

Package ‘assemblerr’

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Title Assembly of Pharmacometric Models

Description Construct pharmacometric nonlinear mixed effect models by combining predefined model components and automatically generate model code for NONMEM. Models are created by combining parameter and observation models, algebraic relationships, compartments, and flows. Pharmacokinetic models can be assembled from the higher-order components: absorption, distribution, and elimination. The generated code is optimized for performance by recognizing, for example, linear differential equations or differential equations with an analytic solution.

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'pk_model.R' 'rendering.R' 'statement.R' 'nm_model.R'
'declaration-creation.R' 'declaration.R' 'algebraics.R'
'assemblerr-package.R' 'ast.R' 'compartment.R'
'conversion-compartment-nm.R' 'input_variable.R'
'conversion-input_variable-nm.R' 'meta.R'
'conversion-meta-nm.R' 'conversion-nm.R' 'observation.R'
'conversion-observation-nm.R' 'parameter.R'
'conversion-parameter-nm.R' 'node-classes.R' 'tasks.R'
'conversion-tasks-nm.R' 'description.R' 'issues.R'
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'test-helpers.R' 'util.R' 'variables.R'

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BugReports <https://github.com/UUPharmacometrics/assemblerr/issues>

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assemblerr-package *assemblerr: Assembly of Pharmacometric Models*

Description

Construct pharmacometric nonlinear mixed effect models by combining predefined model components and automatically generate model code for NONMEM. Models are created by combining parameter and observation models, algebraic relationships, compartments, and flows. Pharmacokinetic models can be assembled from the higher-order components: absorption, distribution, and elimination. The generated code is optimized for performance by recognizing, for example, linear differential equations or differential equations with an analytic solution.

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- João A. Abrantes [reviewer]
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See Also

Useful links:

- <https://github.com/UUPharmacometrics/assemblerr>
- Report bugs at <https://github.com/UUPharmacometrics/assemblerr/issues>

 algebraic

Algebraic relationship

Description

This building block defines a model variable as a function of other variables.

Usage

```
algebraic(definition)
```

Arguments

definition A definition of the model variable

Details

Algebraic relationships are equations where one variable is defined as a function of multiple other variables. `assemblerr` uses R formulas to implement these equations. For example, the Emax dose response model

$$effect = emax * dose / (ed50 + dose)$$

could be declared as

```
algebraic(effect~emax*dose/(ed50+dose))
```

where the tilde `~` replaced the equal sign `=` in the definition.

Value

A building block of type 'algebraic'

Examples

```
m <- model() +
  input_variable("dose") +
  prm_log_normal("emax", 10, 0.3) +
  prm_no_var("ed50", 5) +
  algebraic(effect~emax*dose/(ed50+dose)) +
  obs_additive(~effect)
```

assemblerr_options *Options*

Description

This function creates a list of options for the use with the render function.

Usage

```
assemblerr_options(  
  prm.use_mu_referencing = FALSE,  
  ode.use_special_advans = TRUE,  
  ode.use_general_linear_advans = TRUE,  
  ode.general_nonlinear_advan = "advan13",  
  ode.general_linear_advan = "advan5",  
  ode.preferred_trans_routines = c("trans2", "trans4"),  
  issues.missing_variables = c("fix-warn", "fix", "ignore", "fail")  
)
```

Arguments

prm.use_mu_referencing
 Use mu-referencing?

ode.use_special_advans
 Use analytic solution ADVANs?

ode.use_general_linear_advans
 Use ADVANs for linear ODEs?

ode.general_nonlinear_advan
 ADVAN to be used for non-linear ODEs

ode.general_linear_advan
 ADVAN to be used for linear ODEs

ode.preferred_trans_routines
 Order of TRANS routines to be tried

issues.missing_variables
 How to handle missing variables

Details

The function helps to create properly formatted list that can serve as input to the options= argument of the render() function.

Value

A list of options

`check`*Checking for issues*

Description

This function checks a model for existing issues.

Usage

```
check(model)
```

Arguments

`model` Model to check

Details

The function accepts a model object and returns a list of issues that can help to identify problems in a model. If no issues are found, a message and an empty list are produced. Issues can either be critical or non-critical, depending on whether a valid model could still be rendered.

The function currently detects the following issues:

- Undefined variables
- Lack of parameters
- Lack of observations
- Lack of distribution/elimination components (`pk_model`)
- Inconsistent capitalization of variable names

Value

An issue list (printed to the console by default)

Examples

```
m <- model() +
  prm_log_normal("emax") +
  prm_log_normal("ed50") +
  obs_additive(eff~emax*dose/(ed50+dose))
check(m)

# fix issue
m <- m + input_variable("dose")
check(m)
```

compartment	<i>Compartment</i>
-------------	--------------------

Description

Defines name and volume of a compartment.

Usage

```
compartment(name, volume = 1)
```

```
cmp(name, volume = 1)
```

Arguments

name	Name of the compartment
volume	Volume as a number, formula or parameter name

Details

In most applications, compartments contain kinetically homogeneous amount of drug (applications where the compartment content represents other quantities are also possible). In `assemblerr`, a compartment is defined by providing a name and the compartment volume.

