

Package ‘mrgsolve’

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Type Package

Title Simulate from ODE-Based Models

Version 1.6.1

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Description Fast simulation from ordinary differential equation (ODE) based models typically employed in quantitative pharmacology and systems biology.

License GPL (>= 2)

URL <https://mrgsolve.org/docs/>,
<https://github.com/metrumresearchgroup/mrgsolve>

BugReports <https://github.com/metrumresearchgroup/mrgsolve/issues>

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'class_ev.R' 'class_derived.R' 'class_mrgmod.R'
'class_mrgsims.R' 'Aaaa.R' 'annot.R' 'chain.R' 'class_build.R'

'class_evd.R' 'events.R' 'class_rx.R' 'compile.R'
 'custom-tol.R' 'data_set.R' 'datasets.R' 'env.R' 'funset.R'
 'handle_spec_block.R' 'idata_set.R' 'init.R' 'inven.R'
 'knobs.R' 'matlist.R' 'matrix.R' 'mcode.R' 'model_include.R'
 'modlib.R' 'modspec.R' 'mread.R' 'mrgindata.R' 'mrgsim_q.R'
 'mrgsims.R' 'mrgsolve.R' 'mwrite.R' 'nm-mode.R' 'nmxml.R'
 'param.R' 'print.R' 'r_to_cpp.R' 'realize_addl.R' 'relabel.R'
 'render.R' 'update.R' 'workflows.R'

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Contents

mrgsolve-package	4
aboutsolver	7
as.ev	8
as.list,mrgmod-method	9
as.list,mrgsims-method	11
as_bmat	11
as_data_set	13
as_deslist	14
blocks	15
BLOCK_PARSE	16
c,matlist-method	19
c,tgrid-method	19
carry_out	20
check_data_names	21
cmtn	22
code	23
collapse_matrix	23
collapse_omega	24
custom_tol	25
data_set	26
design	29

details	30
env_eval	31
env_get	31
env_ls	32
env_update	32
ev	33
evd	35
ev_assign	36
ev_days	37
ev_rep	38
ev_repeat	39
ev_rx	39
ev_seq	41
exdatasets	43
expand.idata	44
expand_observations	45
get_tol	46
idata_set	46
init	48
inventory	50
is.mrgmod	51
is.mrgsims	51
lctran	52
loadso	53
matrix_helpers	54
mcode	55
mcRNG	56
modlib	56
modlib_details	57
modlib_pk	58
modlib_pkpd	59
modlib_tmdd	60
modlib_viral	61
mread	62
mread_yaml	65
mrgsim	66
mrgsims_dplyr	70
mrgsims_modify	72
mrgsim_q	73
mrgsim_variants	75
mutate.ev	76
mwrite_cpp	76
mwrite_yaml	77
names,mrgmod-method	78
nmext	79
nmxml	80
numerics_only	82
obsaug	82

obsonly	83
omega	83
outvars	85
param	85
param_tags	87
PKMODEL	88
plot,batch_mrgsims,missing-method	90
plot_mrgsims	91
plot_sims	92
qsim	93
read_nmext	95
realize_addl	96
render	97
Req	98
reserved	99
reset_tol	100
revar	101
see	101
sigma	102
simargs	103
soloc	104
solversettings	105
summary.mrgmod	105
tscale	106
update	107
use_custom_tol	109
valid_data_set	110
valid_idata_set	111
within	112
zero_re	113
\$.ev-method	113
\$.mrgmod-method	114

Index**115**

mrgsolve-package

*mrgsolve: Simulate from ODE-Based Models***Description**

mrgsolve is an R package maintained under the auspices of Metrum Research Group that facilitates simulation from models based on systems of ordinary differential equations (ODE) that are typically employed for understanding pharmacokinetics, pharmacodynamics, and systems biology and pharmacology. mrgsolve consists of computer code written in the R and C++ languages, providing an interface to a C++ translation of the lsoda differential equation solver. See [aboutsolver](#) for more information.

Resources

- Main mrgsolve resource page: <https://mrgsolve.org>
- User guide: https://mrgsolve.org/user_guide/
- Package documentation and vignettes: <https://mrgsolve.org/docs/>

Package-wide options

- `mrgsolve.project`: sets the default project director (`mread()`)
- `mrgsolve.soloc`: sets the default package build directory (`mread()`)
- `mrgsolve_mread_quiet`: don't print messages during `mread()`
- `mrgsolve.update.strict`: this option has been deprecated; use the `strict` argument to `update()` instead

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See Also

Useful links:

- <https://mrgsolve.org/docs/>
- <https://github.com/metrumresearchgroup/mrgsolve>
- Report bugs at <https://github.com/metrumresearchgroup/mrgsolve/issues>

Examples

```
## example("mrgsolve")

mod <- mrgsolve::house(delta=0.1) %>% param(CL=0.5)

events <- ev(amt=1000, cmt=1, addl=5, ii=24)

events
```

```

mod

see(mod)

## Not run:
stime(mod)

## End(Not run)
param(mod)

init(mod)

out <- mod %>% ev(events) %>% mrgsim(end=168)

head(out)
tail(out)
dim(out)

plot(out, GUT+CP~.)

sims <- as.data.frame(out)

t72 <- dplyr::filter(sims, time==72)

str(t72)

idata <- data.frame(ID=c(1,2,3), CL=c(0.5,1,2),VC=12)
out <- mod %>% ev(events) %>% mrgsim(end=168, idata=idata, req="")
plot(out)

out <- mod %>% ev(events) %>% mrgsim(carry_out="amt,evid,cmt,CL")
head(out)

ev1 <- ev(amt=500, cmt=2,rate=10)
ev2 <- ev(amt=100, cmt=1, time=54, ii=8, addl=10)
events <- c(ev1+ev2)
events

out <- mod %>% ev(events) %>% mrgsim(end=180, req="")
plot(out)

## "Condensed" data set
data(extran1)
extran1

out <- mod %>% data_set(extran1) %>% mrgsim(end=200)

plot(out,CP~time|factor(ID))

## idata

```

```

data(exidata)

out <-
  mod %>%
    ev(amt=1000, cmt=1) %>%
    idata_set(exidata) %>%
    mrgsim(end=72)

plot(out, CP~., as="log10")

# Internal model library
## Not run:
mod <- mread("irm1", modlib())

mod

x <- mod %>% ev(amt=300, ii=12, addl=3) %>% mrgsim

## End(Not run)

```

aboutsolver

*About the lsoda differential equation solver used by mrgsolve***Description**

The differential equation solver is a C++ translation of DLSODA from ODEPACK. The C++ translation was created by Dilawar Singh and hosted here <https://github.com/dilawar/liblsoda-cxx/>. As we understand the history of the code, Heng Li was also involved in early versions of the code written in C. There was a potentially-related project hosted here <https://github.com/sdwfrost/liblsoda/>.

Details

The C++ translation by Dilawar Singh contains functions that appear to be based on BLAS and LAPACK routines. These functions have been renamed to be distinct from the respective BLAS and LAPACK function names. References are given in the section below.

History

The following history was recorded in the source code published by Dilawar Singh:

```

/*
* HISTORY:
* This is a CPP version of the LSODA library for integration into MOOSE
  somulator.
* The original was aquired from

```

```

* http://www.ccl.net/cca/software/SOURCES/C/kinetics2/index.shtml and modified
by
* Heng Li <lh3lh3@gmail.com>. Heng merged several C files into one and added a
* simpler interface. [Available
here](http://lh3lh3.users.sourceforge.net/download/lsoda.c)

* The original source code came with no license or copyright
* information. Heng Li released his modification under the MIT/X11 license. I
* maintain the same license. I have removed quite a lot of text/comments from
* this library. Please refer to the standard documentation.
*
* Contact: Dilawar Singh <dilawars@ncbs.res.in>
*/

```

References

1. LAPACK: <https://netlib.org/lapack/>
2. BLAS: <https://netlib.org/blas/>

as.ev

Coerce an object to class ev

Description

Use this function to convert a data frame to an event object.

Usage

```

as.ev(x, ...)

## S4 method for signature 'data.frame'
as.ev(x, keep_id = TRUE, clean = FALSE, ...)

## S4 method for signature 'ev'
as.ev(x, ...)

```

Arguments

x	an object to coerce.
...	not used.
keep_id	if TRUE, ID column is retained if it exists.
clean	if TRUE, only dosing or ID information is retained in the result.

Details

If CMT (or cmt) is missing from the input, it will be set to 1 in the event object.

If TIME (or time) is missing from the input, it will be set to 0 in the event object.

If EVID (or evid) is missing from the input, it will be set to 1 in the event object.

Value

An object with class `ev`.

Examples

```
data <- data.frame(AMT = 100)

as.ev(data)

as.ev(data, clean = TRUE)
```

`as.list, mrgmod-method` *Coerce a model object to list*

Description

Coerce a model object to list

Usage

```
## S4 method for signature 'mrgmod'
as.list(x, deep = FALSE, ...)
```

Arguments

<code>x</code>	a model object.
<code>deep</code>	if TRUE, extra information is returned in the output list (see Details).
<code>...</code>	not used.

Details

If `deep` is TRUE, then the values for `trans`, `advan`, and `mindt` are returned as well as a summary of internal model functions (with a call to `mrgsolve::funset()`).

Value

A named list containing formatted contents from `x`.

Slots

- `npar`: number of parameters
- `neq`: number of compartments or differential equations
- `pars`: names of model parameters
- `covariates`: names of parameters identified as covariates
- `cmt`: names of model compartments

- param: the parameter list
- init: initial condition list
- omega: \$OMEGA matrices, as a matlist object
- sigma: \$SIGMA matrices, as a matlist object
- fixed: named list of \$FIXED values
- model: model name
- project: model project directory
- soloc: directory where the model is being built
- sodll: complete path to the model shared object
- cfile: path for the model source code file
- shlib: list of compilation information
- start: simulation start time
- end: simulation end time
- delta: simulation time step
- add: additional simulation times
- capture: names of captured data items
- request: compartments requested upon simulation
- cmti: named indices for current output compartments
- capturei: named indices for current output capture
- random: names and labels of \$OMEGA and \$SIGMA
- code: model source code from cfile
- details: model details data frame
- nm_import: a character vector listing the names of nonmem output files that were read to import estimates from a completed nonmem run
- cpp_variables: a data frame listing variables internal to the model cpp file
- atol: see [solversettings](#)
- rtol: see [solversettings](#)
- ss_atol: absolute tolerance to use when advancing to PK steady state
- ss_rtol: relative tolerance to use when advancing to PK steady state
- custom_atol: absolute tolerances, one for each compartment
- custom_rtol: relative tolerances, one for each compartment
- maxsteps: see [solversettings](#)
- hmin: see [solversettings](#)
- hmax: see [solversettings](#)
- itol: 1 (use scalar values) or 4 (use customized values)
- itol_type: either "scalar" (itol = 1) or "custom" (itol = 4)
- envir: the model environment

- plugins: plugins invoked in the model
- digits: number of digits to request in simulated data
- tscale: multiplicative scalar for time in results only
- mindt: simulation output time below which there model will assume to have not advanced
- preclean: logical indicating to clean up compilation artifacts prior to compiling
- debug: print debugging information during simulation run
- verbose: print extra information during setup for model run

Examples

```
mod <- mrgsolve::house()
l <- as.list(mod)
```

as.list, mrgsims-method

Coerce an mrgsims object to list

Description

Coerce an mrgsims object to list

Usage

```
## S4 method for signature 'mrgsims'
as.list(x, ...)
```

Arguments

x	an mrgsims object.
...	not used.

as_bmat

Coerce R objects to block or diagonal matrices

Description

These are simple functions that may be helpful to create the matrix objects that mrgsolve expects. Functions are named based on whether they create a diagonal matrix (d), a block matrix (b), or a correlation matrix (c).

Usage

```
as_bmat(x, ...)  
  
## S4 method for signature 'list'  
as_bmat(x, ...)  
  
## S4 method for signature 'numeric'  
as_bmat(x, pat = "*", ...)  
  
## S4 method for signature 'data.frame'  
as_bmat(x, pat = "*", cols = NULL, ...)  
  
## S4 method for signature 'ANY'  
as_bmat(x, ...)  
  
as_dmat(x, ...)  
  
## S4 method for signature 'list'  
as_dmat(x, ...)  
  
## S4 method for signature 'ANY'  
as_dmat(x, ...)  
  
## S4 method for signature 'numeric'  
as_dmat(x, pat = "*", ...)  
  
## S4 method for signature 'data.frame'  
as_dmat(x, pat = "*", cols = NULL, ...)  
  
as_cmat(x, ...)
```

Arguments

x	data frame or list.
...	arguments passed to dmat() or cmat() .
pat	regular expression, character.
cols	column names to use instead of pat.

Details

Use `as_dmat()` to create a diagonal matrix, `as_bmat()` to create a block matrix, and `as_cmat()` to create a block matrix where off-diagonal elements are understood to be correlations rather than covariances. `as_cmat()` uses `as_bmat()` to form the matrix and then converts off-diagonal elements to covariances before returning.

The methods for `data.frame` will work down the rows of the data frame and make the appropriate matrix from the data in each row. The result is a list of matrices.

Value

A numeric matrix for list and numeric methods. For data.frames, a list of matrices are returned.

See Also

[bmat\(\)](#), [dmat\(\)](#), [cmat\(\)](#)

Examples

```
df <- data.frame(
  OMEGA1.1 = c(1,2),
  OMEGA2.1 = c(11,22),
  OMEGA2.2 = c(3,4),
  SIGMA1.1 = 1,
  FOO=-1
)

as_bmat(df, "OMEGA")
as_dmat(df, "SIGMA")
as_dmat(df[1,], "OMEGA")
```

as_data_set

Create a simulation data set from ev objects or data frames

Description

The goal is to take a series of event objects or data frames and combine them into a single data frame that can be passed to [data_set\(\)](#).

Usage

```
as_data_set(x, ...)
```

```
## S4 method for signature 'ev'
as_data_set(x, ...)
```

```
## S4 method for signature 'data.frame'
as_data_set(x, ...)
```

Arguments

x an ev object or data frame.

... additional ev objects or data frames.

Details

Each event object or data frame is added to the data frame as an ID or set of IDs that are distinct from the IDs in the other event objects. Note that including ID argument to the `ev()` call where `length(ID)` is greater than one will render that set of events for all of IDs that are requested.

When determining the case for output names, the case attribute for the first ev object passed will be used to set the case for the output data.frame. In the event x is a data frame, the case of special column names (like amt/AMT or cmt/CMT) in the first data frame will be assessed and the case in the output data frame will be determined based on the relative numbers of lower or upper names.

To get a data frame with one row (event) per ID, look at `expand.ev()`.

Value

A data frame suitable for passing into `data_set()`. The columns will appear in a standardized order.

See Also

`expand.ev()`, `expand.evd()`, `ev()`, `evd()`, `uctran()`, `lctran()`

Examples

```
a <- ev(amt = c(100,200), cmt=1, ID = seq(3))
b <- ev(amt = 300, time = 24, ID = seq(2))
c <- ev(amt = 1000, ii = 8, addl = 10, ID = seq(3))

as_data_set(a, b, c)

d <- evd(amt = 500)

as_data_set(d, a)

# Output will have upper case nmtran names
as_data_set(
  data.frame(AMT = 100, ID = 1:2),
  data.frame(amt = 200, rate = 5, cmt = 2)
)

# Instead of this, use expand.ev
as_data_set(ev(amt = 100), ev(amt = 200), ev(amt = 300))
```

as_deslist

Create a list of designs from a data frame

Description

Create a list of designs from a data frame

Usage

```
as_deslist(data, descol = "ID")
```

Arguments

data	input data set; see Details .
descol	character column name to be used for design groups.

Details

The input data set must have a column with the same name as the value of descol. Other column names should be start (the time of the first observation), end (the time of the last observation), delta (the time steps to take between start and end), and add (other, ad-hoc times). Note that add might be a list-column to get a vector of times for each time grid object.

Value

The function returns a list of tgrid objects, one for each unique value found in descol.

Examples

```
idata <- tibble::tibble(ID=1:4, end=seq(24,96,24), delta=6,
  add=list(c(122,124,135),c(111), c(99),c(88)))

idata <- dplyr::mutate(idata, GRP = ID %%2)

idata

l <- as_deslist(idata,"GRP")

l

lapply(l,stime)

lapply(as_deslist(idata, "ID"),stime)
```

blocks

Return the code blocks from a model specification file

Description

Return the code blocks from a model specification file

Usage

```
blocks(x, ...)

## S4 method for signature 'mrgmod'
blocks(x, ...)

## S4 method for signature 'character'
blocks(x, ...)
```

Arguments

```
x          model object or path to model specification file
...        passed along
```

Examples

```
mod <- mrgsolve::house()
mod %>% blocks
mod %>% blocks(PARAM, TABLE)
```

BLOCK_PARSE

Functions to parse code blocks

Description

Most of the basic blocks are listed in this help topic. But see also [PKMODEL\(\)](#) which has more-involved options and is documented separately.

