 0.5;

  omega_ka = 0;
  omega_Vd = 0;
  omega_Cl = 0;

  Cc_b = 0;
  ka_tv = exp(ka_pop);
  Vd_tv = exp(Vd_pop);
  Cl_tv = exp(Cl_pop);

  ka = ka_tv * exp(omega_ka);
  Vd = Vd_tv * exp(omega_Vd);
  Cl = Cl_tv * exp(omega_Cl);

  Cc = Ac / Vd;

  Ad(0) = 0;
  Ac(0) = 0;

  d/dt(Ad) = -ka * Ad;
  d/dt(Ac) = ka * Ad - Cl * Cc;

  Cc_ResErr = Cc * (1 + Cc_b);
})

sg_vpc_sim(fpath_i, model=mod, outputs = "Cc_ResErr")
# Make VPC plot with the generalized control object
model <- system.file("extdata", "models", "rxode", "model_PK_1c.txt", package = "SimuRg")
data <- system.file("extdata", "datasets", "dspk-warf.csv", package = "SimuRg")
headers <- list(list(name = "ID", use = "identifier", type = NULL),
               list(name = "TIME", use = "time", type = NULL),
                 list(name = "DV", use = "observation", type = "continuous"),
                 list(name = "DVID", use = "observationtype", type = NULL),
                 list(name = "ADM", use = "administration", type = NULL),
                 list(name = "AMT", use = "amount", type = NULL),
                 list(name = "EVID", use = "eventidentifier", type = NULL),
                 list(name = "MDV", use = "missingdependentvariable", type = NULL),
                 list(name = "AGE", use = "covariate", type = "continuous"),
                 list(name = "AGE_centered", use = "covariate", type = "continuous"),

```

```

        list(name = "SEX", use = "covariate", type = "categorical"),
        list(name = "WEIGHT", use = "covariate", type = "continuous"),
        list(name = "BMI", use = "covariate", type = "continuous"))

theta <- tribble(~NAME, ~TRANS, ~INIT, ~LB, ~UB, ~EST,
                "Cl", "logNormal", 0.2, NA, NA, TRUE,
                "V", "logNormal", 20, NA, NA, TRUE,
                "ka", "logNormal", 0.2, NA, NA, TRUE
)

ruv <- list(YNAME = "y1", DVID = 1, TRANS = "normal", PRED = "Cc",
           ERR = "combined1", INIT = c(1, 1), EST = c(TRUE, TRUE), BLQM = NULL)

re <- list(init = tribble(~Cl, ~V, ~ka,
                        1, 0, 0,
                        0, 1, 0,
                        0, 0, 1) %>% as.matrix(),
          est = tribble(~Cl, ~V, ~ka,
                        TRUE, NA, NA,
                        NA, TRUE, NA,
                        NA, NA, TRUE) %>% as.matrix())

occ <- list(init = tribble(~Cl, ~V, ~ka,
                          0, 0, 0,
                          0, 0, 0,
                          0, 0, 0) %>% as.matrix(),
          est = tribble(~Cl, ~V, ~ka,
                        NA, NA, NA,
                        NA, NA, NA,
                        NA, NA, NA) %>% as.matrix())

covs <- list(list(PAR = "V", COVNAME = "AGE", FUNC = "linear",
                 TRANS = "median", INIT = 1, EST = TRUE),
            list(PAR = "ka", COVNAME = "SEX", REF = 0, INIT = 1, EST = TRUE))

gco <- list(headers = headers, data = data, model = model, task_opt = "",
           covs = covs, project_name = "test-proj", theta = theta,
           ruv = ruv, re = re, occ = occ, modelText = "")

res <- sg_vpc_sim(fpath_i, gco = gco, output = "Cc")

```

sg_vpc_vis

Visual Predictive Check (VPC) Function

Description

Generates VPC plots to assess model performance by comparing observed data with prediction intervals derived from simulated data.

Usage