Compartment names:

Every compartment must have a valid name. A compartment name can contain letters, numbers as well as the underscore character, and needs to start with a letter. Adding a compartment with an already existing name will replace the definition of the compartment.

Compartment volumes:

The compartment volume can be provided as a number, R formula, or a parameter name. It will be used by `assemblerr` to replace references to the compartment concentration (e.g., `~C["central"]`) with the corresponding amount divided by volume (e.g., `~A["central"]/vc`).

Value

A building block of type 'compartment'

See Also

[flow](#) for how to describe compartment kinetics

Examples

```
# model with depot and central compartment
m <- model() +
  compartment("depot", volume = 1) +
  compartment("central", volume = "vc") +
  flow(~ka*A, from = "depot", to = "central") +
  flow(~cl*C, from = "central") +
  prm_log_normal("ka") +
  prm_log_normal("cl") +
  prm_log_normal("vc") +
  obs_additive(conc~C["central"])

render(
  model = m,
  options = assemblerr_options(
    ode.use_special_advans = FALSE,
    ode.use_general_linear_advans = FALSE
  )
)
```

flow

Flow between compartments

Description

This building block describes a flow between compartments.

Usage

```
flow(definition, from = NA_character_, to = NA_character_)
```

Arguments

definition	Equation describing the flow
from	Name of the source compartment (NA for an inflow without source)
to	Name of the sink compartment (NA for an outflow without sink)

Details

Flows define the connections between compartments and the equations according to which exchanges occur.

Flow equations:

The first function argument is the flow equation. It is defined using R formulas that can start with the tilde \sim operator and do not need to have a left-hand side (i.e., $\sim k\theta$ is a valid flow definition).

Flow equations can contain the special variables A and C which can be used to refer to the amount and concentration in the compartment specified via the `from=` argument. For example, the following code creates a flow building block describing the first-order transfer from the depot to the central compartment


```
flow(~ka*A, "depot", "central")
```

When the model is rendered, A and C will get replaced with the corresponding compartment reference. `assemblerr` will raise an error if A or C are used without specifying the `from=` compartment (such as in an inflow).

Compartment connections:

The connection between compartments can be specified using the `from=` and `to=` arguments of the function. Setting either `from=` or `to=` to NA allows the definition of in and outflows without a source or sink. Setting both arguments to NA results in error.

Conversion to differential equations:

When flows are rendered they are converted to ordinary differential equations (ODEs). The connection between compartments together with the flow equations allow `assemblerr` to determine whether an analytic solution can be generated. This automatic optimization of differential equations can be disabled via the rendering options.

Value

A building block of type 'flow'

Examples

```
# one-compartment model with first-order elimination
m <- model() +
  prm_log_normal("v") +
  prm_log_normal("cl") +
  compartment("central", volume = ~v) +
  flow(declaration(~cl*C), from = "central") +
  obs_additive(~C["central"])
# an analytic solution is generated
render(m)

# one-compartment model with Michaelis-Menten elimination
m2 <- model() +
  prm_log_normal("v") +
  prm_log_normal("vmax") +
  prm_no_var("km") +
  compartment("central", volume = ~v) +
  flow(declaration(~vmax*C/(km+C)), from = "central") +
  obs_additive(~C["central"])

# an ODE is generated
render(m2)
```

input_variable *Input variables*

Description

These building block declare input variables, i.e., variables that are defined in the dataset.

Usage

```
input_variable(name)
```

```
dataset(path)
```

Arguments

name	Variable name
path	Dataset path

Details

An input variable is defined in the dataset and is declared so that it can be used in the rest of the model definition. The function `input_variable()` declares a single variable whereas the `dataset()` function reads the header of the file provided and declares all variables found.

Value

A building block of type 'input_variable'

Examples

```
m <- model() +  
  input_variable("dose") +  
  prm_log_normal("emax") +  
  prm_log_normal("ed50") +  
  obs_additive(eff~emax*dose/(ed50+dose))  
render(m)
```

model *General model*

Description

This function creates the basis for a general pharmacometric model, a flexible but verbose model type.

Usage

```
model()
```

Details

The function creates the fundament for a general pharmacometric model to which different building blocks can be added. The following building blocks are relevant for this model type:

- Parameters: [prm_log_normal](#), [prm_logit_normal](#), [prm_no_var](#), [prm_normal](#)
- Observations: [obs_additive](#), [obs_combined](#), [obs_proportional](#)
- Algebraic relationships: [algebraic](#)
- Compartments: [compartment](#)
- Flows: [flow](#)
- Input variables: [input_variable](#), [dataset](#)

The more specialized [pk_model\(\)](#) is converted to a general model during the rendering process.

Value

A general pharmacometric model

Examples

```
m <- model() +  
  input_variable("dose") +  
  prm_log_normal("emax") +  
  prm_log_normal("ed50") +  
  obs_additive(eff~emax*dose/(ed50+dose))  
render(m)
```

model-variable-selection

Selecting model variables

Description

The output task allows to select model variables using a concise mini language. You can select variables by name or using one of the helper functions described below.