Usage

```
PARAM(
  x,
  env,
  pos = 1,
  annotated = FALSE,
  object = NULL,
  as_object = FALSE,
  covariates = FALSE,
  input = FALSE,
  tag = NULL,
  ...
)

FIXED(x, env, pos = 1, annotated = FALSE, ...)

THETA(
```



```
    x,  
    env,  
    pos = 1,  
    annotated = FALSE,  
    object = NULL,  
    as_object = FALSE,  
    name = "THETA",  
    fill = NULL,  
    ...  
)  
  
INIT(x, env, pos = 1, annotated = FALSE, object = NULL, as_object = FALSE, ...)  
  
CMT(  
  x,  
  env,  
  pos = 1,  
  annotated = FALSE,  
  object = NULL,  
  as_object = FALSE,  
  number = NULL,  
  prefix = "A",  
  ...  
)  
  
CAPTURE(x, env, pos = 1, annotated = FALSE, etas = NULL, ...)  
  
HANDLEMATRIX(  
  x,  
  env,  
  pos = 1,  
  annotated = FALSE,  
  object = NULL,  
  as_object = FALSE,  
  name = "...",  
  type = NULL,  
  oclass = "",  
  prefix = "",  
  labels = NULL,  
  unlinked = FALSE,  
  ...  
)
```

Arguments

x	data
env	parse environment
pos	block position

annotated	logical
object	the name of an object in ENV
as_object	indicates that object code is being provided
covariates	logical; mark as covariates and potentially required data input
input	logical; mark as potentially required data input
tag	space or comma-separated user-defined tags for the parameter block
...	passed
name	block name
fill	deprecated; not used
number	number of compartments to create
prefix	a prefix to add to the label
etas	allows for block capture of ETAs in the simulated output; this should be R code that will get parsed and evaluated; the result should be an integer-like vector which identifies which ETAs will be captured.
type	internal use
oclass	internal use
labels	aliases to use for simulated ETA values
unlinked	internal use

Details

When using `object` or `as_object` populate the block contents, the following types are required

- `PARAM`: a named list
- `INIT` : a named list
- `THETA` : a numeric vector; names are ignored
- `CMT`: a character vector
- `OMEGA`: matrix; set rownames on the matrix to create ETA labels; setting rownames is the only way to specify labels when working through the `object` or `as_object` directives
- `SIGMA`: matrix; set rownames on the matrix to create EPS labels; setting rownames is the only way to specify labels when working through the `object` or `as_object` directives

See Also

[PKMODEL\(\)](#)

c,matlist-method	<i>Operations with matlist objects</i>
------------------	--

Description

Operations with matlist objects

Usage

```
## S4 method for signature 'matlist'
c(x, ..., recursive = FALSE)
```

Arguments

x	a matlist object.
...	other matlist objects.
recursive	not used.

c,tgrid-method	<i>Operations with tgrid objects</i>
----------------	--------------------------------------

Description

Operations with tgrid objects

Usage

```
## S4 method for signature 'tgrid'
c(x, ..., recursive = FALSE)

## S4 method for signature 'tgrids'
c(x, ..., recursive = FALSE)

## S4 method for signature 'tgrid,numeric'
e1 + e2

## S4 method for signature 'tgrid,numeric'
e1 * e2

## S4 method for signature 'tgrids,numeric'
e1 + e2

## S4 method for signature 'tgrids,numeric'
e1 * e2
```

Arguments

x	a tgrid object.
...	additional tgrid objects.
recursive	not used.
e1	tgrid or tgrids object
e2	numeric value

carry_out	<i>Select items to carry into simulated output</i>
-----------	--

Description

When items named in this function are found in the input data set (either `data_set()` or `idata_set()`), they are copied into the simulated output. Special items like `evid` or `amt` or the like are not copied from the data set per se, but they are copied from `datarecord` objects that are created during the simulation.

Usage

```
carry_out(x, ...)
```

```
carry.out(x, ...)
```

Arguments

x	model object.
...	unquoted names of data items to copy into the simulated output.

Details

There is also a `carry_out` argument to `mrgsim()` that can be set to accomplish the same thing as a call to `carry_out` in the pipeline.

`carry.out` and `carry_out` both do the same thing; using the underscore version is now preferred.

Examples

```
mod <- mrgsolve::house()

e <- ev(amt = 100, ii = 6, addl = 3, WT = 70, dose = amt)

out <- mod %>% ev(e) %>% carry_out(amt, dose, WT) %>% mrgsim()

head(out)
```

check_data_names*Check input data set names against model parameters*

Description

Use this function to check names of input data sets against parameters that have been assigned different tags. Assignment is made in the model specification file. This is useful to alert the user to misspelled or otherwise misspecified parameter names in input data sets. See [param_tags\(\)](#) for information on associating tags with parameters.

Usage

```
check_data_names(  
  data,  
  x,  
  check_covariates = TRUE,  
  check_inputs = TRUE,  
  tags = NULL,  
  mode = c("warn", "error", "inform"),  
  silent = FALSE  
)
```

Arguments

data	a data frame or other object with names to check.
x	a model object.
check_covariates	logical; if TRUE, check data for parameter names carrying the covariates tag.
check_inputs	logical; if TRUE, check data for parameter names carrying the input tag.
tags	a character vector of user-defined parameter tags to require in data; this may be a comma- or space-separated string (e.g. "tag1, tag2").
mode	the default is to "warn" the user when data is missing some expected column names; alternatively, use "error" to issue an error or "inform" to generate a message when data is missing some expected column names.
silent	silences message on successful check.

Details

By default, data will be checked for parameters with the covariates or input tags; these checks can be bypassed with the `check_covariates` and `check_inputs` arguments. When a parameter name is missing from data the user will be warned by default. Use `mode = "error"` to generate an error instead of a warning and use `mode = "inform"` to simply be informed. When the user has not tagged any parameters for checking, there will either be a warning (default) or an error (when `mode = "error"`).

It is an error to request a parameter tag via the `tags` argument when that tag is not found in the model.

It is an error to call `check_data_names` when no parameters have been tagged in the model specification file (see [param_tags\(\)](#)).

Value

A logical value is returned; TRUE if all expected parameters were found and FALSE otherwise.

See Also

[param_tags\(\)](#)

Examples

```
mod <- mcode("ex-cdn", "$PARAM @input \n CL = 1, KA = 2", compile = FALSE)

param(mod)

# Coding mistake!
data <- expand.evd(amt = 100, cl = 2, KA = 5)

check_data_names(data, mod)

try(check_data_names(data, mod, mode = "error"))

check_data_names(data, mod, mode = "inform")
```

cmtn

Get the compartment number from a compartment name

Description

Get the compartment number from a compartment name

Usage

```
cmtn(x, ...)

## S4 method for signature 'mrgmod'
cmtn(x, tag, ...)
```

Arguments

<code>x</code>	model object.
<code>...</code>	not used.
<code>tag</code>	compartment name.

Examples

```
mod <- mrgsolve::house()
cmtn(mod, "CENT")
```

code

*Extract the code from a model***Description**

This function is currently not exported, so be sure to call it with `mrgsolve:::code(...)`.

Usage

```
code(x)
```

Arguments

`x` a model object.

Value

A character vector of model code.

Examples

```
mod <- mrgsolve::house()
mrgsolve:::code(mod)

# Alternative
as.list(mod)$code
```

collapse_matrix

*Collapse the matrices of a matlist object***Description**

This function is called by `collapse_omega()` and `collapse_sigma()` to convert multiple matrix blocks into a single matrix block. This "collapsing" of the matrix list is irreversible.

Usage

```
collapse_matrix(x, range = NULL, name = NULL)
```

Arguments

x	an object that inherits from <code>matlist</code> ; this object is most frequently extracted from a model object using <code>omat()</code> or <code>smat()</code> for OMEGA and SIGMA, respectively.
range	numeric vector of length 2 specifying the range of matrices to collapse in case there are more than 2. The second element may be NA to indicate the length of the list of matrices.
name	a new name for the collapsed matrix; note that this is the matrix name, not the labels which alias ETA(n) or EPS(n); specifying a name will only alter how this matrix is potentially updated in the future.

Value

An update `matlist` object (either `omegalist` or `sigmalist`).

See Also

`collapse_omega()`, `collapse_sigma()`, `omat()`, `smat()`

Examples

```
omega <- omat(list(dmat(1, 2), dmat(3, 4, 5)))
omega
collapse_matrix(omega)
```

collapse_omega

Collapse OMEGA or SIGMA matrix lists

Description

If multiple OMEGA (or SIGMA) blocks were written into the model, these can be collapsed into a single matrix. This will not change the functionality of the model, but will alter how OMEGA (or SIGMA) are updated, usually making it easier. This "collapsing" of the matrix list is irreversible.

Usage

```
collapse_omega(x, range = NULL, name = NULL)
```

```
collapse_sigma(x, range = NULL, name = NULL)
```

Arguments

x	a model object.
range	numeric vector of length 2 specifying the range of matrices to collapse in case there are more than 2. The second element may be NA to indicate the length of the list of matrices.

name a new name for the collapsed matrix; note that this is the matrix name, not the labels which alias ETA(n) or EPS(n); specifying a name will only alter how this matrix is potentially updated in the future.

Value

A model object with updated OMEGA or SIGMA matrix lists.

See Also

[collapse_matrix\(\)](#)

Examples

```
code <- '
$OMEGA 1 2 3
$OMEGA 4 5
$OMEGA 6 7 8 9
'

mod <- mcode("collapse-example", code, compile = FALSE)
revar(mod)
collapse_omega(mod) %>% omat()
collapse_omega(mod, range = c(2,3), name = "new_matrix") %>% omat()
collapse_omega(mod, range = c(2,NA), name = "new_matrix") %>% omat()
```

custom_tol

Customize tolerances for specific compartments

Description

These functions update the relative or absolute tolerance values only for the custom tolerance configuration.

Usage

```
custom_tol(.x, .rtol = NULL, .atol = NULL)

custom_rtol(.x, .rtol = list(), .default = NULL, .use = TRUE, ...)

custom_atol(.x, .atol = list(), .default = NULL, .use = TRUE, ...)
```

Arguments

.x a model object.

.rtol a named numeric list or vector, where names reference selected model compartments and relative tolerances for those compartments; custom_tol() also accepts the name of an object in the model environment to be used.

<code>.atol</code>	a named numeric list or vector, where names reference selected model compartments and absolute tolerances for those compartments; <code>custom_tol()</code> also accepts the name of an object in the model environment to be used.
<code>.default</code>	the default tolerance value to use for compartments not listed in <code>.rtol</code> , <code>.atol</code> , or <code>...</code> ; if not supplied, the current scalar value in <code>.x</code> will be used.
<code>.use</code>	logical; if TRUE, then a call to <code>use_custom_tol()</code> will be made prior to return; if FALSE, a call to <code>use_scalar_tol()</code> will be made; under expected use, the value for this argument is kept TRUE, so that whenever tolerances are customized, they will be used in the next simulation run.
<code>...</code>	name/value pairs, where name references a model compartment and value is a new, numeric value to use for <code>rtol</code> or <code>atol</code> .

Details

New tolerance values can be supplied by either a named, numeric vector or list via `.rtol` and `.atol` or via `...` or by both. If duplicate compartment names are found in `...` and either `.rtol` or `.atol`, the value passed via `...` will take precedence.

The `custom_tol()` function provides a mechanism for coding customized tolerances into the model file itself. Simply create named numeric lists or vectors for customized `rtol` or `atol` in a `$ENV` block. On loading the model, call `custom_tol()` and supply the names of those objects as `.rtol` and `.atol`.

Value

An updated model object.

See Also

`reset_tol()`, `use_custom_tol()`, `use_scalar_tol()`, `get_tol()`

Examples

```
mod <- house()
mod <- custom_rtol(mod, GUT = 1e-2, CENT = 1e-3)

new_tolerances <- c(GUT = 1e-4, RESP = 1e-5)
mod <- custom_rtol(mod, new_tolerances, RESP = 1e-6)
```

data_set

Select and modify a data set for simulation

Description

The input data set (`data_set`) is a data frame that specifies observations, model events, and / or parameter values for a population of individuals.

Usage

```
data_set(x, data, ...)

## S4 method for signature 'mrgmod,data.frame'
data_set(
  x,
  data,
  .subset = TRUE,
  .select = TRUE,
  object = NULL,
  need = NULL,
  ...
)

## S4 method for signature 'mrgmod,ANY'
data_set(x, data, ...)

## S4 method for signature 'mrgmod,ev'
data_set(x, data, ...)

## S4 method for signature 'mrgmod,missing'
data_set(x, object, ...)
```

Arguments

x	a model object.
data	input data set as a data frame.
...	other arguments passed along when object is a function.
.subset	an unquoted expression passed to <code>dplyr::filter()</code> ; retain only certain rows in the data set
.select	passed to <code>dplyr::select()</code> ; retain only certain columns in the data set; this should be the result of a call to <code>dplyr::vars()</code> .
object	character name of an object existing in \$ENV to use for the data set.
need	passed to <code>inventory()</code> .

Details

Input data sets are R data frames that can include columns with any valid name, however columns with selected names are treated specially by mrgsolve and incorporated into the simulation.

ID specifies the subject ID and is required for every input data set.

When columns have the same name as parameters (\$PARAM or \$INPUT in the model specification file), the values in those columns will be used to update the corresponding parameter as the simulation progresses.

Input data set may include the following columns related to PK dosing events: TIME, CMT, AMT, RATE, II, ADDL, SS. Both ID and TIME are required columns in the input data set unless \$PRED is in use. Lower case PK dosing column names including time, cmt, amt, rate, ii, addl, ss are also

recognized. However, an error will be generated if a mix of both upper case and lower case columns in this family are found. Use the functions `lctran()` and `uctran()` to convert between upper and lower case naming for these data items.

TIME is the observation or event time, CMT is the compartment number (see `init()`), AMT is the dosing amount, RATE is the infusion rate, II is the dosing interval, ADDL specifies additional doses to administer, and ss is a flag indicating that the system should be advanced to a pharmacokinetic steady state prior to administering the dose. These column names operate similarly to other non-linear mixed effects modeling software.

EVID is an integer value specifying the ID of an event record. Values include:

- 0: observation
- 1: dose event, either bolus or infusion
- 2: other-type event; in mrgsolve, this functions like an observation record, but a discontinuity is created in the simulation at the time of the event (i.e., the ODE solver will stop and restart at the time of the event)
- 3: reset the system
- 4: reset the system and dose
- 8: replace the amount in a compartment

For all EVID greater than 0, a discontinuity is created in the simulation, as described for EVID 2.

An error will be generated when mrgsolve detects that the data set is not sorted by time within an individual. mrgsolve does **not** allow time to be reset to zero on records where EVID is set to 4 (reset and dose).

Only numeric data can be brought in to the problem. Any non-numeric data columns will be dropped with warning. See `numerics_only()`, which is used to prepare the data set.

An error will be generated if any parameter columns in the input data set contain missing values (NA). Likewise, an error will be generated if missing values are found in the following columns: ID, time/TIME, rate/RATE.

See [exdatasets](#) for several example data sets that are provided by mrgsolve.

See Also

`idata_set()`, `ev()`, `valid_data_set()`, `valid_idata_set()`, `lctran()`, `uctran()`.

Examples

```
mod <- mrgsolve::house()

data <- expand.ev(ID = seq(3), amt = c(10, 20))

mod %>% data_set(data, ID > 1) %>% mrgsim()

data(extran1)
head(extran1)

mod %>% data_set(extran1) %>% mrgsim()
mod %>% mrgsim(data = extran1)
```

design	<i>Set observation designs for the simulation</i>
--------	---

Description

This function also allows you to assign different designs to different groups or individuals in a population.

Usage

```
design(x, deslist = list(), descol = character(0), ...)
```

Arguments

x	model object
deslist	a list of tgrid or tgrids objects or numeric vector to be used in place of ...
descol	the idata column name (character) for design assignment
...	not used

Details

This setup requires the use of an `idata_set`, with individual-level data passed in one ID per row. For each ID, specify a grouping variable in `idata` (`descol`). For each unique value of the grouping variable, make one `tgrid` object and pass them in order as ... or form them into a list and pass as `deslist`.

You must assign the `idata_set` before assigning the designs in the command chain (see the example below).

Examples

```
peak <- tgrid(0,6,0.1)
sparse <- tgrid(0,24,6)

des1 <- c(peak,sparse)
des2 <- tgrid(0,72,4)

data <- expand.ev(ID = 1:10, amt=c(100,300))
data$GRP <- data$amt/100

idata <- data[,c("ID", "amt")]

mod <- mrgsolve::house()

mod %>%
  omat(dmat(1,1,1,1)) %>%
  carry_out(GRP) %>%
  idata_set(idata) %>%
```

```
design(list(des1, des2), "amt") %>%
data_set(data) %>%
mrgsim() %>%
plot(RES~time|GRP)
```

details	<i>Extract model details</i>
---------	------------------------------

Description

Extract model details

Usage

```
details(x, complete = FALSE, values = TRUE, ...)
```

Arguments

x	a model object
complete	logical; if TRUE, un-annotated parameters and compartments will be added to the output
values	logical; if TRUE, a values column will be added to the output
...	not used

Details

This function is not exported. You will have to call it with `mrgsolve:::details()`.

Examples

```
mod <- mrgsolve::house()

mrgsolve:::details(mod)
```

env_eval	<i>Re-evaluate the code in the ENV block</i>
----------	--

Description

The \$ENV block is a block of R code that can realize any sort of R object that might be used in running a model.

Usage

```
env_eval(x, seed = NULL)
```

Arguments

x	a model object.
seed	passed to set.seed() if a numeric value is supplied.

See Also

[env_get\(\)](#), [env_ls\(\)](#)

env_get	<i>Return model environment</i>
---------	---------------------------------

Description

Return model environment

Usage

```
env_get(x, tolist = TRUE)
```

```
env_get_env(x)
```

Arguments

x	a model object.
tolist	should the environment be coerced to list?

env_ls	<i>List objects in the model environment</i>
--------	--

Description

Each model keeps an internal environment that allows the user to carry any R object along. Objects are coded in \$ENV.

Usage

```
env_ls(x, ...)
```

Arguments

x	a model object.
...	passed to <code>ls()</code> .

env_update	<i>Update objects in model environment</i>
------------	--

Description

Update objects in model environment

Usage

```
env_update(.x, ..., .dots = list())
```

Arguments

.x	a model object.
...	objects to update.
.dots	list of objects to updated.

Description

An event object specifies dosing or other interventions that get implemented during simulation. Event objects do similar things as [data_set](#), but simpler and easier to create.

Usage

```
ev(x, ...)

## S4 method for signature 'mrgmod'
ev(x, object = NULL, ...)

## S4 method for signature 'missing'
ev(
  time = 0,
  amt = 0,
  evid = 1,
  cmt = 1,
  ID = numeric(0),
  replicate = TRUE,
  until = NULL,
  tinf = NULL,
  realize_addl = FALSE,
  ...
)

## S4 method for signature 'ev'
ev(x, realize_addl = FALSE, ...)
```

Arguments

x	a model object.
...	other items to be incorporated into the event object; see Details .
object	an event object to be added to a model object.
time	event time.
amt	dose amount.
evid	event ID.
cmt	compartment number or name.
ID	subject ID.
replicate	logical; if TRUE, events will be replicated for each individual in ID.
until	the expected maximum observation time for this regimen; doses will be scheduled up to, but not including, the until time; see Examples .

`tinf` infusion time; if greater than zero, then the rate item will be derived as `amt/tinf`.
`realize_addl` if FALSE (default), no change to `addl` doses. If TRUE, `addl` doses are made explicit with `realize_addl()`.