```
sg_vpc_vis(
  ds_sim,
  data_i,
  output_names,
  time_col = "TIME",
  dv_col = "DV",
  log_y = FALSE,
  piLow = 0.1,
  piUp = 0.9,
  ciLow = 0.025,
  ciUp = 0.975,
  pred.corr = FALSE,
  lab_y = "DV",
  lab_x = "Time, h",
  theor_perc = TRUE,
  theor_percCI = TRUE,
  emp_perc = TRUE,
  dt_obs_f1 = FALSE,
  legend_f1 = FALSE,
  n_bins = 10,
  method = "kmeans",
  interpolation = TRUE,
  strat_by_dose = NULL
)
```

Arguments

ds_sim	A data frame with simulated data containing columns: <ul style="list-style-type: none"> • ID: simulation replicate identifier • TIME: time points • VAR: output variable name • VALUE: simulated values
data_i	A data frame with observed data containing columns: <ul style="list-style-type: none"> • ID: patient identifier • TIME: time points • DV: observed dependent variable values • DVID: dependent variable identifier
output_names	A data frame mapping VAR values (from ds_sim) to DVID values (from data_i). Must contain columns: <ul style="list-style-type: none"> • output: character, VAR values (e.g., "Cc", "Cp") • dvid: numeric, corresponding DVIDs (e.g., 1, 2)
time_col	String. The column to use as a time column. Currently, can be only TIME. Default is TIME.
dv_col	Character. Name of DV column in data_i. Default is DV

log_y	Logical. If TRUE, a logarithmic scale is applied to y-axis. Default is FALSE.
piLow	Numeric. Lower prediction interval bound. Default is 0.10.
piUp	Numeric. Upper prediction interval bound. Default is 0.90.
ciLow	Numeric. Lower confidence interval bound. Default is 0.025
ciUp	Numeric. Upper confidence interval bound. Default is 0.975
pred.corr	Logical. Apply prediction correction. Default is FALSE.
lab_y	String. Y-axis label. Default is "DV".
lab_x	String. X-axis label. Default is "TIME, h".
theor_perc	Logical. Show theoretical percentiles. Default is TRUE.
theor_percCI	Logical. Show CI around theoretical percentiles. Default is TRUE.
emp_perc	Logical. Show empirical percentiles. Default is TRUE
dt_obs_fl	Logical. Show observed data points. Default is FALSE
legend_fl	Logical. Show legend. Default is FALSE.
n_bins	Integer. Number of bins to use in the histogram. Default is 10.
method	Character. Binning method: "kmeans", "equal", "quantile" (default: "kmeans").
interpolation	Logical. Use line interpolation vs rectangles (default: FALSE).
strat_by_dose	Character. Variable name for dose stratification (default: NULL).

Details

For now, the model in the example is NOT the generalized model object (GMO), as the parameters in generalized fit object are backtransformed, and, therefore, not transformed in the model. This issue will be fixed in the next versions of SimuRg package

Value

List of ggplot objects, one for each output variable

Examples

```
library(rxode2)
fpath_i <- system.file("extdata", "simurg_object", "Warfarin_PK.RData", package = "SimuRg")
mod <- rxode2::rxode2({
  ka_pop = 0.1;
  Vd_pop = 10;
  Cl_pop = 0.5;

  omega_ka = 0;
  omega_Vd = 0;
  omega_Cl = 0;

  Cc_b = 0;
  ka_tv = exp(log(ka_pop));
  Vd_tv = exp(log(Vd_pop));
  Cl_tv = exp(log(Cl_pop));
```

```

ka = ka_tv * exp(omega_ka);
Vd = Vd_tv * exp(omega_Vd);
Cl = Cl_tv * exp(omega_Cl);

Cc = Ac / Vd;

Ad(0) = 0;
Ac(0) = 0;

d/dt(Ad) = -ka * Ad;
d/dt(Ac) = ka * Ad - Cl * Cc;

Cc_ResErr = Cc * (1 + Cc_b);
})

sim_data <- sg_vpc_sim(fpath_i, model=mod, outputs = "Cc_ResErr")
outp_nms <- data.frame(dvid = 1, output = "Cc")
vpc_plots <- sg_vpc_vis(
  ds_sim = sim_data,
  data_i = warfarin,
  output_names = outp_nms,
  lab_x = "Time (hours)",
  lab_y = "Concentration (ng/mL)",
  n_bins = 8,
  pred.corr = TRUE
)

```

sg_vpop_est

Perform generation of synthetic datasets for an empirical distribution

Description

The function operates in two modes:

- **Fixed seed mode** (when seed is specified): Generates a single dataset using the provided seed
- **Search mode** (when seed = NA): Iteratively searches for nds datasets that meet the correlation difference threshold

Usage