Overview of selection features:

The selection of variables builds on the [tidyselect](#) package which implements a powerful variable selection language (see [tidyselect::language](#)). The following features are most relevant for the selection of model variables:

- | for selecting the union of several variables
- c() for combining selections
- ! for taking the complement of a set of variables

In addition, you can select variables using a combination of the following helper functions:

- `vars_prms()` selects all model parameters
- `vars_data()` selects all data defined variables
- `vars_eta()` selects all eta variables
- `vars_nm_std()` selects the standard NONMEM variables DV, PRED, RES, WRES, IPREDI, IWRESI
- `vars_starts_with()` selects variables that start with a prefix
- `vars_matches()` selects variables that match a regular expression

Usage

```
vars_prms(vars)
```

```
vars_data(vars)
```

```
vars_eta(vars)
```

```
vars_nm_std(vars)
```

```
vars_starts_with(match, vars)
```

```
vars_matches(match, vars)
```

Arguments

<code>vars</code>	A character vector of variable names (taken from the selection context)
<code>match</code>	A character vector to match against

Value

A selection context

Examples

```
m <- model() +  
  input_variable("dose") +  
  prm_log_normal("emax", median = 10, var_log = 0.09) +  
  prm_log_normal("ed50", median = 50, var_log = 0.09) +  
  algebraic(effect~emax*dose/(ed50 + dose)) +  
  obs_proportional(~effect, var_prop = 1)  
  
# output all model parameter and eta variables  
render(m, tasks = tsk_output("prms", variables = vars_prms() | vars_eta()))
```

obs_additive	<i>Observation with additive error</i>
--------------	--

Description

This building block declares an observation model with an additive residual error model ($y = f + \epsilon_1$).

Usage

```
obs_additive(prediction, name, var_add = 1)
```

Arguments

prediction	A definition of the model prediction
name	A name for the observation (automatically derived if missing)
var_add	Variance of the additive error

Details

Observation models specify the observed variable, how an observation is expected to diverge from the model (i.e, the residual unexplained variability model), and parameter values. The observation model type is selected through the function name. The observed variable as well as the parameters are specified as function arguments.

Specifying predictions:

The actual prediction from the model is the first argument of the function. It can be specified in a number of different ways:

- A name of a variable in the model: `obs_additive("effect")`
- A compartment concentration: `obs_additive(~C["central"])`
- An equation: `obs_additive(~base+slp*time)`

If the definition contains a variable name on the left-hand side (as in `conc~C["central"]`), the variable will appear in the generated model code. This can be useful to make the model code more readable if the prediction is defined as a long equation.

Observation names:

The observation name can be specified via the `name=` argument and is automatically derived if the argument is left empty. Adding an observation model with an already existing name will replace the previous definition.

Error variance:

The variance of the error components are specified via the `var_add=` and `var_prop=` arguments of the function.

Value

A building block of type 'observation'

See Also

Other observation models: [obs_combined\(\)](#), [obs_proportional\(\)](#)

Examples

```
# additive RUV model for observing the variable WT
m <- model() +
  prm_log_normal("wt") +
  obs_additive(~wt)

# EMAX dose-response model with proportional RUV
m2 <- model() +
  input_variable("dose") +
  prm_no_var("emax") +
  prm_no_var("ed50") +
  obs_proportional(effect~emax*dose/(ed50+dose))
```

obs_combined

Observation with combined error

Description

This building block declares an observation model with a combined residual error model ($y = f + f\epsilon_1 + \epsilon_2$).

Usage

```
obs_combined(prediction, name, var_prop = 0.1, var_add = 1)
```

Arguments

prediction	A definition of the model prediction
name	A name for the observation (automatically derived if missing)
var_prop	Variance of the proportional error component
var_add	Variance of the additive error component

Details

Observation models specify the observed variable, how an observation is expected to diverge from the model (i.e. the residual unexplained variability model), and parameter values. The observation model type is selected through the function name. The observed variable as well as the parameters are specified as function arguments.

Specifying predictions:

The actual prediction from the model is the first argument of the function. It can be specified in a number of different ways:

- A name of a variable in the model: `obs_additive("effect")`

- A compartment concentration: `obs_additive(~C["central"])`
- An equation: `obs_additive(~base+slp*time)`

If the definition contains a variable name on the left-hand side (as in `conc~C["central"]`), the variable will appear in the generated model code. This can be useful to make the model code more readable if the prediction is defined as a long equation.

Observation names:

The observation name can be specified via the `name=` argument and is automatically derived if the argument is left empty. Adding an observation model with an already existing name will replace the previous definition.

Error variance:

The variance of the error components are specified via the `var_add=` and `var_prop=` arguments of the function.

Value

A building block of type 'observation'

See Also

Other observation models: [obs_additive\(\)](#), [obs_proportional\(\)](#)

Examples

```
# additive RUV model for observing the variable WT
m <- model() +
  prm_log_normal("wt") +
  obs_additive(~wt)

# EMAX dose-response model with proportional RUV
m2 <- model() +
  input_variable("dose") +
  prm_no_var("emax") +
  prm_no_var("ed50") +
  obs_proportional(effect~emax*dose/(ed50+dose))
```

obs_proportional	<i>Observation with proportional error</i>
------------------	--

Description

This building block declares an observation model with a proportional residual error model ($y = f + f\epsilon_1$).