Details

- Required items in events objects include `time`, `amt`, `evid` and `cmt`.
- If not supplied, `evid` is assumed to be 1.
- If not supplied, `cmt` is assumed to be 1.
- If not supplied, `time` is assumed to be 0.
- If `amt` is not supplied, an error will be generated.
- If `total` is supplied, then `addl` will be set to `total-1`.
- Other items can include `ii`, `ss`, and `addl` (see [data_set](#) for details on all of these items).
- ID may be specified as a vector.
- If `replicate` is TRUE (default), then the events regimen is replicated for each ID; otherwise, the number of event rows must match the number of IDs entered.

Value

`ev()` returns an event object.

See Also

[evd\(\)](#), [ev_rep\(\)](#), [ev_days\(\)](#), [ev_repeat\(\)](#), [ev_assign\(\)](#), [ev_seq\(\)](#), [mutate.ev\(\)](#), [as.ev\(\)](#), [as.evd\(\)](#), [ev_methods](#).

Examples

```
mod <- mrgsolve::house()

mod <- mod %>% ev(amt = 1000, time = 0, cmt = 1)

loading <- ev(time = 0, cmt = 1, amt = 1000)

maint <- ev(time = 12, cmt = 1, amt = 500, ii = 12, addl = 10)

c(loading, maint)

reduced_load <- dplyr::mutate(loading, amt = 750)

# Three additional doses in this case
e <- ev(amt = 100, ii = 4*7, until = 16*7)
e
# Last dose is given at 84
realize_addl(e)

# Four additional doses with last at 112 in this case
e <- ev(amt = 100, ii = 4*7, until = 16*7 + 0.001)
realize_addl(e)
```

evd

*Create an event object with data-like names***Description**

This function calls `ev()` to create an event object and then sets the case attribute so that it renders nmtran data names in upper case. An object created with `evd()` can be used in the same way as an object created with `ev()`.

Usage

```
evd(x, ...)

## S4 method for signature 'mrgmod'
evd(x, ...)

## S4 method for signature 'missing'
evd(x, ...)

## S4 method for signature 'ev'
evd(x, ...)

as.evd(x)
```

Arguments

<code>x</code>	an event object.
<code>...</code>	arguments passed to <code>ev()</code> .

Details

Note that `evd` isn't a separate class; it is just an `ev` object with a specific case attribute. See examples which illustrate the difference.

See Also

`ev()`, `lctran()`, `uctran()`

Examples

```
a <- evd(amt = 100)
b <- ev(amt = 300)
a
as.data.frame(a)
as_data_set(a, b)
as_data_set(b, a)
as.data.frame(seq(a, b))
```

ev_assign

Replicate a list of events into a data set

Description

Replicate a list of events into a data set

Usage

```
ev_assign(l, idata, evgroup, join = FALSE)
```

```
assign_ev(...)
```

Arguments

l	list of event objects.
idata	an idata set (one ID per row).
evgroup	the character name of the column in idata that specifies event object to implement.
join	if TRUE, join idata to the data set before returning.
...	used to pass arguments from assign_ev(). to ev_assign().

Details

ev_assign() connects events in a list passed in as the l argument to values in the data set identified in the evgroup argument. For making assignments, the unique values in the evgroup column are first sorted so that the first sorted unique value in evgroup is assigned to the first event in l, the second sorted value in evgroup column is assigned to the second event in l, and so on. This is a change from previous behavior, which did not sort the unique values in evgroup prior to making the assignments.

Examples

```
ev1 <- ev(amt = 100)
ev2 <- ev(amt = 300, rate = 100, ii = 12, addl = 10)

idata <- data.frame(ID = seq(10))
idata$arm <- 1+(idata$ID %%2)

ev_assign(list(ev1, ev2), idata, "arm", join = TRUE)
```

ev_days

Schedule dosing events on days of the week

Description

This function lets you schedule doses on specific days of the week, allowing you to create dosing regimens on Monday/Wednesday/Friday, or Tuesday/Thursday, or every other day (however you want to define that) etc.

Usage

```
ev_days(  
  ev = NULL,  
  days = "",  
  addl = 0,  
  ii = 168,  
  unit = c("hours", "days"),  
  ...  
)
```

Arguments

ev	an event object.
days	comma- or space-separated character string of valid days of the the week (see details).
addl	additional doses to administer.
ii	inter-dose interval; intended use is to keep this at the default value.
unit	time unit; the function can only currently handle hours or days.
...	event objects named by one of the valid days of the week (see Details).

Details

Valid names of the week are:

- m for Monday
- t for Tuesday
- w for Wednesday
- th for Thursday
- f for Friday
- sa for Saturday
- s for Sunday

The whole purpose of this function is to schedule doses on specific days of the week, in a repeating weekly schedule. Please do use caution when changing `ii` from its default value.

Examples

```
# Monday, Wednesday, Friday x 4 weeks
e1 <- ev(amt = 100)
ev_days(e1, days="m,w,f", addl = 3)

# 50 mg Tuesdays, 100 mg Thursdays x 6 months
e2 <- ev(amt = 50)
ev_days(t = e2, th = e1, addl = 23)
```

ev_rep

*Replicate an event object***Description**

An event sequence can be replicated a certain number of times in a certain number of IDs.

Usage

```
ev_rep(x, ID = 1, n = NULL, wait = 0, as.ev = FALSE, id = NULL)
```

Arguments

x	event object.
ID	numeric vector if IDs.
n	passed to ev_repeat() .
wait	passed to ev_repeat() .
as.ev	if TRUE an event object is returned.
id	deprecated; use ID instead.

Value

A single data.frame or event object as determined by the value of [as.ev\(\)](#).

See Also

[ev_repeat\(\)](#)

Examples

```
e1 <- c(ev(amt=100), ev(amt=200, ii=24, addl=2, time=72))

ev_rep(e1, 1:5)
```

ev_repeat	<i>Repeat a block of dosing events</i>
-----------	--

Description

Repeat a block of dosing events

Usage

```
ev_repeat(x, n, wait = 0, as.ev = FALSE)
```

Arguments

x	event object or dosing data frame.
n	number of times to repeat.
wait	time to wait between repeats.
as.ev	if TRUE, an event object is returned; otherwise a data.frame is returned.

Value

See as.ev argument.

Examples

```
e1 <- ev(amt = 100, ii = 24, addl = 20)
e4 <- ev_repeat(e1, n = 4, wait = 168)
mod <- mrgsolve::house()
out <- mrgsim(mod, events = e4, end = 3200)
plot(out, "CP")
```

ev_rx	<i>Create intervention objects from Rx input</i>
-------	--

Description

See details below for Rx specification. Actual parsing is done by [parse_rx\(\)](#); this function can be used to debug Rx inputs.

Usage

```

ev_rx(x, y, ...)

## S4 method for signature 'mrgmod,character'
ev_rx(x, y, ...)

## S4 method for signature 'character,missing'
ev_rx(x, df = FALSE, ...)

parse_rx(x)

```

Arguments

x	a model object or character Rx input.
y	character Rx input; see details.
...	not used at this time.
df	if TRUE then a data frame is returned.

Value

The method dispatched on model object (mrgmod) returns another model object. The character method returns an event object. The parse_rx function return a list named with arguments for the event object constructor [ev\(\)](#).

Rx specification

- The dose is found at the start of the string by sequential digits; this may be integer, decimal, or specified in scientific notation
- Use in to identify the dosing compartment number; must be integer
- Use q to identify the dosing interval; must be integer or decimal number (but not scientific notation)
- Use over to indicate an infusion and its duration; integer or decimal number
- Use x to indicate total number of doses; must be integer
- Use then or , to separate dosing periods
- Use after to insert a lag in the start of a period; integer or decimal number (but not scientific notation)
- Use & to implement multiple doses at the same time

Examples

```

# example("ev_rx")

ev_rx("100")

ev_rx("100 in 2")

```



```

ev_rx("100 q12 x 3")
ev_rx("100 over 2")
ev_rx("100 q 24 x 3 then 50 q12 x 2")
ev_rx("100 then 50 q 24 after 12")
ev_rx("100.2E-2 q4")
ev_rx("100 over 2.23")
ev_rx("100 q 12 x 3")
ev_rx("100 in 1 & 200 in 2")
parse_rx("100 mg q 24 then 200 mg q12")

```

ev_seq

Schedule a series of event objects

Description

Use this function when you want to schedule two or more event objects in time according the dosing interval (*ii*) and additional doses (*add1*).

Usage

```
ev_seq(..., ID = NULL, .dots = NULL, id = NULL)
```

```
## S3 method for class 'ev'
seq(...)
```

Arguments

<code>...</code>	event objects or numeric arguments named <code>wait</code> or <code>ii</code> to implement a period of no-dosing activity in the sequence (see Details).
<code>ID</code>	numeric vector of subject IDs.
<code>.dots</code>	a list of event objects that replaces <code>...</code>
<code>id</code>	deprecated; use <code>ID</code> .

Details

Use the generic [seq\(\)](#) when the first argument is an event object. If a waiting period (`wait` or `ii`) is the first event, you will need to use [ev_seq\(\)](#). When an event object has multiple rows, the end time for that sequence is taken to be one dosing interval after the event that takes place on the last row of the event object.

The doses for the next event line start after all of the doses from the previous event line plus one dosing interval from the previous event line (see **Examples**).

When numerics named `wait` or `ii` are mixed in with the event objects, a period with no dosing activity is incorporated into the sequence, between the adjacent dosing event objects. `wait` and `ii` accomplish a similar result, but differ by the starting point for the inactive period.

- Use `wait` to schedule the next dose relative to the end of the dosing interval for the previous dose.
- Use `ii` to schedule the next dose relative to the time of the the previous dose.

So `wait` acts like similar to an event object, by starting the waiting period from one dosing interval after the last dose while `ii` starts the waiting period from the time of the last dose itself. Both `wait` and `ii` can accomplish identical behavior depending on whether the last dosing interval is included (or not) in the value. Values for `wait` or `ii` can be negative.

NOTE: `.ii` had been available historically as an undocumented feature. Starting with `mrgsolve` version 0.11.3, the argument will be called `ii`. For now, both `ii` and `.ii` will be accepted but you will get a deprecation warning if you use `.ii`. Please use `ii` instead.

Values for `time` in any event object act like a prefix time spacer wherever that event occurs in the event sequence (see **Examples**).

Value

A single event object sorted by time.

Examples

```
e1 <- ev(amt = 100, ii = 12, addl = 1)

e2 <- ev(amt = 200)

seq(e1, e2)

seq(e1, ii = 8, e2)

seq(e1, wait = 8, e2)

seq(e1, ii = 8, e2, ID = seq(10))

ev_seq(ii = 12, e1, ii = 120, e2, ii = 120, e1)

seq(ev(amt = 100, ii = 12), ev(time = 8, amt = 200))
```

exdatasets	<i>Example input data sets</i>
------------	--------------------------------

Description

Example input data sets

Usage

```
data(exidata)
```

```
data(extran1)
```

```
data(extran2)
```

```
data(extran3)
```

```
data(exTheoph)
```

```
data(exBoot)
```

Details

- exidata holds individual-level parameters and other data items, one per row
- extran1 is a "condensed" data set
- extran2 is a full dataset
- extran3 is a full dataset with parameters
- exTheoph is the theophylline data set, ready for input into mrgsolve
- exBoot a set of bootstrap parameter estimates

Examples

```
mod <- mrgsolve::house() %>% update(end=240) %>% Req(CP)
```

```
## Full data set
```

```
data(exTheoph)
```

```
out <- mod %>% data_set(exTheoph) %>% mrgsim
```

```
out
```

```
plot(out)
```

```
## Condensed: mrgsolve fills in the observations
```

```
data(extran1)
```

```
out <- mod %>% data_set(extran1) %>% mrgsim
```

```
out
```

```
plot(out)
```

```
## Add a parameter to the data set
```

```

stopifnot(require(dplyr))
data <- extran1 %>% distinct(ID) %>% select(ID) %>%
  mutate(CL=exp(log(1.5) + rnorm(nrow(.), 0,sqrt(0.1)))) %>%
  left_join(extran1,.)

data

out <- mod %>% data_set(data) %>% carry_out(CL) %>% mrgsim
out
plot(out)

## idata
data(exidata)
out <- mod %>% idata_set(exidata) %>% ev(amt=100,ii=24,addl=10) %>% mrgsim
plot(out, CP~time|ID)

```

expand.idata

Create template data sets for simulation

Description

These functions expand all combinations of arguments using [expand.grid\(\)](#). `expand.idata()` generates an idata set; the others generate a full data set. The result always has only one row for one individual. Use `expand.evd()` or `evd_expand()` to render NMTRAN names (e.g. AMT or CMT) in upper case.

Usage

```
expand.idata(...)
```

```
expand.ev(...)
```

```
expand.evd(...)
```

```
ev_expand(...)
```

```
evd_expand(...)
```

Arguments

... passed to [expand.grid\(\)](#).

Details

An ID column is added as if not supplied by the user. In the output data frame, ID is always re-written as the row number.

For `expand.ev()`, defaults also added include `cmt = 1`, `time = 0`, `evid = 1`. If `total` is included, then `addl` is derived as `total-1`. If `tinf` is included, then an infusion rate is derived for row where `tinf` is greater than zero.

ev_expand() is a synonym for expand.ev() and evd_expand() is a synonym for expand.evd().

Value

A data frame containing one row for each combination of the items passed in . . . The result always has ID set to the row number.

Examples

```
idata <- expand.idata(CL = c(1,2,3), VC = c(10,20,30))

doses <- expand.ev(amt = c(300,100), ii = c(12,24), cmt = 1)

infusion <- expand.ev(amt = 100, tinf = 2)
```

expand_observations	<i>Insert observations into a data set</i>
---------------------	--

Description

Insert observations into a data set

Usage

```
expand_observations(data, times, unique = FALSE, obs_pos = -1L)
```

Arguments

data	a data set or event object.
times	a vector of observation times.
unique	logical; if TRUE then values for time are dropped if they are found anywhere in data.
obs_pos	determines sorting order for observations; use -1 (default) to put observations first; otherwise, use large integer to ensure observations are placed after doses.

Details

Non-numeric columns will be dropped with a warning.

Value

A data frame with additional rows for added observation records.

Examples

```
data <- expand.ev(amt = c(100, 200, 300))

expand_observations(data, times = seq(0, 48, 2))
```

get_tol	<i>Extract rtol and atol information from a model object</i>
---------	--

Description

Extract rtol and atol information from a model object

Usage

```
get_tol(x)
get_tol_list(x)
```

Arguments

`x` a model object.

Value

A data frame (`get_tol()`) or a named list (`get_tol_list()`).

See Also

[reset_tol\(\)](#), [custom_tol\(\)](#), [use_custom_tol\(\)](#), [use_scalar_tol\(\)](#)

Examples

```
mod <- house()
get_tol(mod)
get_tol_list(mod)
```

idata_set	<i>Select and modify a idata set for simulation</i>
-----------	---

Description

The individual data set (`idata_set`) is a data frame with one row for each individual in a population, specifying parameters and other individual-level data.

Usage

```

idata_set(x, data, ...)

## S4 method for signature 'mrgmod,data.frame'
idata_set(
  x,
  data,
  .subset = TRUE,
  .select = TRUE,
  object = NULL,
  need = NULL,
  ...
)

## S4 method for signature 'mrgmod,ANY'
idata_set(x, data, ...)

## S4 method for signature 'mrgmod,missing'
idata_set(x, object, ...)

```

Arguments

x	model object.
data	a data set that can be coerced to data.frame.
...	other arguments passed along when object is a function.
.subset	an unquoted expression passed to <code>dplyr::filter()</code> ; retain only certain rows in the data set.
.select	passed to <code>dplyr::select()</code> ; retain only certain columns in the data set; this should be the result of a call to <code>dplyr::vars()</code> .
object	character name of an object existing in \$ENV to use for the data set.
need	passed to <code>inventory()</code> .

Details

The `idata_set` is a data frame that specifies individual-level data for the problem. An ID column is required and there can be no more than one row in the data frame for each individual.

In most cases, the columns in the `idata_set` have the same names as parameters in the `param()` list. When this is the case, the parameter set is updated as the simulation proceeds once at the start of each individual. The `idata_set` can also be used to set initial conditions for each individual: for a compartment called CMT, make a column in `idata_set` called `CMT_0` and make the value the desired initial value for that compartment. Note that this initial condition will be overridden if you also set the `CMT_0` in `$MAIN ($PK)`.

The most common application of `idata_set` is to specify a population or batch of simulations to do. We commonly use `idata_set` with an event object (see `ev()`). In that case, the event gets applied to each individual in the `idata_set`.

It is also possible to provide both a `data_set` and a `idata_set`. In this case, the `idata_set` is used as a parameter lookup for IDs found in the `data_set`. Remember in this case, it is the `data_set` (not the `idata_set`) that determines the number of individuals in the simulation.

An error will be generated if any parameter columns in the input `idata` set contain NA.

See Also

[data_set\(\)](#), [ev\(\)](#)

Examples

```
mod <- mrgsolve::house()

data(exidata)

exidata

mod %>%
  idata_set(exidata, ID <= 2) %>%
  ev(amt = 100) %>%
  mrgsim() %>%
  plot()

mod %>%
  idata_set(exidata) %>%
  ev(amt = 100) %>%
  mrgsim()

mod %>% ev(amt = 100) %>% mrgsim(idata=exidata)
```

init

Methods for working with the model compartment list

Description

Calling `init()` with the model object as the first argument will return the model initial conditions as a [numericlist](#) object. See [numericlist](#) for methods to deal with `cmt_list` objects.

Usage

```
init(.x, ...)
```

S4 method for signature 'mrgmod'

```
init(.x, .y = list(), ..., .pat = "*")
```

S4 method for signature 'mrgsims'

```
init(.x, ...)
```



```
## S4 method for signature 'missing'
init(.x, ...)

## S4 method for signature 'list'
init(.x, ...)

## S4 method for signature 'ANY'
init(.x, ...)
```

Arguments

<code>.x</code>	the model object.
<code>...</code>	name = value assignments to update the initial conditions list.
<code>.y</code>	list to be merged into parameter list.
<code>.pat</code>	a regular expression (character) to be applied as a filter when printing compartments to the screen.

Details

Can be used to either get a compartment list object from a `mrghmod` model object or to update the compartment initial conditions in a model object. For both uses, the return value is a `cmt_list` object. For the former use, `init()` is usually called to print the compartment initial conditions to the screen, but the `cmt_list` object can also be coerced to a list or numeric R object.

Value

An object of class `cmt_list` (see [numericlist](#)).