```

sg_vpop_est(
  data_i,
  nobj = NA,
  id_col = NULL,
  minnumlev = 3,
  npop = 1,
  excl_col = NULL,

```

```

    seed = NA,
    seed_umap = 123,
    palette = NULL,
    diag_plots = FALSE,
    remove_duplicates = TRUE,
    noise_level = 0.1,
    nds = 1,
    tg_corr dif = 0.1
  )

```

Arguments

<code>data_i</code>	data frame. Input data frame containing the original dataset to be synthesized (required)
<code>nobj</code>	integer. Specify the exact number of rows to generate in the synthetic dataset. When provided, overrides <code>npop</code> (optional, default: NA)
<code>id_col</code>	Character string. Specify the name of the identifier column to exclude from synthesis. Default: NULL.
<code>minnumlev</code>	integer. Threshold; numeric variables with \leq <code>minnumlev</code> unique values are converted to factors (optional, default: 3)
<code>npop</code>	Integer. Number of population replicates. Default is 1.
<code>excl_col</code>	Character vector. Contains column names to exclude from synthesis. Default: NULL
<code>seed</code>	integer. Random seed for synthetic data generation. If provided (not NA), generates a single dataset with this seed (fixed seed mode). If NA, uses search mode to find <code>nds</code> datasets meeting correlation threshold (optional, default: NA)
<code>seed_umap</code>	integer. Random seed for UMAP algorithm reproducibility (optional, default: 123)
<code>palette</code>	character vector. Contains color codes (hex format) for custom plot color schemes. If provided, should contain at least 2 colors. Used for histograms, bar plots, and UMAP visualizations (optional, default: <code>c("#3a6eba", "#efdd3c", "#1a1866", "#f2b93b")</code>)
<code>diag_plots</code>	logical flag. If TRUE, generates diagnostic plots and UMAP visualizations (optional, default: TRUE)
<code>remove_duplicates</code>	logical flag. If TRUE, automatically removes exact duplicates between original and synthetic data by adding controlled noise (optional, default: TRUE)
<code>noise_level</code>	numeric. Proportion of standard deviation to use when adding noise to continuous variables for duplicate removal. E.g., 0.10 means 10% of SD (optional, default: 0.10)
<code>nds</code>	integer. Number of synthetic datasets to generate in search mode. Ignored in fixed seed mode (optional, default: 1)
<code>tg_corr dif</code>	numeric. Target maximum absolute correlation difference threshold for dataset selection in search mode. Ignored in fixed seed mode (optional, default: 0.1)

Value

A list of lists, where each element contains results for one generated dataset:

- `datagen` - Synthetic dataset returned by `synthpop::syn()` (data.frame)
- `seed` - Random seed used to generate this dataset (integer or NA)
- `exact_dupl_check` - Logical indicating if exact duplicates exist between original and synthetic data
- `dplot_umap` - ggplot object with combined UMAP visualization comparing original and synthetic data (or NULL if `diag_plots=FALSE`)
- `ks_test` - Tibble with Kolmogorov-Smirnov p-values and statuses for continuous variables (or NULL if no continuous variables)
- `jsd_res` - Weighted mean Jensen-Shannon divergence (JSD) value for categorical variables (numeric or NULL if no categorical variables)
- `corr_diff_mean` - Mean absolute difference between original and synthetic correlation matrices (numeric or NULL if no continuous variables)
- `corr_diff_max` - Maximum absolute difference between original and synthetic correlation matrices (numeric or NULL if no continuous variables)
- `dplot_corr_diff` - ggplot heatmap object showing correlation difference matrix (Synthetic - Original) (or NULL if `diag_plots=FALSE` or no continuous variables)
- `dplot_cont` - List of ggplot histograms for continuous variables (or NULL if `diag_plots=FALSE` or no continuous variables)
- `dplot_cat` - List of ggplot barplots for categorical variables (or NULL if `diag_plots=FALSE` or no categorical variables)

Examples