Usage

```
obs_proportional(prediction, name, var_prop = 0.1)
```

Arguments

prediction	A definition of the model prediction
name	A name for the observation (automatically derived if missing)
var_prop	Variance of the proportional error

Details

Observation models specify the observed variable, how an observation is expected to diverge from the model (i.e, the residual unexplained variability model), and parameter values. The observation model type is selected through the function name. The observed variable as well as the parameters are specified as function arguments.

Specifying predictions:

The actual prediction from the model is the first argument of the function. It can be specified in a number of different ways:

- A name of a variable in the model: `obs_additive("effect")`
- A compartment concentration: `obs_additive(~C["central"])`
- An equation: `obs_additive(~base+slp*time)`

If the definition contains a variable name on the left-hand side (as in `conc~C["central"]`), the variable will appear in the generated model code. This can be useful to make the model code more readable if the prediction is defined as a long equation.

Observation names:

The observation name can be specified via the `name=` argument and is automatically derived if the argument is left empty. Adding an observation model with an already existing name will replace the previous definition.

Error variance:

The variance of the error components are specified via the `var_add=` and `var_prop=` arguments of the function.

Value

A building block of type 'observation'

See Also

Other observation models: [obs_additive\(\)](#), [obs_combined\(\)](#)

Examples

```
# additive RUV model for observing the variable WT
m <- model() +
  prm_log_normal("wt") +
  obs_additive(~wt)

# EMAX dose-response model with proportional RUV
m2 <- model() +
```



```

input_variable("dose") +
prm_no_var("emax") +
prm_no_var("ed50") +
obs_proportional(effect~emax*dose/(ed50+dose))

```

pk_absorption_fo *PK absorption first-order*

Description

This building block declares a first-order absorption component for a pharmacokinetic model.

Usage

```
pk_absorption_fo(prm_mat = prm_log_normal("mat", median = 0.5, var_log = 0.1))
```

Arguments

prm_mat Parameter model for the mean absorption time (MAT)

Details

PK components:

PK components can be added to a [pk_model](#) and exist in three different types: absorption, distribution, and elimination. The absorption component is optional, distribution and elimination are not and need to be added for the PK model to be valid.

A PK model can only have one component of each type and adding a component with an already existing type will replace the previous definition. For example, the distribution component will be a two compartment model in the following snippet:

```

pkm <- pk_model() +
  pk_absorption_fo() +
  pk_distribution_1cmp() +
  pk_distribution_2cmp() +
  pk_elimination_linear() +
  obs_additive(conc~C["central"])
pkm

```

Parameter models:

All PK component functions allow the specification of the parameter model via their arguments. Arguments that refer to a parameter start with the prefix `prm_`. The default parameter model can be deduced from the default arguments in the usage section of the help entry. The parameter name, specified via the `name=` argument of the parameter model building block allows the renaming of the model parameters.

For example, the parameter `prm_vc=` refers to the central volumen of distribution parameter in the one compartment distribution PK component and the default parameter model is a log-normal distribution. The following code block specifies a normal distribution parameter model and names the parameter `v`:

```
pk_distribution_1cmp(  
  prm_vc = prm_normal("v", mean = 50, var = 25)  
)
```

Value

A building block of type 'pk_component'

See Also

[pk_model\(\)](#) for the creation of PK models

Other absorption components: [pk_absorption_fo_lag\(\)](#), [pk_absorption_fo_transit\(\)](#), [pk_absorption_fo_zo\(\)](#), [pk_absorption_zo_lag\(\)](#), [pk_absorption_zo\(\)](#)

pk_absorption_fo_lag *PK absorption first-order, lag-time*

Description

This building block declares a first-order absorption with lag-time component for a pharmacokinetic model.

Usage

```
pk_absorption_fo_lag(  
  prm_mat = prm_log_normal("mat", median = 0.5, var_log = 0.1),  
  prm_mdt = prm_log_normal("mdt", median = 0.5, var_log = 0.1)  
)
```

Arguments

prm_mat	Parameter model for the mean absorption time (MAT)
prm_mdt	Parameter model for the mean delay time (MDT)

Details

PK components:

PK components can be added to a [pk_model](#) and exist in three different types: absorption, distribution, and elimination. The absorption component is optional, distribution and elimination are not and need to be added for the PK model to be valid.

A PK model can only have one component of each type and adding a component with an already existing type will replace the previous definition. For example, the distribution component will be a two compartment model in the following snippet:

```

pkm <- pk_model() +
  pk_absorption_fo() +
  pk_distribution_1cmp() +
  pk_distribution_2cmp() +
  pk_elimination_linear() +
  obs_additive(conc~C["central"])
pkm

```

Parameter models:

All PK component functions allow the specification of the parameter model via their arguments. Arguments that refer to a parameter start with the prefix `prm_`. The default parameter model can be deduced from the default arguments in the usage section of the help entry. The parameter name, specified via the `name=` argument of the parameter model building block allows the renaming of the model parameters.