Examples

```
## example("init")
mod <- mrgsolve::house()

init(mod)

init(mod, .pat="^C") ## may be useful for large models

class(init(mod))

init(mod)$CENT

as.list(init(mod))

as.data.frame(init(mod))
```

inventory	<i>Check whether all required parameters needed in a model are present in an object</i>
-----------	---

Description

This function has largely been superseded by [check_data_names\(\)](#).

Usage

```
inventory(x, obj, ..., .strict = FALSE)
```

Arguments

x	model object.
obj	data.frame to pass to idata_set() or data_set() .
...	capture dplyr-style parameter requirements.
.strict	whether to stop execution if all requirements are present (TRUE) or just warn (FALSE); see Details .

Details

If parameter requirements are not explicitly stated, the requirement defaults to all parameter names in x. Note that, by default, the inventory is not .strict unless the user explicitly states the parameter requirement. That is, if parameter requirements are explicitly stated, .strict will be set to TRUE if a value .strict was not passed in the call.

Value

x is returned invisibly.

See Also

[check_data_names\(\)](#)

Examples

```
## Not run:
  inventory(mod, idata, CL:V) # parameters defined, inclusively, CL through Volume
  inventory(mod, idata, everything()) # all parameters
  inventory(mod, idata, contains("OCC")) # all parameters containing OCC
  inventory(mod, idata, -F) # all parameters except F

## End(Not run)
```

`is.mrgmod`*Check if an object is a model object*

Description

The function checks to see if the object is either mrgmod or packmod.

Usage

```
is.mrgmod(x)
```

Arguments

`x` any object

Value

TRUE if the object inherits from either mrgmod or packmod class.

Examples

```
mod <- mrgsolve::house()
is.mrgmod(mod)
```

`is.mrgsims`*Check if an object is mrgsims output*

Description

Check if an object is mrgsims output

Usage

```
is.mrgsims(x)
```

Arguments

`x` any object.

Value

TRUE if x inherits mrgsims.

lctran

*Change the case of nmtran-like data items***Description**

Previous data set requirements included lower case names for data items like AMT and EVID. Lower case is no longer required. However, it is still a requirement that nmtran like data column names are either all lower case or all upper case.

Usage

```
lctran(data, ...)

## S3 method for class 'data.frame'
lctran(data, warn = TRUE, ...)

## S3 method for class 'ev'
lctran(data, ...)

uctran(data, ...)

## S3 method for class 'data.frame'
uctran(data, warn = TRUE, ...)

## S3 method for class 'ev'
uctran(data, ...)
```

Arguments

data	a data set with nmtran-like format or an event object.
...	for potential future use.
warn	if TRUE, a warning will be issued when there are both upper and lower case versions of any nmtran-like column in the data frame.

Details

Columns that will be renamed with lower or upper case versions:

- AMT / amt
- II / ii
- SS / ss
- CMT / cmt
- ADDL / addl
- RATE / rate
- EVID / evid

- TIME / time

If both lower and upper case versions of the name are present in the data frame, no changes will be made.

Value

A data frame or event object, with column names possibly converted to upper or lower case.

Examples

```
data <- data.frame(TIME = 0, AMT = 5, II = 24, addl = 2, WT = 80)
lctran(data)

data <- data.frame(TIME = 0, AMT = 5, II = 24, addl = 2, wt = 80)
uctran(data)

ev <- evd(amt = 100, evid = 3)
uctran(ev)

# warning
data <- data.frame(TIME = 1, time = 2, CMT = 5)
lctran(data)
```

loadso	<i>Load the model shared object</i>
--------	-------------------------------------

Description

Once the model is compiled, the model object can be used to re-load the model shared object (the compiled code underlying the mode) when the simulation is to be done in a different R process.

Usage

```
loadso(x, ...)
```

S3 method for class 'mrgmod'

```
loadso(x, ...)
```

Arguments

x	a model object.
...	not used.

Details

The loadso function most frequently needs to be used when parallelizing simulations across worker nodes. The model can be run after calling loadso, without requiring that it is re-compiled on worker nodes. It is likely required that the model is built (and the shared object stored) in a local directory off of the working R directory (see the second example).

Value

The model object (invisibly).

Examples

```
## Not run:
mod <- mread("pk1", modlib())
loadso(mod)

mod2 <- mread("pk2", modlib(), soloc = "build")
loadso(mod2)

## End(Not run)
```

matrix_helpers

Create matrices from vector input

Description

These functions are simple utilities for creating diagonal, block or correlation matrices.

Usage

```
bmat(..., correlation = FALSE, digits = -1)

cmat(..., digits = -1)

dmat(...)
```

Arguments

...	matrix data.
correlation	logical; if TRUE, off-diagonal elements are assumed to be correlations and converted to covariances.
digits	if greater than zero, matrix is passed to signif() (along with digits) prior to returning.

Details

`bmat()` makes a block matrix. `cmat()` makes a correlation matrix. `dmat()` makes a diagonal matrix.

Value

A matrix.

See Also

[as_bmat\(\)](#), [as_dmat\(\)](#)

Examples

```
dmat(1,2,3)/10

bmat(0.5,0.01,0.2)

cmat(0.5, 0.87,0.2)
```

mcode

Write, compile, and load model code

Description

This is a convenience function that ultimately calls [mread\(\)](#). Model code is written to a file and read back in using [mread\(\)](#).

Usage

```
mcode(model, code, project = getOption("mrgsolve.project", tempdir()), ...)

mcode_cache(
  model,
  code,
  project = getOption("mrgsolve.project", tempdir()),
  ...
)
```

Arguments

model	model name.
code	character string specifying a mrgsolve model.
project	project directory for the model.
...	passed to mread() ; see that help topic for other arguments that can be set.

Details

Note that the arguments are in slightly different order than [mread\(\)](#). The default project is [tempdir\(\)](#).

See the [mread\(\)](#) help topic for discussion about caching compilation results with [mcode_cache\(\)](#).

See Also

[mread\(\)](#), [mread_cache\(\)](#)

Examples

```
## Not run:
code <- '
$CMT DEPOT CENT
$PKMODEL ncmt=1, depot=TRUE
$MAIN
double CL = 1;
double V = 20;
double KA = 1;
'

mod <- mcode("example", code, compile = FALSE)

## End(Not run)
```

mcRNG	<i>Set RNG to use L'Ecuyer-CMRG</i>
-------	-------------------------------------

Description

Set RNG to use L'Ecuyer-CMRG

Usage

```
mcRNG()
```

modlib	<i>Internal model library</i>
--------	-------------------------------

Description

Pre-coded models are included in the mrgsolve installation; these can be compiled and loaded with `modlib()`. These models are usually most useful for exploratory simulation or learning mrgsolve. Production simulation work is typically accomplished by a custom-coded model.

Usage

```
modlib(model = NULL, ..., list = FALSE)
```

Arguments

- `model` character name of a model in the library.
- `...` passed to `mread_cache()`.
- `list` logical; if TRUE, a list of available models is returned.

Details

See [modlib_details](#), [modlib_pk](#), [modlib_pkpd](#), [modlib_tmdd](#), [modlib_viral](#) for details.

Call `modlib("<modelname>")` to compile and load a mode from the library.

Call `modlib(list=TRUE)` to list available models. Once the model is loaded (see examples below), call `as.list(mod)$code` to extract model code and equations.

Examples

```
## Not run:
mod <- mread("pk1cmt", modlib())
mod <- mread("pk2cmt", modlib())
mod <- mread("pk3cmt", modlib())
mod <- mread("pk1", modlib())
mod <- mread("pk2", modlib())
mod <- mread("popex", modlib())
mod <- mread("irm1", modlib())
mod <- mread("irm2", modlib())
mod <- mread("irm3", modlib())
mod <- mread("irm4", modlib())
mod <- mread("emax", modlib())
mod <- mread("effect", modlib())
mod <- mread("tmdd", modlib())
mod <- mread("viral1", modlib())
mod <- mread("viral2", modlib())
mod <- mread("pred1", modlib())
mod <- mread("pbpk", modlib())
mod <- mread("1005", modlib()) # embedded NONMEM result
mod <- mread("nm-like", modlib()) # model with nonmem-like syntax
mod <- mread("evtools", modlib())

as.list(mod)$code

## End(Not run)
```

modlib_details	<i>modlib: PK/PD Model parameters, compartments, and output variables</i>
----------------	---

Description

modlib: PK/PD Model parameters, compartments, and output variables

Compartments

- EV1, EV2: extravascular dosing compartments
- CENT: central PK compartment
- PERIPH: peripheral PK compartment
- PERIPH2: peripheral PK compartment 2
- RESP: response PD compartment (irm models)

Output variables

- CP: concentration in the central compartment (CENT/VC)
- RESP: response (emax model)

PK parameters

- KA1, KA2: first order absorption rate constants from first and second extravascular compartment (1/time)
- CL: clearance (volume/time)
- VC: volume of distribution, central compartment (volume)
- VP: volume of distribution, peripheral compartment (volume)
- VP2: volume of distribution, peripheral compartment 2 (volume)
- Q: intercompartmental clearance (volume/time)
- Q2: intercompartmental clearance 2 (volume/time)
- VMAX: maximum rate, nonlinear process (mass/time)
- KM: Michaelis constant (mass/volume)
- K10: elimination rate constant (1/time); CL/VC
- K12: rate constant for transfer to peripheral compartment from central (1/time); Q/VC
- K21: rate constant for transfer to central compartment from peripheral (1/time); Q/VP

PD parameters

- E0: baseline effect (emax model)
- EMAX, IMAX: maximum effect (response)
- EC50, IC50: concentration producing 50 percent of effect (mass/volume)
- KIN: zero-order response production rate (irm models) (response/time)
- KOUT: first-order response elimination rate (irm models) (1/time)
- n: sigmoidicity factor
- KE0: rate constant for transfer to effect compartment (1/time)

modlib_pk	<i>modlib: Pharmacokinetic models</i>
-----------	---------------------------------------

Description

modlib: Pharmacokinetic models

Arguments

... passed to update

Details

See [modlib_details](#) for more detailed descriptions of parameters and compartments.

The pk1cmt model is parameterized in terms of CL, VC, KA1 and KA2 and uses compartments EV1, EV2, and CENT. The pk2cmt model adds a PERIPH compartment and parameters Q and VP to that of the one-compartment model. Likewise, the three-compartment model (pk3cmt) adds PERIPH2 and parameters Q2 and VP2 to that of the two-compartment models. All pk models also have parameters VMAX (defaulting to zero, no non-linear clearance) and KM.

Value

an object of class packmod

Model description

All pk models have two extravascular dosing compartments and potential for linear and nonlinear clearance.

- pk1cmt: one compartment pk model using ODEs
- pk2cmt: two compartment pk model using ODEs
- pk3cmt: three compartment pk model using ODEs
- pk1: one compartment pk model in closed-form
- pk2: two compartment pk model in closed-form
- popex: a simple population pk model

modlib_pkpd

modlib: Pharmacokinetic / pharmacodynamic models

Description

modlib: Pharmacokinetic / pharmacodynamic models

Details

See [modlib_details](#) for more detailed descriptions of parameters and compartments.

All PK/PD models include 2-compartment PK model with absorption from 2 extravascular compartments and linear + nonlinear clearance. The PK models are parameterized with CL, VC, Q, VMAX, KM, KA1 and KA2 and implement compartments EV1, EV2, CENT, PERIPH. The indirect response models have compartment RESP and the emax model has output variable RESP. PD parameters include KIN, KOUT, IC50, EC50, IMAX, EMAX, E0, and n.

Also, once the model is loaded, use [see](#) method for mrgmod to view the model code.

Model description

- irm1 inhibition of response production
- irm2 inhibition of response loss
- irm3 stimulation of response production
- irm4 stimulation of response loss
- pd_effect effect compartment model
- emax sigmoid emax model

modlib_tmdd

*modlib: Target mediated disposition model***Description**

modlib: Target mediated disposition model

Arguments

... passed to update

Parameters

- KEL: elimination rate constant
- KTP: tissue to plasma rate constant
- KPT: plasma to tissue rate constant
- VC: volume of distribution
- KA1, KA2: absorption rate constants
- KINT: internalization rate constant
- KON: association rate constant
- KOFF: dissociation rate constant
- KSYN: target synthesis rate
- KDEG: target degradation rate constant

Compartments

- CENT: unbound drug in central compartment
- TISS: unbound drug in tissue compartment
- REC: concentration of target
- RC: concentration of drug-target complex
- EV1, EV2: extravascular dosing compartments

Output variables

- CP: unbound drug in the central compartment
- TOTAL: total concentration of target (complexed and uncomplexed)

`modlib_viral`*modlib: HCV viral dynamics models*

Description

modlib: HCV viral dynamics models

Models

- viral1: viral dynamics model with single HCV species
- viral2: viral dynamics model with wild-type and mutant HCV species

Parameters

- s: new hepatocyte synthesis rate (cells/ml/day)
- d: hepatocyte death rate constant (1/day)
- p: viral production rate constant (copies/cell/day)
- beta: new infection rate constant (ml/copy/day)
- delta: infected cell death rate constant (1/day)
- c: viral clearance rate constant (1/day)
- fit: mutant virus fitness
- N: non-target hepatocytes
- mu: forward mutation rate
- Tmax: maximum number of target hepatocytes (cells/ml)
- rho: maximum hepatocyte regeneration rate (1/day)

Compartments

- T: uninfected target hepatocytes (cells/ml)
- I: productively infected hepatocytes (cells/ml)
- V: hepatitis C virus (copies/ml)
- IM: mutant infected hepatocytes (cells/ml)
- VM: mutant hepatitis C virus (copies/ml)
- expos: exposure metric to drive pharmacodynamic model

mread*Read a model specification file*

Description

`mread()` reads and parses the `mrgsolve` model specification file, builds the model, and returns a model object for simulation. `mread_cache()` does the same, but caches the compilation result for later use. `mread_file()` can be used for convenience, taking the model file name as the first argument.

Usage

```
mread(  
  model,  
  project = getOption("mrgsolve.project", getwd()),  
  code = NULL,  
  file = NULL,  
  udll = TRUE,  
  ignore.stdout = TRUE,  
  raw = FALSE,  
  compile = TRUE,  
  audit = TRUE,  
  quiet = getOption("mrgsolve_mread_quiet", FALSE),  
  check.bounds = FALSE,  
  warn = TRUE,  
  soloc = getOption("mrgsolve.soloc", tempdir()),  
  capture = NULL,  
  preclean = FALSE,  
  recover = FALSE,  
  ...  
)  
  
mread_cache(  
  model = NULL,  
  project = getOption("mrgsolve.project", getwd()),  
  file = paste0(model, ".cpp"),  
  code = NULL,  
  soloc = getOption("mrgsolve.soloc", tempdir()),  
  quiet = FALSE,  
  preclean = FALSE,  
  capture = NULL,  
  ...  
)  
  
mread_file(file, ...)
```

Arguments

<code>model</code>	model name.
<code>project</code>	location of the model specification file and any headers to be included; see also the discussion about <code>model</code> ; this argument can be set via <code>options()</code> . library under details as well as the <code>modlib()</code> help topic.
<code>code</code>	a character string with model specification code to be used instead of a model file.
<code>file</code>	the full file name (with extension, but without path) where the model is specified.
<code>udll</code>	use unique name for shared object.
<code>ignore.stdout</code>	passed to system call when compiling the model; set this to <code>FALSE</code> to print output to the R console.
<code>raw</code>	if <code>TRUE</code> , return model content as a list, bypassing the compile step; this argument is typically used for debugging problems with the model build.
<code>compile</code>	logical; if <code>TRUE</code> , the model will be built.
<code>audit</code>	check the model specification file for errors.
<code>quiet</code>	don't print messages from mrgsolve when compiling.
<code>check.bounds</code>	check boundaries of parameter list.
<code>warn</code>	logical; if <code>TRUE</code> , print warning messages that may arise while building the model.
<code>soloc</code>	the directory location where the model shared object is built and stored; see details; this argument can be set via <code>options()</code> ; if the directory does not exist, <code>mread()</code> will attempt to create it.
<code>capture</code>	a character vector or comma-separated string of additional model variables to capture; these variables will be added to the capture list for the current call to <code>mread()</code> only.
<code>preclean</code>	logical; if <code>TRUE</code> , compilation artifacts are cleaned up first.
<code>recover</code>	if <code>TRUE</code> , a list of build will be returned in case the model shared object fails to compile; use this option to and the returned object to collect information assist in debugging.
<code>...</code>	passed to <code>update()</code> ; also arguments passed to <code>mread()</code> from <code>mread_cache()</code> .

Details

The `model` argument is required. For typical use, the `file` argument is omitted and the value for `file` is generated from the value for `model`. To determine the source file name, mrgsolve will look for a file extension in `model`. A file extension is assumed when it finds a period followed by one to three alpha-numeric characters at the end of the string (e.g. `mymodel.txt` but not `my.model`). If no file extension is found, the extension `.cpp` is assumed (e.g. file is `<model-name>.cpp`). If a file extension is found, file is `<model-name>`.

Best practice is to avoid using `.` in `model` unless you are using `model` to point to the model specification file name. Otherwise, use `mread_file()`.

Use the `soloc` argument to specify a directory location for building the model. This is the location where the model shared object will be stored on disk. The default is a temporary directory, so compilation artifacts are lost when R restarts when the default is used. Changing `soloc` to a persistent

directory location will preserve those artifacts across R restarts. Also, if simulation from a single model is being done in separate processes on separate compute nodes, it might be necessary to store these compilation artifacts in a local directory to make them accessible to the different nodes. If the `soloc` directory does not exist, `mread()` will attempt to create it.

Similarly, using `mread_cache()` will cache results in the temporary directory and the cache cannot be accessed after the R process is restarted.

Model Library

`mrsgsolve` comes bundled with several pre-coded PK, PK/PD, and other systems models that are accessible via the `mread()` interface.

Models available in the library include:

- PK models: `pk1cmt`, `pk2cmt`, `pk3cmt`, `pk1`, `pk2`, `popex`, `tmdd`
- PKPD models: `irm1`, `irm2`, `irm3`, `irm4`, `emax`, `effect`
- Other models: `viral1`, `viral2`

When the library model is accessed, `mrsgsolve` will compile and load the model as you would for any other model. It is only necessary to reference the correct model name and point the project argument to the `mrsgsolve` model library location via `modlib()`.

For more details, see [modlib_pk](#), [modlib_pkpd](#), [modlib_tmdd](#), [modlib_viral](#), and [modlib_details](#) for more information about the state variables and parameters in each model.