```
library(dplyr)

# Generate example dataset
data <- data.frame(
  id = 1:150,
  ALB = rnorm(150, mean = 3.5, sd = 0.5),
  ALT = rnorm(150, mean = 50, sd = 20),
  SEX = factor(sample(c("M", "F"), 150, replace = TRUE)),
  RACE = factor(sample(c("White", "Black", "Asian", "Other"), 150, replace = TRUE))
)

# EXAMPLE 1: Fixed seed mode - generate single dataset with specified seed
output_fixed <- sg_vpop_est(data_i = data,
  id_col = "id",
  seed = 123,          # Fixed seed mode
  seed_umap = 40,
  diag_plots = TRUE)

# Access the dataset and its diagnostics
print(head(output_fixed[[1]]$datagen, 10))
print(output_fixed[[1]]$ks_test)
```

```

print(output_fixed[[1]]$seed)          # Will be 123
print(output_fixed[[1]]$corr_diff_max)
print(output_fixed[[1]]$dplot_umap)

# EXAMPLE 2: Search mode - generate 3 datasets meeting correlation threshold
output_search <- sg_vpop_est(data_i = data,
                             id_col = "id",
                             seed = NA,          # Search mode (default)
                             seed_umap = 40,
                             diag_plots = TRUE,
                             nds = 3,          # Generate 3 datasets
                             tg_corr dif = 0.1) # Target correlation difference

# Compare metrics across all datasets
sapply(output_search, function(x) x$corr_diff_max)
sapply(output_search, function(x) x$jsd_res)
sapply(output_search, function(x) x$seed) # Different seeds for each dataset

```

warfarin

Warfarin PKPD dataset

Description

Generated dataset with warfarin PK/PD measurements and covariates for 100 patients.

Usage

```
warfarin
```

Format

warfarin:

A data frame with 1700 rows and 17 columns:

ID identifier of the individual

TIME time of the dose or observation record (nominal)

DV records the measurement data

DVID identifier for the observation type (to distinguish different types of observations, e.g PK and PD)

DVNAME name for the observation type (to distinguish different types of observations, e.g PK and PD)

CMT compartment of the event

ADM identifier for the type of dose

AMT dose amount

EVID identifier to indicate if the line is a dose-line or a response-line

MDV identifier to ignore the observation information of that line

AGE age of the individual

SEX sex of the individual

WEIGHT weight of the individual

BMI BMI of the individual

CLCR creatinine clearance of the individual

CYP2C9_gentyp CYP2C9 genotype of the individual

VKORC1_gentyp VKORC1 genotype of the individual

Index

* datasets

data_pbc, [2](#)
ds_covval, [3](#)
gfo4cov, [4](#)
parest, [5](#)
warfarin, [81](#)

data_pbc, [2](#)
ds_covval, [3](#)

gfo4cov, [4](#)

parest, [5](#)

read_smrg_obj, [10](#)

sg_converter, [6](#)
sg_covsens_sim, [7](#), [13](#), [14](#)
sg_covsens_vis, [12](#)
sg_dummy, [17](#)
sg_fit, [26](#)
sg_globalsens_sim, [32](#)
sg_globalsens_vis, [35](#)
sg_gof_obpr, [37](#)
sg_gof_par_cov, [39](#)
sg_gof_par_dist, [40](#)
sg_gof_res, [42](#)
sg_gof_res_dist, [44](#)
sg_gof_tp, [46](#)
sg_localsens_sim, [47](#)
sg_localsens_vis, [50](#)
sg_modbuild, [53](#)
sg_modcomp, [57](#)
sg_multistart, [57](#)
sg_parsum, [61](#)
sg_predist_sim, [62](#)
sg_sim, [10](#), [63](#)
sg_sim_tp, [69](#)
sg_translator, [71](#)
sg_vpc_sim, [73](#)
sg_vpc_vis, [75](#)

sg_vpop_est, [78](#)

warfarin, [81](#)