For example, the parameter `prm_vc=` refers to the central volumen of distribution parameter in the one compartment distribution PK component and the default parameter model is a log-normal distribution. The following code block specifies a normal distribution parameter model and names the parameter `v`:

```

pk_distribution_1cmp(
  prm_vc = prm_normal("v", mean = 50, var = 25)
)

```

Value

A building block of type 'pk_component'

See Also

[pk_model\(\)](#) for the creation of PK models

Other absorption components: [pk_absorption_fo_transit\(\)](#), [pk_absorption_fo_zo\(\)](#), [pk_absorption_fo\(\)](#), [pk_absorption_zo_lag\(\)](#), [pk_absorption_zo\(\)](#)

pk_absorption_fo_transit

PK absorption first-order, transit compartment

Description

This building block declares a first-order absorption with transit compartments component for a pharmacokinetic model.

Usage

```

pk_absorption_fo_transit(
  prm_mat = prm_log_normal("mat", median = 0.5, var_log = 0.1),
  transit_compartments = 1L,
  prm_mdt = prm_log_normal("mdt", median = 0.5, var_log = 0.1)
)

```

Arguments

prm_mat	Parameter model for the mean absorption time (MAT)
transit_compartments	Number of transit compartments
prm_mdt	Parameter model for the mean delay time (MDT)

Details**PK components:**

PK components can be added to a [pk_model](#) and exist in three different types: absorption, distribution, and elimination. The absorption component is optional, distribution and elimination are not and need to be added for the PK model to be valid.

A PK model can only have one component of each type and adding a component with an already existing type will replace the previous definition. For example, the distribution component will be a two compartment model in the following snippet:

```
pkm <- pk_model() +
  pk_absorption_fo() +
  pk_distribution_1cmp() +
  pk_distribution_2cmp() +
  pk_elimination_linear() +
  obs_additive(conc~C["central"])
pkm
```

Parameter models:

All PK component functions allow the specification of the parameter model via their arguments. Arguments that refer to a parameter start with the prefix `prm_`. The default parameter model can be deduced from the default arguments in the usage section of the help entry. The parameter name, specified via the `name=` argument of the parameter model building block allows the renaming of the model parameters.

For example, the parameter `prm_vc=` refers to the central volumen of distribution parameter in the one compartment distribution PK component and the default parameter model is a log-normal distribution. The following code block specifies a normal distribution parameter model and names the parameter `v`:

```
pk_distribution_1cmp(
  prm_vc = prm_normal("v", mean = 50, var = 25)
)
```

Value

A building block of type 'pk_component'

See Also

[pk_model\(\)](#) for the creation of PK models

Other absorption components: [pk_absorption_fo_lag\(\)](#), [pk_absorption_fo_zo\(\)](#), [pk_absorption_fo\(\)](#), [pk_absorption_zo_lag\(\)](#), [pk_absorption_zo\(\)](#)

pk_absorption_fo_zo *PK absorption first-order, zero-order delay*

Description

This building block declares a first-order absorption with zero-order delay component for a pharmacokinetic model.

Usage

```
pk_absorption_fo_zo(
  prm_mat = prm_log_normal("mat", median = 0.5, var_log = 0.1),
  prm_mdt = prm_log_normal("mdt", median = 0.5, var_log = 0.1)
)
```

Arguments

prm_mat	Parameter model for the mean absorption time (MAT)
prm_mdt	Parameter model for the mean delay time (MDT)

Details

PK components:

PK components can be added to a [pk_model](#) and exist in three different types: absorption, distribution, and elimination. The absorption component is optional, distribution and elimination are not and need to be added for the PK model to be valid.

A PK model can only have one component of each type and adding a component with an already existing type will replace the previous definition. For example, the distribution component will be a two compartment model in the following snippet:

```
pkm <- pk_model() +
  pk_absorption_fo() +
  pk_distribution_1cmp() +
  pk_distribution_2cmp() +
  pk_elimination_linear() +
  obs_additive(conc~C["central"])
pkm
```

Parameter models:

All PK component functions allow the specification of the parameter model via their arguments. Arguments that refer to a parameter start with the prefix `prm_`. The default parameter model can be deduced from the default arguments in the usage section of the help entry. The parameter name, specified via the `name=` argument of the parameter model building block allows the renaming of the model parameters.

For example, the parameter `prm_vc=` refers to the central volumen of distribution parameter in the one compartment distribution PK component and the default parameter model is a log-normal distribution. The following code block specifies a normal distribution parameter model and names the parameter `v`:

```
pk_distribution_1cmp(
  prm_vc = prm_normal("v", mean = 50, var = 25)
)
```

Value

A building block of type 'pk_component'

See Also

[pk_model\(\)](#) for the creation of PK models

Other absorption components: [pk_absorption_fo_lag\(\)](#), [pk_absorption_fo_transit\(\)](#), [pk_absorption_fo\(\)](#), [pk_absorption_zo_lag\(\)](#), [pk_absorption_zo\(\)](#)

pk_absorption_zo	<i>PK absorption zero-order</i>
------------------	---------------------------------

Description

This building block declares a zero-order absorption component for a pharmacokinetic model.

Usage

```
pk_absorption_zo(prm_mat = prm_log_normal("mat", median = 0.5, var_log = 0.1))
```

Arguments

prm_mat Parameter model for the mean absorption time (MAT)

Details**PK components:**

PK components can be added to a [pk_model](#) and exist in three different types: absorption, distribution, and elimination. The absorption component is optional, distribution and elimination are not and need to be added for the PK model to be valid.