See Also

[mcode\(\)](#), [mcode_cache\(\)](#)

Examples

```
## Not run:
code <- '
$PARAM CL = 1, VC = 5
$CMT CENT
$ODE dxdt_CENT = -(CL/VC)*CENT;
'

mod <- mcode("ex_mread", code)
mod

mod %>% init(CENT=1000) %>% mrsgsim() %>% plot()

mod <- mread("irm3", modlib())

# if the model is in the file mymodel.cpp
mod <- mread("mymodel")

# if the model is in the file mymodel.txt
mod <- mread(file = "mymodel.txt")

or
```



```
mod <- mread_file("mymodel.txt")

## End(Not run)
```

mread_yaml*Read a model from yaml format*

Description

Read back models written to file using [mwrite_yaml\(\)](#). Function [yaml_to_cpp\(\)](#) is also provided to convert the yaml file to mrgsolve cpp file format.

Usage

```
mread_yaml(
  file,
  model = basename(file),
  project = tempdir(),
  update = FALSE,
  ...
)

yaml_to_cpp(file, model = basename(file), project = getwd(), update = TRUE)
```

Arguments

file	the yaml file name.
model	a new model name to use when calling mread_yaml() .
project	the directory where the model should be built.
update	TRUE if model settings should be written into the cpp file in a <code>\$SET</code> block.
...	passed to mread() .

Details

Note that [yaml_to_cpp\(\)](#) by default writes model settings into the cpp file. [mread_yaml\(\)](#) does not write model settings into the file but rather update the model object directly with data read back from the yaml file.

Value

A model object.

See Also

[mwrite_yaml\(\)](#)

Examples

```
mod <- house()

temp <- tempfile(fileext = ".yaml")

mwrite_yaml(mod, file = temp)

# Note: this model is not compiled
mod <- mread_yaml(temp, model = "new-house", compile = FALSE)
mod

cppfile <- yaml_to_cpp(temp, project = tempdir())

readLines(cppfile)
```

mrgsim

Simulate from a model object

Description

This function sets up the simulation run from data stored in the model object as well as arguments passed in. Use [mrgsim_q\(\)](#) instead to benchmark mrgsolve or to do repeated quick simulation for tasks like parameter optimization, sensitivity analyses, or optimal design. See [mrgsim_variants](#) for other mrgsim-like functions that have more focused inputs. `mrgsim_df` coerces output to `data.frame` prior to returning.

Usage

```
mrgsim(x, data = NULL, idata = NULL, events = NULL, nid = NULL, ...)

mrgsim_df(..., output = "df")

do_mrgsim(
  x,
  data,
  idata = no_idata_set(),
  carry_out = carry.out,
  carry.out = character(0),
  recover = character(0),
  seed = as.integer(NA),
  Request = character(0),
  output = NULL,
  capture = NULL,
  obsonly = FALSE,
  obsaug = FALSE,
  tgrid = NULL,
  etasrc = "omega",
```

```

    recsort = 1,
    deslist = list(),
    descol = character(0),
    filbak = TRUE,
    tad = FALSE,
    nocb = TRUE,
    skip_init_calc = FALSE,
    ss_n = 500,
    ss_fixed = FALSE,
    interrupt = 256,
    ...
)

```

Arguments

<code>x</code>	the model object.
<code>data</code>	NMTRAN-like data set (see data_set()).
<code>idata</code>	a matrix or data frame of model parameters, one parameter per row (see idata_set()).
<code>events</code>	an event object.
<code>nid</code>	integer number of individuals to simulate; only used if <code>idata</code> and <code>data</code> are missing.
<code>...</code>	passed to update() and do_mrgsim() .
<code>output</code>	if NULL (the default) a <code>mrgsim</code> s object is returned; otherwise, pass <code>df</code> to return a data.frame or <code>matrix</code> to return a matrix.
<code>carry_out</code>	numeric data items to copy into the output.
<code>carry.out</code>	soon to be deprecated; use <code>carry_out</code> instead.
<code>recover</code>	character column names in either <code>data</code> or <code>idata</code> to join back (recover) to simulated data; may be any class (e.g. numeric, character, factor, etc).
<code>seed</code>	deprecated.
<code>Request</code>	compartments or captured variables to retain in the simulated output; this is different than the <code>request</code> slot in the model object, which refers only to model compartments.
<code>capture</code>	character file name used for debugging (not related to <code>\$CAPTURE</code>).
<code>obsonly</code>	if TRUE, dosing records are not included in the output.
<code>obsaug</code>	augment the data set with time grid observations; when TRUE and a full data set is used, the simulated output is augmented with an observation at each time in stime() . When using <code>obsaug</code> , a flag indicating augmented observations can be requested by including <code>a.u.g</code> in <code>carry_out</code> .
<code>tgrid</code>	a <code>tgrid</code> object; or a numeric vector of simulation times or another object with an <code>stime</code> method.
<code>etasrc</code>	source for <code>ETA()</code> values in the model; values can include: <code>"omega"</code> , <code>"data"</code> , <code>"data.all"</code> , <code>"idata"</code> , or <code>"idata.all"</code> ; see 'Details'.

recsort	record sorting flag. Default value is 1. Possible values are 1,2,3,4: 1 and 2 put doses in a data set after padded observations at the same time; 3 and 4 put those doses before padded observations at the same time. 2 and 4 will put doses scheduled through addl after observations at the same time; 1 and 3 put doses scheduled through addl before observations at the same time. recsort will not change the order of your input data set if both doses and observations are given.
deslist	a list of tgrid objects.
descol	the name of a column for assigning designs.
filbak	carry data items backward when the first data set row has time greater than zero.
tad	when TRUE a column is added to simulated output is added showing the time since the last dose. Only data records with evid == 1 will be considered doses for the purposes of tad calculation. The tad can be properly calculated with a dosing lag time in the model as long as the dosing lag time (specified in \$MAIN) is always appropriate for any subsequent doses scheduled through addl. This will always be true if the lag time doesn't change over time. But it might (possibly) not hold if the lag time changes prior to the last dose in the addl sequence. This known limitation shouldn't affect tad calculation in most common dosing lag time implementations.
nocb	if TRUE, use next observation carry backward method; otherwise, use locf.
skip_init_calc	don't use \$MAIN to calculate initial conditions.
ss_n	maximum number of iterations for determining steady state for the PK system; a warning will be issued if steady state is not achieved within ss_n iterations when ss_fixed is TRUE.
ss_fixed	if FALSE (the default), then a warning will be issued if the system does not reach steady state within ss_n iterations given the model tolerances rtol and atol; if TRUE, the number of iterations for determining steady state are capped at ss_n and no warning will be issued if steady state has not been reached within ss_n dosing iterations. To silence warnings related to steady state, set ss_fixed to TRUE and set ss_n as the maximum number of iterations to try when advancing the system for steady state determination.
interrupt	integer check user interrupt interval; when interrupt is a positive integer, the simulation will check for the user interrupt signal every interrupt simulation records; pass a negative number to never check for the user interrupt interval.

Details

- Use `mrgsim_df()` to return a data frame rather than `mrgsim` object.
- Both `data` and `idata` will be coerced to numeric matrix
- `carry_out` can be used to insert data columns into the output data set. This is partially dependent on the nature of the data brought into the problem.
- When using `data` and `idata` together, an error is generated if an ID occurs in `data` but not `idata`. Also, when looking up data in `idata`, ID in `idata` is assumed to be uniquely keyed to ID in `data`. No error is generated if ID is duplicated in `data`; parameters will be used from the first occurrence found in `idata`.

- `carry_out`: `idata` is assumed to be individual-level and variables that are carried from `idata` are repeated throughout the individual's simulated data. Variables carried from `data` are carried via last-observation carry forward. NA is returned from observations that are inserted into simulated output that occur prior to the first record in `data`.
- `recover`: this is similar to `carry_out` with respect to end result, but it uses a different process. Columns to be recovered are cached prior to running the simulation, and then joined back on to the simulated data. So, whereas `carry_out` will only accept numeric data items, `recover` can handle data frame columns of any type. There is a small decrease in performance with `recover` compared to `carry_out`, but it is likely that the performance difference is difficult to perceive (when the simulation runs very fast) or only a small fractional increase in run time when the simulation is very large. And any performance hit is likely to be well worth it in light of the convenience gain. Just think carefully about using this feature when every millisecond counts.
- `etasrc`: this argument lets you control where $ETA(n)$ come from in the model. When `etasrc` is set to "omega" (the default), ETAs will be simulated from a multivariate normal distribution defined by the `$OMEGA` blocks in the model. Alternatively, input data or `idata` sets can be used to pass in fixed $ETA(n)$ by setting `etasrc` to "data", "idata", "data.all" or "idata.all". When `etasrc` is set to "data" or "data.all", the input data set will be scanned for columns called `ETA1`, `ETA2`, ..., `ETAn` and those values will be copied into the appropriate slot in the `ETA()` vector. Only the first record for each individual will be copied into `ETA()`; all records after the first will be ignored. When there are more than 9 ETAs in a model, `NONMEM` will start naming the outputs `ET10`, `ET11` etc rather than `ETA10` and `ETA11`. When `mrgsolve` is looking for these columns, it will first search, for example, `ET10` and use that value if it is found. If `ET10` isn't found and there are more than 9 ETAs, then it will *also* search for `ETA10`. An error will be generated in case `mrgsolve` finds both the `ETA` and `ET` name variant for the tenth and higher `ETA` (e.g. it is an error to have both `ETA10` and `ET10` in the data set). When `mrgsolve` is searching for `ETA` columns in the data set, it will *only* look for `ETAn` up to the number of rows (or columns) in all the model `$OMEGA` blocks. For example, if `$OMEGA` is `5x5`, only `ETA1` through `ETA5` will be searched. An error will be generated in case `mrgsolve` finds *no* columns with `ETAn` names and something other than `etasrc = "omega"` was passed. When `etasrc = "data"` and an `ETAn` column is missing from the data set, the missing `ETA()` will be set to 0. Alternatively, the user can pass `etasrc = "data.all"` which causes an error to be generated if any `ETAn` is missing from the data set. Use this option when you intend to have *all* ETAs attached to the data set and want an error generated if `mrgsolve` finds one or more of them is missing. Using `etasrc = "idata"` or "idata.all", the behavior is identical to "data" (or "data.all"), except `mrgsolve` will look at the `idata` set rather than `data` set.

Value

An object of class `mrgsims`.

See Also

`mrgsim_variants`, `mrgsim_q()`

Examples

```
## example("mrgsim")
```

```

e <- ev(amt = 1000)

mod <- mrgsolve::house()

out <- mod %>% ev(e) %>% mrgsim()

plot(out)

out <- mod %>% ev(e) %>% mrgsim(end=22)

out

data(exTheoph)

out <- mod %>% data_set(exTheoph) %>% mrgsim()

out

out <- mod %>% mrgsim(data=exTheoph)

out <- mrgsim(mod, data=exTheoph, obsonly=TRUE)

out

out <- mod %>% mrgsim(data=exTheoph, obsaug=TRUE, carry_out="a.u.g")

out

out <- mod %>% ev(e) %>% mrgsim(outvars="CP,RESP")

out

a <- ev(amt = 1000, group = 'a')
b <- ev(amt = 750, group = 'b')
data <- as_data_set(a,b)

out <- mrgsim_d(mod, data, recover="group")

out

```

Description

These methods modify the data in a mrgsim object and return a data frame. Contrast with the functions in [mrgsim_modify](#).

Usage

```
## S3 method for class 'mrgsim'
pull(.data, ...)

## S3 method for class 'mrgsim'
filter(.data, ...)

## S3 method for class 'mrgsim'
group_by(.data, ..., add = FALSE, .add = FALSE)

## S3 method for class 'mrgsim'
distinct(.data, ..., .keep_all = FALSE)

## S3 method for class 'mrgsim'
mutate(.data, ...)

## S3 method for class 'each'
summarise(.data, fns, ...)

## S3 method for class 'mrgsim'
summarise(.data, ...)

## S3 method for class 'mrgsim'
do(.data, ..., .dots)

## S3 method for class 'mrgsim'
select(.data, ...)

## S3 method for class 'mrgsim'
slice(.data, ...)

as_data_frame.mrgsim(x, ...)

## S3 method for class 'mrgsim'
as_tibble(x, ...)

as.tbl.mrgsim(x, ...)
```

Arguments

<code>.data</code>	an mrgsim object; passed to various dplyr functions
<code>...</code>	passed to other methods
<code>add</code>	passed to <code>dplyr::group_by</code> (for dplyr < 1.0.0)
<code>.add</code>	passed to <code>dplyr::group_by</code> (for dplyr >= 1.0.0)
<code>.keep_all</code>	passed to <code>dplyr::distinct</code>
<code>fns</code>	passed to <code>dplyr::summarise_each</code>
<code>.dots</code>	passed to various dplyr functions

x mrgsims object.

Details

For the `select_sims` function, the dots ... must be either compartment names or variables in `$CAPTURE`. An error will be generated if no valid names are selected or the names for selection are not found in the simulated output.

See Also

[mrgsims_modify](#)

Examples

```
out <- mrgsim(house(), events = ev(amt = 100), end = 5, delta=1)

dplyr::filter(out, time==2)

dplyr::mutate(out, label = "abc")

dplyr::select(out, time, RESP, CP)
```

mrgsims_modify

Methods for modifying mrgsims objects

Description

These functions modify the simulated data in an mrgsims object and return the modified object. Contrast with the functions in [mrgsims_dplyr](#).

Usage

```
mutate_sims(.data, ...)

select_sims(.data, ...)

filter_sims(.data, ...)
```

Arguments

.data a mrgsims object.
 ... other arguments passed to the dplyr functions.

See Also

[mrgsims_dplyr](#)

Examples

```

out <- mrgsim(house(), events = ev(amt = 100))

filter_sims(out, time > 2)

mutate_sims(out, label = "abc")

select_sims(out, RESP, CP)

```

mrgsim_q	<i>Simulate from a model object with quicker turnaround</i>
----------	---

Description

Use the function when you would usually use `mrgsim_d()`, but you need a quicker turnaround time. The timing differences might be difficult to detect for a single simulation run but could become appreciable with repeated simulation. See **Details** for important differences in how `mrgsim_q()` is invoked compared to `mrgsim()` and `mrgsim_d()`. This function should always be used for benchmarking simulation time with `mrgsolve`.

Usage

```

mrgsim_q(
  x,
  data,
  recsort = 1,
  stime = numeric(0),
  output = "mrgsims",
  skip_init_calc = FALSE,
  simcall = 0,
  etasrc = "omega"
)

```

Arguments

<code>x</code>	a model object.
<code>data</code>	a simulation data set.
<code>recsort</code>	record sorting flag.
<code>stime</code>	a numeric vector of observation times; these observation times will only be added to the output if there are no observation records in data.
<code>output</code>	output data type; if "mrgsims", then the default output object is returned; if "df" then a data frame is returned.
<code>skip_init_calc</code>	don't use \$MAIN to calculate initial conditions.
<code>simcall</code>	not used; only the default value of 0 is allowed.
<code>etasrc</code>	source for ETA() values in the model; values can include: "omega", "data", "data.all", "idata", or "idata.all"; see 'Details' in <code>mrgsim()</code> .

Details

`mrgsim_q()` mainly cuts some of the overhead from the simulation. So, the primary efficiency gain from using `mrgsim_q()` comes when the simulation executes very quickly. It is unlikely you will see a big performance difference between `mrgsim_q()` and `mrgsim()` when the model is difficult to solve or if there is a large input data set.

This function does not support the piped simulation workflow. All arguments must be passed into the function except for `x`.

A data set is required for this simulation workflow. The data set can have only dosing records or doses with observations. When the data set only includes doses, a single numeric vector of observation times should be passed in.

This simulation workflow does not support Req (request) functionality. All compartments and captured variables will always be returned in the simulation output.

This simulation workflow does not support carry-out functionality.

This simulation workflow does not accept arguments to be passed to `update()`. This must be done by a separate call to `update()`.

This simulation workflow does not support use of event objects. If an event object is needed, it should be converted to a data set prior to the simulation run (see `as_data_set()` or `as.data.frame()`).

This simulation workflow does not support idata sets or any feature enabled by idata set use. Individual level parameters should be joined onto the data set prior to simulation. Otherwise `mrgsim_i()` or `mrgsim_ei()` should be used.

By default, a `mrgsims` object is returned (as with `mrgsim()`). Use the `output = "df"` argument to request a plain `data.frame` of simulated data on return.

Value

By default, an object of class `mrgsims`. Use `output = "df"` to return a data frame.

See Also

`mrgsim()`, `mrgsim_variants`, `qsim()`

Examples

```
mod <- mrgsolve::house()

data <- expand.ev(amt = c(100, 300, 1000))

out <- mrgsim_q(mod, data)

out
```

mrgsim_variants	<i>mrgsim variant functions</i>
-----------------	---------------------------------

Description

These functions are called by `mrgsim()` and have explicit input requirements written into the function name. The motivation behind these variants is to give the user a clear workflow with specific, required inputs as indicated by the function name. Use `mrgsim_q()` instead to benchmark mrgsolve or to do repeated quick simulation for tasks like parameter optimization, sensitivity analyses, or optimal design.

Usage

```
mrgsim_e(x, events, idata = NULL, data = NULL, ...)
mrgsim_d(x, data, idata = NULL, events = NULL, ...)
mrgsim_ei(x, events, idata, data = NULL, ...)
mrgsim_di(x, data, idata, events = NULL, ...)
mrgsim_i(x, idata, data = NULL, events = NULL, ...)
mrgsim_0(x, idata = NULL, data = NULL, events = NULL, ...)
```

Arguments

<code>x</code>	the model object.
<code>events</code>	an event object.
<code>idata</code>	a matrix or data frame of model parameters, one parameter per row (see <code>idata_set()</code>).
<code>data</code>	NMTRAN-like data set (see <code>data_set()</code>).
<code>...</code>	passed to <code>update()</code> and <code>do_mrgsim()</code> .

Details

Important: all of these functions require that data, idata, and/or events be pass directly to the functions. They will not recognize these inputs from a pipeline.