A PK model can only have one component of each type and adding a component with an already existing type will replace the previous definition. For example, the distribution component will be a two compartment model in the following snippet:

```
pkm <- pk_model() +
  pk_absorption_fo() +
  pk_distribution_1cmp() +
  pk_distribution_2cmp() +
  pk_elimination_linear() +
  obs_additive(conc~C["central"])
pkm
```

Parameter models:

All PK component functions allow the specification of the parameter model via their arguments. Arguments that refer to a parameter start with the prefix `prm_`. The default parameter model can be deduced from the default arguments in the usage section of the help entry. The parameter name, specified via the `name=` argument of the parameter model building block allows the renaming of the model parameters.

For example, the parameter `prm_vc=` refers to the central volumen of distribution parameter in the one compartment distribution PK component and the default parameter model is a log-normal distribution. The following code block specifies a normal distribution parameter model and names the parameter `v`:

```
pk_distribution_1cmp(
  prm_vc = prm_normal("v", mean = 50, var = 25)
)
```

Value

A building block of type 'pk_component'

See Also

[pk_model\(\)](#) for the creation of PK models

Other absorption components: [pk_absorption_fo_lag\(\)](#), [pk_absorption_fo_transit\(\)](#), [pk_absorption_fo_zo\(\)](#), [pk_absorption_fo\(\)](#), [pk_absorption_zo_lag\(\)](#)

pk_absorption_zo_lag *PK absorption zero-order, lag-time*

Description

This building block declares a zero-order absorption with lag-time component for a pharmacokinetic model.

Usage

```
pk_absorption_zo_lag(
  prm_mat = prm_log_normal("mat", median = 0.5, var_log = 0.1),
  prm_mdt = prm_log_normal("mdt", median = 0.5, var_log = 0.1)
)
```

Arguments

<code>prm_mat</code>	Parameter model for the mean absorption time (MAT)
<code>prm_mdt</code>	Parameter model for the mean delay time (MDT)

Details

PK components:

PK components can be added to a [pk_model](#) and exist in three different types: absorption, distribution, and elimination. The absorption component is optional, distribution and elimination are not and need to be added for the PK model to be valid.

A PK model can only have one component of each type and adding a component with an already existing type will replace the previous definition. For example, the distribution component will be a two compartment model in the following snippet:

```
pkm <- pk_model() +
  pk_absorption_fo() +
  pk_distribution_1cmp() +
  pk_distribution_2cmp() +
  pk_elimination_linear() +
  obs_additive(conc~C["central"])
pkm
```

Parameter models:

All PK component functions allow the specification of the parameter model via their arguments. Arguments that refer to a parameter start with the prefix `prm_`. The default parameter model can be deduced from the default arguments in the usage section of the help entry. The parameter name, specified via the `name=` argument of the parameter model building block allows the renaming of the model parameters.

For example, the parameter `prm_vc=` refers to the central volumen of distribution parameter in the one compartment distribution PK component and the default parameter model is a log-normal distribution. The following code block specifies a normal distribution parameter model and names the parameter `v`:

```
pk_distribution_1cmp(
  prm_vc = prm_normal("v", mean = 50, var = 25)
)
```

Value

A building block of type 'pk_component'

See Also

[pk_model\(\)](#) for the creation of PK models

Other absorption components: [pk_absorption_fo_lag\(\)](#), [pk_absorption_fo_transit\(\)](#), [pk_absorption_fo_zo\(\)](#), [pk_absorption_fo\(\)](#), [pk_absorption_zo\(\)](#)

pk_distribution_1cmp *PK distribution 1 compartment*

Description

This building block declares a one compartment distribution component for a pharmacokinetic model.

Usage

```
pk_distribution_1cmp(  
  prm_vc = prm_log_normal("vc", median = 100, var_log = 0.1)  
)
```

Arguments

prm_vc Parameter model for the central volume of distribution

Details

PK components:

PK components can be added to a [pk_model](#) and exist in three different types: absorption, distribution, and elimination. The absorption component is optional, distribution and elimination are not and need to be added for the PK model to be valid.

A PK model can only have one component of each type and adding a component with an already existing type will replace the previous definition. For example, the distribution component will be a two compartment model in the following snippet:

```
pkm <- pk_model() +  
  pk_absorption_fo() +  
  pk_distribution_1cmp() +  
  pk_distribution_2cmp() +  
  pk_elimination_linear() +  
  obs_additive(conc~C["central"])  
pkm
```

Parameter models:

All PK component functions allow the specification of the parameter model via their arguments. Arguments that refer to a parameter start with the prefix `prm_`. The default parameter model can be deduced from the default arguments in the usage section of the help entry. The parameter name, specified via the `name=` argument of the parameter model building block allows the renaming of the model parameters.

For example, the parameter `prm_vc=` refers to the central volumen of distribution parameter in the one compartment distribution PK component and the default parameter model is a log-normal distribution. The follwing code block specifies a normal distribution parameter model and names the parameter `v`:

```
pk_distribution_1cmp(
  prm_vc = prm_normal("v", mean = 50, var = 25)
)
```

Value

A building block of type 'pk_component'

See Also

[pk_model\(\)](#) for the creation of PK models

Other distribution components: [pk_distribution_2cmp\(\)](#), [pk_distribution_3cmp\(\)](#)

pk_distribution_2cmp *PK distribution 2 compartments*

Description

This building block declares a two compartment distribution component for a pharmacokinetic model.

Usage

```
pk_distribution_2cmp(
  prm_vc = prm_log_normal("vc", median = 100, var_log = 0.1),
  prm_vp = prm_log_normal("vp", median = 5, var_log = 0.1),
  prm_q = prm_log_normal("q", median = 50, var_log = 0.1)
)
```

Arguments

prm_vc	Parameter model for the central volume of distribution
prm_vp	Parameter model for the peripheral volume of distribution
prm_q	Parameter model for the inter-compartmental clearance

Details**PK components:**

PK components can be added to a [pk_model](#) and exist in three different types: absorption, distribution, and elimination. The absorption component is optional, distribution and elimination are not and need to be added for the PK model to be valid.

A PK model can only have one component of each type and adding a component with an already existing type will replace the previous definition. For example, the distribution component will be a two compartment model in the following snippet:

```

pkm <- pk_model() +
  pk_absorption_fo() +
  pk_distribution_1cmp() +
  pk_distribution_2cmp() +
  pk_elimination_linear() +
  obs_additive(conc~C["central"])
pkm

```

Parameter models:

All PK component functions allow the specification of the parameter model via their arguments. Arguments that refer to a parameter start with the prefix `prm_`. The default parameter model can be deduced from the default arguments in the usage section of the help entry. The parameter name, specified via the `name=` argument of the parameter model building block allows the renaming of the model parameters.

For example, the parameter `prm_vc=` refers to the central volumen of distribution parameter in the one compartment distribution PK component and the default parameter model is a log-normal distribution. The following code block specifies a normal distribution parameter model and names the parameter `v`:

```

pk_distribution_1cmp(
  prm_vc = prm_normal("v", mean = 50, var = 25)
)

```

Value

A building block of type 'pk_component'

See Also

[pk_model\(\)](#) for the creation of PK models

Other distribution components: [pk_distribution_1cmp\(\)](#), [pk_distribution_3cmp\(\)](#)

pk_distribution_3cmp *PK distribution 3 compartments*

Description

This building block declares a three compartment distribution component for a pharmacokinetic model.

Usage

```

pk_distribution_3cmp(
  prm_vc = prm_log_normal("vc", median = 100, var_log = 0.1),
  prm_vp1 = prm_log_normal("vp1", median = 5, var_log = 0.1),
  prm_vp2 = prm_log_normal("vp2", median = 5, var_log = 0.1),
  prm_q1 = prm_log_normal("q1", median = 25, var_log = 0.1),
  prm_q2 = prm_log_normal("q2", median = 25, var_log = 0.1)
)

```

Arguments

prm_vc	Parameter model for the central volume of distribution
prm_vp1	Parameter model for the volume of the first peripheral compartment
prm_vp2	Parameter model for the volume of the second peripheral compartment
prm_q1	Parameter model for the inter-compartmental clearance between central and first peripheral compartment
prm_q2	Parameter model for the inter-compartmental clearance between central and second peripheral compartment

Details**PK components:**

PK components can be added to a [pk_model](#) and exist in three different types: absorption, distribution, and elimination. The absorption component is optional, distribution and elimination are not and need to be added for the PK model to be valid.

A PK model can only have one component of each type and adding a component with an already existing type will replace the previous definition. For example, the distribution component will be a two compartment model in the following snippet:

```
pkm <- pk_model() +
  pk_absorption_fo() +
  pk_distribution_1cmp() +
  pk_distribution_2cmp() +
  pk_elimination_linear() +
  obs_additive(conc~C["central"])
pkm
```

Parameter models:

All PK component functions allow the specification of the parameter model via their arguments. Arguments that refer to a parameter start with the prefix `prm_`. The default parameter model can be deduced from the default arguments in the usage section of the help entry. The parameter name, specified via the `name=` argument of the parameter model building block allows the renaming of the model parameters.

For example, the parameter `prm_vc=` refers to the central volumen of distribution parameter in the one compartment distribution PK component and the default parameter model is a log-normal distribution. The follwing code block specifies a normal distribution parameter model and names the parameter `v`:

```
pk_distribution_1cmp(
  prm_vc = prm_normal("v", mean = 50, var = 25)
)
```

Value

A building block of type 'pk_component'

See Also

[pk_model\(\)](#) for the creation of PK models

Other distribution components: [pk_distribution_1cmp\(\)](#), [pk_distribution_2cmp\(\)](#)

pk_elimination_linear *PK elimination linear*

Description

This building block declares a linear elimination component for a pharmacokinetic model.

Usage

```
pk_elimination_linear(  
  prm_cl = prm_log_normal("cl", median = 50, var_log = 0.1)  
)
```

Arguments

prm_cl Parameter model for the clearance

Details**PK components:**

PK components can be added to a [pk_model](#) and exist in three different types: absorption, distribution, and elimination. The absorption component is optional, distribution and elimination are not and need to be added for the PK model to be valid.