- `mrgsim_e` simulate using an event object
- `mrgsim_ei` simulate using an event object and `idata_set`
- `mrgsim_d` simulate using a `data_set`
- `mrgsim_di` simulate using a `data_set` and `idata_set`
- `mrgsim_i` simulate using a `idata_set`
- `mrgsim_0` simulate using just the model
- `mrgsim_q` simulate from a data set with quicker turnaround (see `mrgsim_q()`)

See Also

```
mrgsim(), mrgsim_q(), qsim()
```

mutate.ev	<i>dplyr verbs for event objects</i>
-----------	--------------------------------------

Description

dplyr verbs for event objects

Usage

```
## S3 method for class 'ev'  
mutate(.data, ...)  
  
## S3 method for class 'ev'  
select(.data, ...)  
  
## S3 method for class 'ev'  
filter(.data, ...)
```

Arguments

- .data the event object.
- ... passed to the dplyr function.

mwrite_cpp	<i>Write a model to native mrgsolve format</i>
------------	--

Description

Model code is written to a file in native mrgsolve format. This can be useful for (1) breaking connection to NONMEM modeling outputs that are imported by \$NMXML or \$NMEXT and (2) saving model updates (e.g., an updated parameter list). Models can be read back using [mread\(\)](#).

Usage

```
mwrite_cpp(x, file, update = TRUE)
```

Arguments

- x a model object.
- file output file name; if non-character (e.g., NULL), no output will be written to file.
- update TRUE if model settings should be written into the cpp file in a \$SET block.

Details

See important details in [mwrite_yaml\(\)](#).

Value

A list containing data that was written out to the cpp file, with added item file, is returned invisibly.

See Also

[mwrite_yaml\(\)](#), [yaml_to_cpp\(\)](#)

Examples

```
temp <- tempfile(fileext = ".mod")

mod <- modlib("pk1", compile = FALSE)

x <- mwrite_cpp(mod, file = temp)

mod <- mread(x$file, compile = FALSE)

mod
```

mwrite_yaml

Write model code to yaml format

Description

Model code is written to a readable, transport format. This transport format can be useful for (1) breaking connection to NONMEM modeling outputs that are imported by \$NMXML or \$NMEXT and (2) saving model updates (e.g., an updated parameter list). Models can be read back using [mread_yaml\(\)](#) or converted to mrgsolve cpp format with [yaml_to_cpp\(\)](#).

Usage

```
mwrite_yaml(x, file, digits = 8)
```

Arguments

x	a model object.
file	output file name; if non-character (e.g., NULL), no output will be written to file.
digits	precision to use when writing outputs.

Details

Parameters and omega and sigma matrices that were imported via \$NMXML or \$NMEXT will be written into the yaml file and the NONMEM import blocks will be dropped. This allows the user to load a model based on a NONMEM run without having a connection to that output (e.g., root.xml or root.ext). Given that the connection to the NONMEM modeling outputs is broken when writing to yaml, any update to the NONMEM run will only be propagated to the yaml file when `mwrite_yaml()` is run again.

The yaml file does not currently have the ability to track other external dependencies, such as user-defined header files or other code that might be sourced in by the user when the model is loaded via `mread()`. NONMEM xml and ext files imported by \$NMXML or \$NMEXT are the *only* external dependencies that are accounted for in the yaml transport file.

Value

A list containing data that was written out to the yaml file, with added item file, is returned invisibly.

See Also

`mread_yaml()`, `yaml_to_cpp()`

Examples

```
mod <- house()

temp1 <- tempfile(fileext = ".yaml")

x <- mwrite_yaml(mod, temp1)

readLines(temp1)
```

names,mrgmod-method	<i>Get all names from a model object</i>
---------------------	--

Description

Get all names from a model object

Usage

```
## S4 method for signature 'mrgmod'
names(x)
```

Arguments

x the model object

Examples

```
mod <- mrgsolve::house()
names(mod)
```

nmext

Import model estimates from a NONMEM ext file

Description

Import model estimates from a NONMEM ext file

Usage

```
nmext(
  run = NA_real_,
  project = getwd(),
  file = paste0(run, ".ext"),
  path = NULL,
  root = c("working", "cppfile"),
  index = "last",
  theta = TRUE,
  omega = TRUE,
  sigma = TRUE,
  olabels = NULL,
  slabels = NULL,
  oprefix = "",
  sprefix = "",
  tname = "THETA",
  oname = "...",
  sname = "...",
  read_fun = "data.table",
  env = NULL
)
```

Arguments

run	run number.
project	project directory.
file	deprecated; use path instead.
path	full path to NONMEM ext file.
root	the directory that path and project are relative to; this is currently limited to the working directory or cppdir, the directory where the model file is located.
index	the estimation number to return; "last" will return the last estimation results; otherwise, pass an integer indicating which estimation results to return.

theta	logical; if TRUE, the \$THETA vector is returned.
omega	logical; if TRUE, the \$OMEGA matrix is returned.
sigma	logical; if TRUE, the \$SIGMA matrix is returned.
olabels	labels for \$OMEGA.
slabels	labels for \$SIGMA.
oprefix	prefix for \$OMEGA labels.
sprefix	prefix for \$SIGMA labels.
tname	name for \$THETA.
oname	name for \$OMEGA.
sname	name for \$SIGMA.
read_fun	function to use when reading the ext file.
env	internal use only.

See Also

[nmxml\(\)](#), [read_nmext\(\)](#)

nmxml

Import model estimates from a NONMEM xml file

Description

Import model estimates from a NONMEM xml file

Usage

```
nmxml(
  run = numeric(0),
  project = character(0),
  file = character(0),
  path = character(0),
  root = c("working", "cppfile"),
  theta = TRUE,
  omega = TRUE,
  sigma = TRUE,
  olabels = NULL,
  slabels = NULL,
  oprefix = "",
  sprefix = "",
  tname = "THETA",
  oname = "...",
  sname = "...",
  index = "last",
  xpath = "../nm:estimation",
  env = NULL
)
```


Arguments

run	run number.
project	project directory.
file	deprecated; use path instead.
path	the complete path to the run.xml file.
root	the directory that path and project are relative to; this is currently limited to the working directory or cppdir, the directory where the model file is located.
theta	logical; if TRUE, the \$THETA vector is returned.
omega	logical; if TRUE, the \$OMEGA matrix is returned.
sigma	logical; if TRUE, the \$SIGMA matrix is returned.
olabels	labels for \$OMEGA.
slabels	labels for \$SIGMA.
oprefix	prefix for \$OMEGA labels.
sprefix	prefix for \$SIGMA labels.
tname	name for \$THETA.
oname	name for \$OMEGA.
sname	name for \$SIGMA.
index	the estimation number to return; "last" will return the last estimation results; otherwise, pass an integer indicating which estimation results to return.
xpath	xml path containing run results; if the default doesn't work, consider using <code>./estimation</code> as an alternative; see details.
env	internal use only.

Details

If run and project are supplied, the .xml file is assumed to be located in run.xml, in directory run off the project directory. If file is supplied, run and project arguments are ignored.

This function requires that the xml2 package be installed and loadable. If `requireNamespace("xml2")` fails, an error will be generated.

nmxml usually expects to find run results in the xpath called `./nm:estimation`. Occasionally, the run results are not stored in this namespace but no namespaces are found in the xml file. In this case, the user can specify the xpath containing run results. Consider trying `./estimation` as an alternative if the default fails.

Value

A list with theta, omega and sigma elements, depending on what was requested.

See Also

nmext

Examples

```
if(requireNamespace("xml2")) {
  proj <- system.file("nonmem", package = "mrgsolve")
  mrgsolve::nmxml(run = 1005, project = proj)
}
```

numerics_only	<i>Prepare data.frame for input to mrgsim()</i>
---------------	---

Description

Prepare data.frame for input to mrgsim()

Usage

```
numerics_only(x, quiet = FALSE, convert_lgl = FALSE)
```

Arguments

x	a input data set.
quiet	logical indicating whether or not warnings should be printed.
convert_lgl	if TRUE, convert logical columns with as.integer() .

obsaug	<i>Augment observations in the simulated output</i>
--------	---

Description

Augment observations in the simulated output

Usage

```
obsaug(x, value = TRUE, ...)
```

Arguments

x	model object
value	the value for obsaug
...	passed along There is also a obsaug argument to mrgsim that can be set to accomplish the same thing as a call to obsaug in the pipeline.

obsonly	<i>Collect only observation records in the simulated output</i>
---------	---

Description

Collect only observation records in the simulated output

Usage

```
obsonly(x, value = TRUE, ...)
```

Arguments

x	model object.
value	use 'TRUE' to collect and return observation records only.
...	not used.

Details

There is also an 'obsonly' argument to [mrgsim()] that can be set to accomplish the same thing as a call to 'obsonly()' in the pipeline.

omega	<i>Manipulate OMEGA matrices</i>
-------	----------------------------------

Description

The primary function is `omat()` that can be used to both get the \$OMEGA matrices out of a model object and to update \$OMEGA matrices in a model object.

Usage

```
omat(.x, ...)
```

```
## S4 method for signature 'missing'
```

```
omat(.x, ...)
```

```
## S4 method for signature 'matrix'
```

```
omat(.x, ..., labels = list())
```

```
## S4 method for signature 'NULL'
```

```
omat(.x, ...)
```

```
## S4 method for signature 'list'
```

```
omat(.x, ...)
```

```
## S4 method for signature 'omegalist'
omat(.x, ...)

## S4 method for signature 'mrgmod'
omat(.x, ..., make = FALSE, open = FALSE)

## S4 method for signature 'mrgsims'
omat(.x, make = FALSE, ...)
```

Arguments

<code>.x</code>	a matrix, list of matrices or <code>matlist</code> object.
<code>...</code>	passed to other functions, including <code>modMATRIX()</code> .
<code>labels</code>	character vector of names for Ω elements; must be equal to number of rows/columns in the matrix.
<code>make</code>	logical; if TRUE, matrix list is rendered into a single matrix.
<code>open</code>	passed to <code>merge.list()</code> .
<code>x</code>	<code>matlist</code> object.

See Also

`smat()`, `dmat()`, `bmat()`, `cmat()`

Examples

```
# example("omega")
mat1 <- matrix(1)
mat2 <- diag(c(1,2,3))
mat3 <- matrix(c(0.1, 0.002, 0.002, 0.5), 2,2)
mat4 <- dmat(0.1, 0.2, 0.3, 0.4)

omat(mat1)
omat(mat1, mat2, mat3)
omat(A = mat1, B = mat2, C = mat3)

mod <- mrgsolve::house() %>% omat(mat4)

omat(mod)
omat(mod, make = TRUE)
as.matrix(omat(mod))
```

outvars	<i>Show names of current output variables</i>
---------	---

Description

Outputs can include model compartments or variables defined in the model that have been marked to capture in simulated output.

Usage

```
outvars(x, unlist = FALSE)
```

Arguments

x	model object.
unlist	if TRUE then a character vector (rather than list) is returned.

Value

When unlist is FALSE (default) : a named list, with cmt showing names of output compartments and capture giving names of output variables in capture. When unlist is TRUE, then a single, unnamed character vector of outvar names is returned.

Examples

```
mod <- mrgsolve::house()
outvars(mod)
```

param	<i>Create and work with parameter objects</i>
-------	---

Description

See [numericlist](#) for methods to deal with parameter_list objects.

Usage

```
param(.x, ...)

## S4 method for signature 'mrgmod'
param(.x, .y = NULL, ..., .pat = "*", .strict = FALSE)

## S4 method for signature 'mrgsims'
param(.x, ...)
```

```
## S4 method for signature 'missing'
param(..., .strict = TRUE)

## S4 method for signature 'list'
param(.x, ...)

## S4 method for signature 'ANY'
param(.x, ...)

allparam(.x)
```

Arguments

<code>.x</code>	the model object.
<code>...</code>	passed along or name/value pairs to update the parameters in a model object; when passing new values this way, all values must be numeric and all names must exist in the parameter list for <code>.x</code> .
<code>.y</code>	an object to be merged into parameter list; non-NULL values must be named list, data.frame, numeric vector, or <code>parameter_list</code> object; named items that do not exist in the parameter list are allowed and will be silently ignored; use the <code>.strict</code> argument to require that all names in <code>.y</code> exist already in the parameter list.
<code>.pat</code>	a regular expression (character) to be applied as a filter for which parameters to show when printing.
<code>.strict</code>	if TRUE, all names to be updated must be found in the parameter list.

Details

Can be used to either get a parameter list object from a `mrgmod` model object or to update the parameters in a model object. For both uses, the return value is a `parameter_list` object. For the former use, `param()` is usually called to print the parameters to the screen, but the `parameter_list` object can also be coerced to a list or numeric R object.

Use `allparam()` to get a `parameter_list` object including both model parameters and data items listed in `$FIXED`.

The update to parameters can be permissive (candidates with names that don't exist in the parameter list are silently ignored) or strict (all candidates must already exist in the parameter list). When passing candidate values via `...`, the update is strict and an error is generated if you pass a name that isn't found in the parameter list. When candidate values are passed as a named object via `.y`, then the update is permissive. Any permissive update can be made strict (error if foreign names are found in the candidates) by passing `.strict = TRUE`.

An alternative is to assess the incoming names using `inventory()`.

Value

An object of class `parameter_list` (see [numericlist](#)).

See Also[inventory\(\)](#)**Examples**

```
## example("param")

mod <- house()

param(mod)

param(mod, .pat="^(C|F)") ## may be useful when large number of parameters

class(param(mod))

param(mod)$KA

param(mod)[["KA"]]

as.list(param(mod))

as.data.frame(param(mod))

mod <- param(mod, CL = 1.2)

new_values <- list(CL = 1.3, VC = 20.5)

mod <- param(mod, new_values)
```

param_tags

Return parameter tags

Description

Use this function if you added the @covariates or @input attributes or specified a user-defined tag (via @tag) in one or more parameter blocks and need to extract that information. Also, using the \$INPUT block to declare parameters will automatically add the input tag (via @input). Once these attributes / tags are added, you can use [check_data_names\(\)](#) to reconcile names of input data sets against tagged model parameters.

Usage

```
param_tags(x)
```

Arguments

x mrgsolve model object.

Value

A data frame listing parameter names and their tags.

Model specification

Note: it is good practice to tag parameters where appropriate with input or covariates as these will automatically be expected on input data when you call `check_data_names()`. User-defined tags are also possible, but you will need to alert `check_data_names()` to look for them.

Model Specification Examples

You can use the \$INPUT block to add the input tag on these parameters

```
$INPUT  
STUDY = 101, WT = 70, DVID = 1
```

Tag some covariates in the model

```
$PARAM @covariates  
WT = 70, SEX = 1, EGFR = 110
```

A user-defined tag

```
$PARAM @tag flags  
FFLAG = 1, DFLAG = 0
```

See Also

[check_data_names\(\)](#)

Examples

```
mod <- mrgsolve::house()  
  
param_tags(mod)
```

PKMODEL

Parse PKMODEL BLOCK data

Description

Parse PKMODEL BLOCK data

Usage

```
PKMODEL(
  ncmt = 1,
  depot = FALSE,
  cmt = NULL,
  trans = pick_trans(ncmt, depot),
  env = list(),
  pos = 1,
  ...
)
```

Arguments

ncmt	number of compartments; must be 1 (one-compartment, not including a depot dosing compartment) or 2 (two-compartment model, not including a depot dosing compartment)
depot	logical indicating whether to add depot compartment
cmt	compartment names as comma-delimited character
trans	the parameterization for the PK model; must be 1, 2, 4, or 11
env	parse environment
pos	block position number
...	not used

Details

When using `$PKMODEL`, certain symbols must be defined in the model specification depending on the value of `ncmt`, `depot` and `trans`.

- `ncmt 1, depot FALSE, trans 2`: `CL, V`
- `ncmt 1, depot TRUE, trans 2`: `CL, V, KA`
- `ncmt 2, depot FALSE, trans 4`: `CL, V1, Q, V2`
- `ncmt 2, depot TRUE, trans 4`: `CL, V2, Q, V3, KA`

If `trans=11` is specified, use the symbols listed above for the `ncmt / depot` combination, but append `i` at the end (e.g. `CLi` or `Qi` or `KAi`).

If `trans=1`, the user must utilize the following symbols:

- `pred_CL` for clearance
- `pred_V` or `pred_V2` for central compartment volume of distribution
- `pred_Q` for intercompartmental clearance
- `pred_V3` for peripheral compartment volume of distribution
- `pred_KA` for absorption rate constant

See Also

[BLOCK_PARSE](#)

plot, batch_mrgsims, missing-method

Plot method for mrgsims objects

Description

Plot method for mrgsims objects

Usage

```
## S4 method for signature 'batch_mrgsims,missing'
plot(x, yval = variables(x), auto.key = list(), mincol = 3, ...)

## S4 method for signature 'batch_mrgsims,formula'
plot(
  x,
  y,
  show.grid = TRUE,
  lwd = 2,
  type = "l",
  yval = variables(x),
  auto.key = list(columns = 1),
  scales = list(y = list(relation = "free")),
  ...
)
```

Arguments

x	mrgsims object
yval	y variables to plot
auto.key	passed to xyplot
mincol	minimum number of columns in key
...	arguments passed to xyplot
y	a formula passed to xyplot
show.grid	print grid in the plot
lwd	passed to xyplot
type	passed to xyplot
scales	passed to xyplot

plot_mrgsims	<i>Generate a quick plot of simulated data</i>
--------------	--

Description

Generate a quick plot of simulated data

Usage

```
## S4 method for signature 'mrgsims,missing'
plot(x, limit = 16, ...)

## S4 method for signature 'mrgsims,formula'
plot(
  x,
  y,
  limit = 16,
  show.grid = TRUE,
  outer = TRUE,
  type = "l",
  lwd = 2,
  ylab = "value",
  groups = ID,
  scales = list(y = list(relation = "free")),
  logy = FALSE,
  logbr = 1,
  ...
)

## S4 method for signature 'mrgsims,character'
plot(x, y, ...)
```

Arguments

x	mrgsims object
limit	limit the the number of panels to create
...	other arguments passed to xyplot
y	formula used for plotting
show.grid	logical indicating whether or not to draw panel.grid
outer	passed to xyplot
type	passed to xyplot
lwd	passed to xyplot
ylab	passed to xyplot
groups	passed to xyplot

scales	passed to xyplot
logy	plot the y variables on log scale
logbr	log scale breaks indicator; use 1 for breaks every log unit; use 3 for breaks every half log unit; use 0 for default breaks

Examples

```
mod <- mrgsolve::house(end=48, delta=0.2) %>% init(GUT=1000)

out <- mrgsim(mod)

plot(out)

plot(out, subset=time <=24)

plot(out, GUT+CP~.)

plot(out, CP+RESP~time, col="black", scales="same", lty=2)

## Not run:
plot(out, "CP RESP, GUT")

## End(Not run)
```

plot_sims	<i>Plot data as an mrgsims object</i>
-----------	---------------------------------------

Description

Plot data as an mrgsims object

Usage

```
plot_sims(.data, ..., .f = NULL, .dots = list())
```

Arguments

.data	a data frame
...	unquoted column names to plot on y-axis
.f	a formula to plot
.dots	extra arguments passed to lattice::xyplot

Details

This function is only intended for use with data frames that were created by modifying an mrgsims object.

Examples

```
mod <- mrgsolve::house() %>% ev(amt = 100)

out <- mrgsim(mod)
out_df <- dplyr::mutate(out, time <= 72)

plot(out)
plot_sims(out, CP, RESP)

## Not run:
plot_sims(out, .f = ~ CP + RESP)
plot_sims(out, .f = CP + RESP ~ time)

## End(Not run)
```

qsim

Basic, simple simulation from model object

Description

This is just a lighter version of `mrgsim()`, with fewer options but with better efficiency in certain cases. See **Details**.

Usage

```
qsim(
  x,
  data,
  idata = no_idata_set(),
  obsonly = FALSE,
  tgrid = NULL,
  recsort = 1,
  tad = FALSE,
  Req = NULL,
  outvars = Req,
  skip_init_calc = FALSE,
  output = "mrgsims"
)
```

Arguments

<code>x</code>	the model object.
<code>data</code>	can be either event object or data set.
<code>idata</code>	a matrix or data frame of model parameters, one parameter per row (see <code>idata_set()</code>).
<code>obsonly</code>	if TRUE, dosing records are not included in the output.

tgrid	a tgrid object; or a numeric vector of simulation times or another object with an stime method.
recsort	record sorting flag. Default value is 1. Possible values are 1,2,3,4: 1 and 2 put doses in a data set after padded observations at the same time; 3 and 4 put those doses before padded observations at the same time. 2 and 4 will put doses scheduled through addl after observations at the same time; 1 and 3 put doses scheduled through addl before observations at the same time. recsort will not change the order of your input data set if both doses and observations are given.
tad	when TRUE a column is added to simulated output is added showing the time since the last dose. Only data records with evid == 1 will be considered doses for the purposes of tad calculation. The tad can be properly calculated with a dosing lag time in the model as long as the dosing lag time (specified in \$MAIN) is always appropriate for any subsequent doses scheduled through addl. This will always be true if the lag time doesn't change over time. But it might (possibly) not hold if the lag time changes prior to the last dose in the addl sequence. This known limitation shouldn't affect tad calculation in most common dosing lag time implementations.
Req	synonym for outvars.
outvars	output items to request; if missing, then only captured items will be returned in the output.
skip_init_calc	don't use \$MAIN to calculate initial conditions.
output	output data type; the default is mrgsim, which returns the default output object; other options include df (for data.frame) or matrix.

Details

qsim() mainly cuts some of the overhead from the simulation. So, the primary efficiency gain from using qsim() comes when the simulation executes very quickly. It is unlikely you will see a big performance difference between qsim() and [mrgsim\(\)](#) when the model is difficult to solve or if there is a large input data set.

There is no pipeline interface for this function; all configuration options (see **Arguments**) must be passed as formal arguments to the function. You can't carry_out, Request specific columns, or pass items in for update. Some other limitations, but only convenience-related. See **Arguments** for available options. Specifically, there is no ... argument for this function. Use the [update\(\)](#) method to update the model object.

See Also

[mrgsim_q\(\)](#), [mrgsim\(\)](#), [mrgsim_variants](#)

Examples

```
mod <- mrgsolve::house()

dose <- ev(amt = 100)

out <- qsim(mod,dose)
```

read_nmext

*Extract estimates from NONMEM ext file***Description**

This function retrieves NONMEM estimates for use in the mrgsolve model when \$NMEXT is invoked. See [nmext\(\)](#).

Usage

```
read_nmext(
  run = NA_real_,
  project = getwd(),
  file = paste0(run, ".ext"),
  path = NULL,
  read_fun = c("data.table", "read.table"),
  index = "last"
)
```

Arguments

run	a run number or run identifier.
project	the NONMEM project directory.
file	the ext file name.
path	full path and file name for ext file.
read_fun	function to read the ext file; data.table::fread() will be used if available; otherwise utils::read.table() is used.
index	selects the table number whose results will be returned; use value "last" to select the last table in the .ext file; or pass an integer specifying the table number; in case there is exactly one table in the .ext file, pass the value "single" to bypass parsing the file to look for sub tables (this might be useful when BAYES analysis was performed as the only estimation method and there are 10000s of posterior samples in the file).

Value

A list with param, omega, and sigma in a format ready to be used to update a model object.

Examples

```
project <- system.file("nonmem", package = "mrgsolve")

est <- read_nmext(1005, project = project)

est$param
```

```

est$omega

est$sigma

est <- read_nmext(2005, project = project, index = 3)

```

realize_addl	<i>Make addl doses explicit in an event object or data set</i>
--------------	--

Description

When doses are scheduled with `ii` and `addl`, the object is expanded to include one record for every dose. In the result, no record will have `ii` or `addl` set to non-zero value.

Usage

```

realize_addl(x, ...)

## S3 method for class 'data.frame'
realize_addl(
  x,
  warn = FALSE,
  mark_new = FALSE,
  fill = c("inherit", "na", "locf"),
  ...
)

## S3 method for class 'ev'
realize_addl(x, ...)

```

Arguments

<code>x</code>	a <code>data_set</code> data frame or an event object (see Details).
<code>...</code>	not used.
<code>warn</code>	if TRUE a warning is issued if no ADDL or <code>addl</code> column is found.
<code>mark_new</code>	if TRUE, a flag is added to indicate new columns.
<code>fill</code>	specifies how to handle non-dose related data columns in new data set records; this option is critical when handling data sets with time-varying, non-dose-related data items; see Details .

Details

If no `addl` column is found the data frame is returned and a warning is issued if `warn` is true. If `ii`, `time`, or `evid` are missing, an error is generated.

If a grouped data.frame (via `dplyr::group_by()`) is passed, it will be ungrouped.

Use caution when passing in data that has non-dose-related data columns that vary within a subject and pay special attention to the `fill` argument. By definition, `realize_addl()` will add new rows to your data frame and it is not obvious how the non-dose-related data should be handled in these new rows. When `inherit` is chosen, the new records have non-dose-related data that is identical to the originating dose record. This should be fine when these data items are not varying with time, but will present a problem when the data are varying with time. When `locf` is chosen, the missing data are filled in with NA and an last observation carry forward operation is applied to **every** column in the data set. This may not be what you want if you already had missing values in the input data set and want to preserve that missingness. When `na` is chosen, the missing data are filled in with NA and no `locf` operation is applied. But note that these missing values may be problematic for a `mrgsolve` simulation run. If you have any time-varying columns or missing data in your data set, be sure to check that the output from this function is what you were expecting.

Value

A `data.frame` or event object, consistent with the type of `x`. The `ii` and `addl` columns will all be set to zero. The result is always ungrouped.

Examples

```
e <- ev(amt = 100, ii = 12, addl = 3)

realize_addl(e)

a <- ev(amt = 100, ii = 12, addl = 2, WT = 69)
b <- ev(amt = 200, ii = 24, addl = 2, WT = 70)
c <- ev(amt = 50, ii = 6, addl = 2, WT = 71)

e <- ev_seq(a,b,c)
realize_addl(e, mark_new = TRUE)
```

render

Render a model to a document

Description

Render a model to a document

Usage

```
render(x, ...)

## S4 method for signature 'character'
render(x, project = NULL, ...)

## S4 method for signature 'mrgmod'
render(x, ...)

dorender(model, project, template = NULL, compile = TRUE, ...)
```

Arguments

x	model object or the model name
...	passed to rmarkdown::render
project	the directory containing the .cpp model file
model	model name
template	template document
compile	logical; if true, the model will be compiled to run

Examples

```
## Not run:
mod <- mrgsolve::house()
mrgsolve::render(mod)
mrgsolve::render("irm2", modlib())

## End(Not run)
```

Req	<i>Request simulated output</i>
-----	---------------------------------

Description

Use this function to select, by name, either compartments or derived variables that have been captured (see [CAPTURE](#)) into the simulated output.

Usage

```
Req(x, ...)

req(x, ...)

## S3 method for class 'mrgmod'
req(x, ...)
```

Arguments

x	model object.
...	unquoted names of compartments or tabled items.

Details

There is also a `Req` argument to `mrgsim()` that can be set to accomplish the same thing as a call to `Req` in the pipeline.

Note the difference between `req` and `Req`: the former only selects compartments to appear in output while the latter selects both compartments and captured items. Also, when there are items explicitly listed in `Req`, all other compartments or captured items not listed there are ignored. But when compartments are selected with `req` all of the captured items are returned. Remember that `req` is strictly for compartments.

Examples

```
mod <- mrgsolve::house()
mod %>% Req(CP,RESP) %>% ev(amt=1000) %>% mrgsim()
```

reserved

Reserved words

Description

Reserved words

Usage

```
reserved()
```

Details

Note: this function is not exported; you must go into the `mrgsolve` namespace by using the `mrgsolve::` prefix.

Examples

```
mrgsolve::reserved()
```

reset_tol	<i>Reset all model tolerances</i>
-----------	-----------------------------------

Description

These functions reset both scalar and customized values for both relative and absolute tolerances. All functions reset tolerances to a single, common `rtol` or `atol`. The functions do *not* change which tolerance configuration is used for simulation (e.g., scalar or customized); see [use_custom_tol\(\)](#) and [use_scalar_tol\(\)](#) to make that change in the model object.

Usage

```
reset_tol(x, rtol = NULL, atol = NULL)

reset_rtol(x, rtol = NULL)

reset_atol(x, atol = NULL)
```

Arguments

<code>x</code>	a model object.
<code>rtol</code>	global relative tolerance for both scalar and customized configurations; if not supplied, the current model's scalar <code>rtol</code> value is used.
<code>atol</code>	global absolute tolerance for both scalar and customized configurations; if not supplied, the current model's scalar <code>atol</code> value is used.

Value

An updated model object.

See Also

[custom_tol\(\)](#), [use_custom_tol\(\)](#), [use_scalar_tol\(\)](#), [get_tol\(\)](#)

Examples

```
mod <- house()
mod <- reset_tol(mod, rtol = 1e-6, atol = 1e-10)
mod
```

`revar`*Get model random effect variances and covariances*

Description

Use this function to extract both OMEGA and SIGMA matrices from a model object. Typical use is for display on the R console.

Usage

```
revar(x, ...)  
  
## S4 method for signature 'mrgmod'  
revar(x, ...)
```

Arguments

<code>x</code>	model object.
<code>...</code>	passed along.

Value

A named list containing omega and sigma matrices.

Examples

```
mod <- mrgsolve::house()  
revar(mod)
```

`see`*Print model code to the console*

Description

This is a simple way to display the model code on the R console using the model object. The `raw` argument will return the model code as a character vector.

Usage

```
see(x, ...)  
  
## S4 method for signature 'mrgmod'  
see(x, raw = FALSE, ...)
```

Arguments

x	model object.
...	not used.
raw	return the raw code.

Value

NULL is returned invisibly when raw is FALSE; when raw is set to TRUE, the model code is returned as a character vector.

sigma	<i>Manipulate SIGMA matrices</i>
-------	----------------------------------

Description

The primary function is `smat()` which can be used to both get the \$SIGMA matrices out of a model object and to update \$SIGMA matrices in a model object.

Usage

```
smat(.x, ...)
```

S4 method for signature 'missing'

```
smat(.x, ...)
```

S4 method for signature 'matrix'

```
smat(.x, ..., labels = list())
```

S4 method for signature 'list'

```
smat(.x, ...)
```

S4 method for signature 'sigmalist'

```
smat(.x, ...)
```

S4 method for signature 'mrgmod'

```
smat(.x, ..., make = FALSE, open = FALSE)
```

S4 method for signature 'NULL'

```
smat(.x, ...)
```

S4 method for signature 'mrgsims'

```
smat(.x, make = FALSE, ...)
```

Arguments

<code>.x</code>	a matrix, list of matrices or <code>matlist</code> object.
<code>...</code>	passed to other functions, including <code>modMATRIX()</code> .
<code>labels</code>	character vector of names for <code>\$SIGMA</code> elements; must be equal to number of rows/columns in the matrix.
<code>make</code>	logical; if <code>TRUE</code> , matrix list is rendered into a single matrix.
<code>open</code>	passed to <code>merge.list()</code> .
<code>x</code>	<code>matlist</code> object.

See Also

`dmat()`, `bmat()`, `cmat()`

Examples

```
## example("sigma")
mat1 <- matrix(1)
mat2 <- diag(c(1,2))
mat3 <- matrix(c(0.1, 0.002, 0.002, 0.5), 2,2)
mat4 <- dmat(0.1, 0.2, 0.3, 0.4)

smat(mat1)
smat(mat1, mat2, mat3)
smat(A=mat1, B=mat2, C=mat3)

mod <- mrgsolve::house() %>% smat(mat1)

smat(mod)
smat(mod, make=TRUE)
```

simargs

Access or clear arguments for calls to `mrgsim()`

Description

As a model object navigates a pipeline prior to simulation, arguments are collected to eventually be passed to `mrgsim()`. `simargs()` lets you intercept and possibly clear those arguments.

Usage

```
simargs(x, which = NULL, clear = FALSE, ...)
```

Arguments

x	model object.
which	character with length 1 naming a single arg to get.
clear	logical indicating whether or not to clear args from the model object.
...	not used.

Value

If clear is TRUE, the argument list is cleared and the model object is returned. Otherwise, the argument list is returned.

Examples

```
mod <- mrgsolve::house()
mod %>% Req(CP, RESP) %>% carry_out(evid, WT, FLAG) %>% simargs()
```

soloc

Return the location of the model shared object

Description

This is also the directory where the model is built, which could be the value of `tempdir()`.

Usage

```
soloc(x, short = FALSE)
```

Arguments

x	model object.
short	logical; if TRUE, solocs will be rendered with a short path name.

Value

A string containing the full path to the model shared object.

Examples

```
mod <- mrgsolve::house()
soloc(mod)
```

solvsettings	<i>Optional inputs for lsoda</i>
--------------	----------------------------------

Description

These are settings for the differential equation solver (lsoda) that can be accessed via the R interface. The code listing below is taken directly from the lsoda source code.

Details

The following items can be set

- hmax (HMAX below); decrease hmax when you want to limit how big of a step the solver can take when integrating from one time to the next time. However be aware that smaller hmax will result in longer run times.
- hmin (HMIN below); don't fiddle with this unless you know what you're doing.
- ixpr (IXPR below)
- maxsteps (MXSTEP below); increase this number when the solver has a long interval between two integration times (e.g. when observation records are far apart).
- mxhnil (MXHNIL below); don't usually modify this one
- atol - the absolute solver tolerance; decrease this number (e.g. to 1E-10 or 1E-20 or 1E-50) when the value in a compartment can get extremely small; without this extra (lower) tolerance, the value can get so low that the number can randomly become negative. However be aware that more precision here will result in longer run times.
- rtol - the relative solver tolerances; decrease this number when you want a more precise solution. However be aware that more precision here will result in longer run times.

See Also

[aboutsolver](#), [update](#)

summary.mrgmod	<i>Print summary of a mrgmod object</i>
----------------	---

Description

Print summary of a mrgmod object

Usage

```
## S3 method for class 'mrgmod'
summary(object, ...)
```

Arguments

object	a mrgmod object
...	not used

tscale	<i>Re-scale time in the simulated output</i>
--------	--

Description

Re-scale time in the simulated output

Usage

```
tscale(x, value = 1, ...)
```

Arguments

x	model object.
value	value by which time will be scaled.
...	not used.

Details

There is also a `tscale` argument to `mrgsim()` that can be set to accomplish the same thing as a call to `tscale` in the pipeline.

Examples

```
# The model is in hours:
mod <- mrgsolve::house()

# The output is in days:
mod %>% tscale(1/24) %>% mrgsim()
```

update	<i>Update the model object</i>
--------	--------------------------------

Description

After the model object is created, update various attributes.

Usage

```
## S4 method for signature 'mrgmod'
update(object, ..., merge = TRUE, open = FALSE, data = NULL, strict = TRUE)

## S4 method for signature 'omegalist'
update(object, y, ...)

## S4 method for signature 'signalist'
update(object, y, ...)

## S4 method for signature 'parameter_list'
update(object, .y, ...)
```

Arguments

object	a model object.
...	named items to update.
merge	logical indicating to merge (rather than replace) new and existing attributes.
open	logical; used only when merge is TRUE and parameter list or initial conditions list is being updated; if FALSE, no new items will be added; if TRUE, the parameter list may expand.
data	a list of items to update; this list is combined with any items passed in via ...
strict	if TRUE, a warning will be issued when there is an attempt to update a non-existent item.
y	another object involved in update
.y	data to update

Details

Slots that can be updated:

- verbose
- debug
- preclean
- mindt
- digits

- atol - absolute solver tolerance; see [solversettings](#)
- rtol - relative solver tolerance; see [solversettings](#)
- ss_rtol - relative tolerance when finding steady state
- ss_atol - absolute tolerance when finding steady state
- ixpr - see IXPR in [solversettings](#)
- mxhnil - see MXHNIL in [solversettings](#)
- hmin - see HMIN in [solversettings](#)
- hmax - see HMAX in [solversettings](#)
- maxsteps - see MXSTEP in [solversettings](#)
- start, end, delta, add
- tscale
- request
- param
- init
- omega
- sigma
- outvars

Value

The updated model object is returned.

See Also

[update](#), [mrgmod-class](#), [within](#)

Examples

```
## Not run:  
mod <- house()  
  
mod <- update(mod, end = 120, delta = 4, param = list(CL = 19.1))  
  
## End(Not run)
```

`use_custom_tol`*Set up a model object to use either scalar or custom tolerances*

Description

Call `use_custom_tol()` to use custom relative and absolute tolerances in a model; call `use_scalar_tol()` to revert to the traditional configuration where a single `rtol` and `atol` are applied to all compartments.

Usage

```
use_custom_tol(x)
```

```
use_scalar_tol(x)
```

Arguments

`x` a model object.

Details

If customized tolerances have not been initialized yet, they will be, assigning the current `rtol` or `atol` for every compartment. These default values can be updated using `custom_rtol()`, `custom_atol()`, or `custom_tol()`.

Value

An updated model object.

See Also

`custom_tol()`, `reset_tol()`, `get_tol()`

Examples

```
mod <- house()

mod <- use_custom_tol(mod)
mod

mod <- use_scalar_tol(mod)
mod
```

valid_data_set	<i>Validate and prepare data sets for simulation</i>
----------------	--

Description

This function is called by `mrgsim()` and friends to check and prepare input data sets for simulation. Users may also call this function to pre-validate data when the same data set is used for repeated simulation.

Usage

```
valid_data_set(x, m = NULL, verbose = FALSE, quiet = FALSE)
```

```
valid_data_set.matrix(x, verbose = FALSE)
```

Arguments

x	data.frame or matrix.
m	a model object.
verbose	logical.
quiet	if TRUE, messages will be suppressed.

Details

An error will be issued when

- non-numeric data is found in columns sharing names with model parameters
- non-numeric data is found in reserved data items related to dosing (see `mrgsolve:::GLOBALS$CARRY_TRAN`)
- a column is found that is "internally classed", including columns that inherit from `integer64` (see `is.object()`)

Value

A matrix with non-numeric columns dropped; if x is a data.frame with character `cmt` column comprised of valid compartment names and m is a model object, the `cmt` column will be converted to the corresponding compartment number.

See Also

`valid_idata_set()`, `idata_set()`, `data_set()`

Examples

```
mod <- mrgsolve::house()

data(exTheoph)

d <- valid_data_set(exTheoph, mod)
```

valid_idata_set	<i>Validate and prepare idata data sets for simulation</i>
-----------------	--

Description

This function is called by `mrgsim()` and friends to check and prepare input data sets for simulation. Users may also call this function to pre-validate data when the same data set is used for repeated simulation.

Usage

```
valid_idata_set(x, m, verbose = FALSE, quiet = FALSE)
```

Arguments

x	data.frame or matrix.
m	a model object.
verbose	logical.
quiet	if TRUE, messages will be suppressed.

Details

An error will be issued when

- non-numeric data is found in columns sharing names with model parameters
- a column is found that is internally classed, including columns that inherit from `integer64` (see `is.object()`)

Value

A numeric matrix with class `valid_idata_set`.

See Also

`valid_data_set()`, `idata_set()`, `data_set()`

within	<i>Update parameters, initials, and settings within a model object</i>
--------	--

Description

The main use case for using [within](#) rather than [update](#) or [param](#) or [init](#) is when you want to update to a new value that is calculated from the existing value. See the example in details

Usage

```
## S3 method for class 'mrgmod'  
within(data, expr, ...)
```

Arguments

data	an object with class mrgmod
expr	expressions evaluated in an environment containing various model object components, including parameters, initial conditions, and others (see details)
...	not used

Details

Other model object slots that can be updated: start, end, delta, add, rtol, atol, hmax, maxsteps. These are include for convenience, but we expect that most of the time these will get updated through the update method.

See Also

[update](#)

Examples

```
mod <- mrgsolve::house()  
  
mod2 <- within(mod, {CL <- CL * 1.5})  
  
mod$CL  
mod2$CL
```

zero_re	<i>Zero out random effects in a model object</i>
---------	--

Description

Sets all elements of the OMEGA or SIGMA matrix to zero.

Usage

```
zero_re(.x, ...)

## S4 method for signature 'mrgmod'
zero_re(.x, ...)
```

Arguments

.x	a model object.
...	which matrix to zero out; pass omega to just zero out omega, sigma to just zero out sigma; passing nothing will zero out both.

Value

An updated object with elements of OMEGA and/or SIGMA set to zero.

Examples

```
mod <- house()
revar(mod)
mod <- zero_re(mod)
revar(mod)

## Not run:
mod <- modlib("popex", compile = FALSE)
mod <- zero_re(mod, omega)
revar(mod)

## End(Not run)
```

\$,ev-method	<i>Select columns from an ev object</i>
--------------	---

Description

Select columns from an ev object

Usage

```
## S4 method for signature 'ev'
x$name

## S4 method for signature 'ev'
x[[i, exact = TRUE]]
```

Arguments

x	ev object
name	column to select
i	an element to select
exact	not used

<code>\$,mrgmod-method</code>	<i>Select parameter values from a model object</i>
-------------------------------	--

Description

The `$` and `[[` operators get the value of a single parameter in the model. The `[` gets several values, returning a named list.

Usage

```
## S4 method for signature 'mrgmod'
x$name

## S4 method for signature 'mrgmod'
x[[i, exact = TRUE]]

## S4 method for signature 'mrgmod'
x[i]
```

Arguments

x	mrgmod object
name	parameter to take
i	an element to select
exact	not used

Index

- * **datasets**
 - exdatasets, [43](#)
- * **param**
 - param, [85](#)
- *, tgrid, numeric-method
 - (c, tgrid-method), [19](#)
- *, tgrids, numeric-method
 - (c, tgrid-method), [19](#)
- +, tgrid, numeric-method
 - (c, tgrid-method), [19](#)
- +, tgrids, numeric-method
 - (c, tgrid-method), [19](#)
- [, mrgmod-method (\$, mrgmod-method), [114](#)
- [[, ev-method (\$, ev-method), [113](#)
- [[, mrgmod-method (\$, mrgmod-method), [114](#)
- \$.ev-method, [113](#)
- \$.mrgmod-method, [114](#)

- aboutsolver, [4](#), [7](#), [105](#)
- allparam (param), [85](#)
- as.data.frame(), [74](#)
- as.ev, [8](#)
- as.ev(), [34](#), [38](#)
- as.ev, data.frame-method (as.ev), [8](#)
- as.ev, ev-method (as.ev), [8](#)
- as.evd (evd), [35](#)
- as.evd(), [34](#)
- as.integer(), [82](#)
- as.list, mrgmod-method, [9](#)
- as.list, mrgsims-method, [11](#)
- as.tbl.mrgsims (mrgsims_dplyr), [70](#)
- as_bmat, [11](#)
- as_bmat(), [55](#)
- as_bmat, ANY-method (as_bmat), [11](#)
- as_bmat, data.frame-method (as_bmat), [11](#)
- as_bmat, list-method (as_bmat), [11](#)
- as_bmat, numeric-method (as_bmat), [11](#)
- as_cmat (as_bmat), [11](#)
- as_data_frame.mrgsims (mrgsims_dplyr), [70](#)

- as_data_set, [13](#)
- as_data_set(), [74](#)
- as_data_set, data.frame-method (as_data_set), [13](#)
- as_deslist, [14](#)
- as_dmat (as_bmat), [11](#)
- as_dmat(), [55](#)
- as_dmat, ANY-method (as_bmat), [11](#)
- as_dmat, data.frame-method (as_bmat), [11](#)
- as_dmat, list-method (as_bmat), [11](#)
- as_dmat, numeric-method (as_bmat), [11](#)
- as_tibble.mrgsims (mrgsims_dplyr), [70](#)
- assign_ev (ev_assign), [36](#)

- BLOCK_PARSE, [16](#), [89](#)
- blocks, [15](#)
- blocks, character-method (blocks), [15](#)
- blocks, mrgmod-method (blocks), [15](#)
- bmat (matrix_helpers), [54](#)
- bmat(), [13](#), [84](#), [103](#)

- c, matlist-method, [19](#)
- c, tgrid-method, [19](#)
- c, tgrids-method (c, tgrid-method), [19](#)
- CAPTURE, [98](#)
- CAPTURE (BLOCK_PARSE), [16](#)
- carry.out (carry_out), [20](#)
- carry_out, [20](#)
- check_data_names, [21](#)
- check_data_names(), [50](#), [87](#), [88](#)
- cmat (matrix_helpers), [54](#)
- cmat(), [12](#), [13](#), [84](#), [103](#)
- CMT (BLOCK_PARSE), [16](#)
- cmtn, [22](#)
- cmtn, mrgmod-method (cmtn), [22](#)
- code, [23](#)
- collapse_matrix, [23](#)
- collapse_matrix(), [25](#)
- collapse_omega, [24](#)

collapse_omega(), 23, 24
collapse_sigma(collapse_omega), 24
collapse_sigma(), 23, 24
custom_atol(custom_tol), 25
custom_atol(), 109
custom_rtol(custom_tol), 25
custom_rtol(), 109
custom_tol, 25
custom_tol(), 46, 100, 109

data.table::fread(), 95
data_set, 26, 33, 34
data_set(), 13, 14, 20, 48, 50, 67, 75, 110, 111
data_set, mrgmod, ANY-method (data_set), 26
data_set, mrgmod, data.frame-method (data_set), 26
data_set, mrgmod, ev-method (data_set), 26
data_set, mrgmod, missing-method (data_set), 26
design, 29
details, 30
distinct.mrgsims (mrgsims_dplyr), 70
dmat (matrix_helpers), 54
dmat(), 12, 13, 84, 103
do.mrgsims (mrgsims_dplyr), 70
do_mrgsim (mrgsim), 66
do_mrgsim(), 67, 75
dorender (render), 97
dplyr::distinct, 71
dplyr::filter(), 27, 47
dplyr::group_by, 71
dplyr::group_by(), 96
dplyr::select(), 27, 47
dplyr::summarise_each, 71
dplyr::vars(), 27, 47

env_eval, 31
env_get, 31
env_get(), 31
env_get_env (env_get), 31
env_ls, 32
env_ls(), 31
env_update, 32
ev, 33
ev(), 14, 28, 35, 40, 47, 48
ev, ev-method (ev), 33
ev, missing-method (ev), 33
ev, mrgmod-method (ev), 33
ev_assign, 36
ev_assign(), 34
ev_days, 37
ev_days(), 34
ev_expand (expand.idata), 44
ev_methods, 34
ev_rep, 38
ev_rep(), 34
ev_repeat, 39
ev_repeat(), 34, 38
ev_rx, 39
ev_rx, character, missing-method (ev_rx), 39
ev_rx, mrgmod, character-method (ev_rx), 39
ev_seq, 41
ev_seq(), 34, 41
evd, 35
evd(), 14, 34, 35
evd, ev-method (evd), 35
evd, missing-method (evd), 35
evd, mrgmod-method (evd), 35
evd_expand (expand.idata), 44
exBoot (exdatasets), 43
exdatasets, 28, 43
exidata (exdatasets), 43
expand.ev (expand.idata), 44
expand.ev(), 14
expand.evd (expand.idata), 44
expand.evd(), 14
expand.grid(), 44
expand.idata, 44
expand_observations, 45
exTheoph (exdatasets), 43
extran1 (exdatasets), 43
extran2 (exdatasets), 43
extran3 (exdatasets), 43

filter.ev (mutate.ev), 76
filter.mrgsims (mrgsims_dplyr), 70
filter_sims (mrgsims_modify), 72
FIXED (BLOCK_PARSE), 16

get_tol, 46
get_tol(), 26, 100, 109
get_tol_list (get_tol), 46
group_by.mrgsims (mrgsims_dplyr), 70

- HANDLEMATRIX (BLOCK_PARSE), 16
- idata_set, 46
- idata_set(), 20, 28, 50, 67, 75, 93, 110, 111
- idata_set, mrgmod, ANY-method (idata_set), 46
- idata_set, mrgmod, data.frame-method (idata_set), 46
- idata_set, mrgmod, missing-method (idata_set), 46
- INIT (BLOCK_PARSE), 16
- init, 48, 112
- init(), 28
- init, ANY-method (init), 48
- init, list-method (init), 48
- init, missing-method (init), 48
- init, mrgmod-method (init), 48
- init, mrgsims-method (init), 48
- inventory, 50
- inventory(), 27, 47, 86, 87
- is.mrgmod, 51
- is.mrgsims, 51
- is.object(), 110, 111
- lctran, 52
- lctran(), 14, 28, 35
- loadso, 53
- ls(), 32
- matrix_helpers, 54
- mcode, 55
- mcode(), 64
- mcode_cache (mcode), 55
- mcode_cache(), 55, 64
- mcRNG, 56
- merge.list(), 84, 103
- modlib, 56
- modlib(), 63, 64
- modlib_details, 57, 57, 59, 64
- modlib_pk, 57, 58, 64
- modlib_pkpd, 57, 59, 64
- modlib_tmdd, 57, 60, 64
- modlib_viral, 57, 61, 64
- modMATRIX(), 84, 103
- mread, 62
- mread(), 5, 55, 65, 76, 78
- mread_cache (mread), 62
- mread_cache(), 55, 56
- mread_file (mread), 62
- mread_yaml, 65
- mread_yaml(), 77, 78
- mrgsim, 66, 82
- mrgsim(), 20, 73–76, 93, 94, 99, 103, 106, 110, 111
- mrgsim_0 (mrgsim_variants), 75
- mrgsim_d (mrgsim_variants), 75
- mrgsim_d(), 73
- mrgsim_df (mrgsim), 66
- mrgsim_df(), 68
- mrgsim_di (mrgsim_variants), 75
- mrgsim_e (mrgsim_variants), 75
- mrgsim_ei (mrgsim_variants), 75
- mrgsim_ei(), 74
- mrgsim_i (mrgsim_variants), 75
- mrgsim_i(), 74
- mrgsim_q, 73
- mrgsim_q(), 66, 69, 75, 76, 94
- mrgsim_variants, 66, 69, 74, 75, 94
- mrgsims, 69
- mrgsims_dplyr, 70, 72
- mrgsims_modify, 70, 72, 72
- mrgsolve (mrgsolve-package), 4
- mrgsolve-package, 4
- mutate.ev, 76
- mutate.ev(), 34
- mutate.mrgsims (mrgsims_dplyr), 70
- mutate_sims (mrgsims_modify), 72
- mwrite_cpp, 76
- mwrite_yaml, 77
- mwrite_yaml(), 65, 77
- names, mrgmod-method, 78
- nmext, 79
- nmext(), 95
- NMXML (nmxml), 80
- nmxml, 80
- nmxml(), 80
- numericlist, 48, 49, 85, 86
- numerics_only, 82
- numerics_only(), 28
- obsaug, 82
- obsonly, 83
- omat (omega), 83
- omat(), 24
- omat, list-method (omega), 83
- omat, matrix-method (omega), 83
- omat, missing-method (omega), 83

- omat, mrgmod-method (omega), 83
- omat, mrgsims-method (omega), 83
- omat, NULL-method (omega), 83
- omat, omegalist-method (omega), 83
- OMEGA (omega), 83
- omega, 83
- options(), 63
- outvars, 85

- PARAM (BLOCK_PARSE), 16
- param, 85, 112
- param(), 47
- param, ANY-method (param), 85
- param, list-method (param), 85
- param, missing-method (param), 85
- param, mrgmod-method (param), 85
- param, mrgsims-method (param), 85
- param_tags, 87
- param_tags(), 21, 22
- parse_rx (ev_rx), 39
- parse_rx(), 39
- PKMODEL, 88
- PKMODEL(), 16, 18
- plot, batch_mrgsims, formula-method
 - (plot, batch_mrgsims, missing-method), 90
- plot, batch_mrgsims, missing-method, 90
- plot, mrgsims, character-method
 - (plot_mrgsims), 91
- plot, mrgsims, formula-method
 - (plot_mrgsims), 91
- plot, mrgsims, missing-method
 - (plot_mrgsims), 91
- plot_mrgsims, 91
- plot_sims, 92
- pull.mrgsims (mrgsims_dplyr), 70

- qsim, 93
- qsim(), 74, 76

- read_nmext, 95
- read_nmext(), 80
- realize_addl, 96
- realize_addl(), 34
- render, 97
- render, character-method (render), 97
- render, mrgmod-method (render), 97
- Req, 98
- req (Req), 98

- reserved, 99
- reset_atol (reset_tol), 100
- reset_rtol (reset_tol), 100
- reset_tol, 100
- reset_tol(), 26, 46, 109
- revar, 101
- revar, mrgmod-method (revar), 101

- see, 59, 101
- see, mrgmod-method (see), 101
- select.ev (mutate.ev), 76
- select.mrgsims (mrgsims_dplyr), 70
- select_sims (mrgsims_modify), 72
- seq(), 41
- seq.ev (ev_seq), 41
- set.seed(), 31
- SIGMA (sigma), 102
- sigma, 102
- signif(), 54
- simargs, 103
- slice.mrgsims (mrgsims_dplyr), 70
- smat (sigma), 102
- smat(), 24, 84
- smat, list-method (sigma), 102
- smat, matrix-method (sigma), 102
- smat, missing-method (sigma), 102
- smat, mrgmod-method (sigma), 102
- smat, mrgsims-method (sigma), 102
- smat, NULL-method (sigma), 102
- smat, signalist-method (sigma), 102
- soloc, 104
- solversettings, 10, 105, 108
- stime(), 67
- summarise.each (mrgsims_dplyr), 70
- summarise.mrgsims (mrgsims_dplyr), 70
- summary.mrgmod, 105

- tempdir(), 55, 104
- tgrid, 29
- tgrid*_numeric (c, tgrid-method), 19
- tgrid+_numeric (c, tgrid-method), 19
- tgrids*_numeric (c, tgrid-method), 19
- tgrids+_numeric (c, tgrid-method), 19
- THETA (BLOCK_PARSE), 16
- tscale, 106

- uctran (lctran), 52
- uctran(), 14, 28, 35
- update, 105, 107, 108, 112

`update()`, [5](#), [63](#), [67](#), [74](#), [75](#), [94](#)
`update`, `mrgmod`-method (`update`), [107](#)
`update`, `omegalist`-method (`update`), [107](#)
`update`, `parameter_list`-method (`update`),
[107](#)
`update`, `sigmalist`-method (`update`), [107](#)
`use_custom_tol`, [109](#)
`use_custom_tol()`, [26](#), [46](#), [100](#)
`use_scalar_tol` (`use_custom_tol`), [109](#)
`use_scalar_tol()`, [26](#), [46](#), [100](#)
`utils::read.table()`, [95](#)

`valid_data_set`, [110](#)
`valid_data_set()`, [28](#), [111](#)
`valid_idata_set`, [111](#)
`valid_idata_set()`, [28](#), [110](#)

`within`, [108](#), [112](#), [112](#)
`within`, `mrgmod`-method (`within`), [112](#)
`within.mrgmod` (`within`), [112](#)

`yaml_to_cpp` (`mread_yaml`), [65](#)
`yaml_to_cpp()`, [77](#), [78](#)

`zero_re`, [113](#)
`zero_re`, `mrgmod`-method (`zero_re`), [113](#)