A PK model can only have one component of each type and adding a component with an already existing type will replace the previous definition. For example, the distribution component will be a two compartment model in the following snippet:

```
pkm <- pk_model() +  
  pk_absorption_fo() +  
  pk_distribution_1cmp() +  
  pk_distribution_2cmp() +  
  pk_elimination_linear() +  
  obs_additive(conc~C["central"])  
pkm
```

Parameter models:

All PK component functions allow the specification of the parameter model via their arguments. Arguments that refer to a parameter start with the prefix `prm_`. The default parameter model can be deduced from the default arguments in the usage section of the help entry. The parameter name, specified via the `name=` argument of the parameter model building block allows the renaming of the model parameters.

For example, the parameter `prm_vc=` refers to the central volumen of distribution parameter in the one compartment distribution PK component and the default parameter model is a log-normal

distribution. The following code block specifies a normal distribution parameter model and names the parameter `v`:

```
pk_distribution_1cmp(
  prm_vc = prm_normal("v", mean = 50, var = 25)
)
```

Value

A building block of type 'pk_component'

See Also

[pk_model\(\)](#) for the creation of PK models

Other elimination components: [pk_elimination_linear_nl\(\)](#), [pk_elimination_nl\(\)](#)

pk_elimination_linear_nl

PK elimination linear & nonlinear

Description

This building block declares a mixed linear and nonlinear elimination component for a pharmacokinetic model.

Usage

```
pk_elimination_linear_nl(
  prm_cllin = prm_log_normal("cllin", median = 50, var_log = 0.1),
  prm_vmax = prm_log_normal("vmax", median = 10, var_log = 0.1),
  prm_km = prm_log_normal("km", median = 0.5, var_log = 0.1)
)
```

Arguments

prm_cllin	Parameter model for the linear clearance
prm_vmax	Parameter model for Vmax (the maximal elimination rate)
prm_km	Parameter model for KM (the half-maximal concentration)

Details

PK components:

PK components can be added to a [pk_model](#) and exist in three different types: absorption, distribution, and elimination. The absorption component is optional, distribution and elimination are not and need to be added for the PK model to be valid.

A PK model can only have one component of each type and adding a component with an already existing type will replace the previous definition. For example, the distribution component will be a two compartment model in the following snippet:

```

pkm <- pk_model() +
  pk_absorption_fo() +
  pk_distribution_1cmp() +
  pk_distribution_2cmp() +
  pk_elimination_linear() +
  obs_additive(conc~C["central"])
pkm

```

Parameter models:

All PK component functions allow the specification of the parameter model via their arguments. Arguments that refer to a parameter start with the prefix `prm_`. The default parameter model can be deduced from the default arguments in the usage section of the help entry. The parameter name, specified via the `name=` argument of the parameter model building block allows the renaming of the model parameters.

For example, the parameter `prm_vc=` refers to the central volumen of distribution parameter in the one compartment distribution PK component and the default parameter model is a log-normal distribution. The following code block specifies a normal distribution parameter model and names the parameter `v`:

```

pk_distribution_1cmp(
  prm_vc = prm_normal("v", mean = 50, var = 25)
)

```

Value

A building block of type 'pk_component'

See Also

[pk_model\(\)](#) for the creation of PK models

Other elimination components: [pk_elimination_linear\(\)](#), [pk_elimination_nl\(\)](#)

pk_elimination_nl *PK elimination nonlinear*

Description

This building block declares a nonlinear elimination component for a pharmacokinetic model.

Usage

```

pk_elimination_nl(
  prm_clmm = prm_log_normal("clmm", median = 25, var_log = 0.1),
  prm_km = prm_log_normal("km", median = 0.5, var_log = 0.1),
  prm_vmax = NULL
)

```

Arguments

prm_clmm	Parameter model for the clearance
prm_km	Parameter model for KM (the half-maximal concentration)
prm_vmax	Parameter model for Vmax (the maximal elimination rate)

Details**PK components:**

PK components can be added to a [pk_model](#) and exist in three different types: absorption, distribution, and elimination. The absorption component is optional, distribution and elimination are not and need to be added for the PK model to be valid.

A PK model can only have one component of each type and adding a component with an already existing type will replace the previous definition. For example, the distribution component will be a two compartment model in the following snippet:

```
pkm <- pk_model() +
  pk_absorption_fo() +
  pk_distribution_1cmp() +
  pk_distribution_2cmp() +
  pk_elimination_linear() +
  obs_additive(conc~C["central"])
pkm
```

Parameter models:

All PK component functions allow the specification of the parameter model via their arguments. Arguments that refer to a parameter start with the prefix `prm_`. The default parameter model can be deduced from the default arguments in the usage section of the help entry. The parameter name, specified via the `name=` argument of the parameter model building block allows the renaming of the model parameters.

For example, the parameter `prm_vc=` refers to the central volumen of distribution parameter in the one compartment distribution PK component and the default parameter model is a log-normal distribution. The following code block specifies a normal distribution parameter model and names the parameter `v`:

```
pk_distribution_1cmp(
  prm_vc = prm_normal("v", mean = 50, var = 25)
)
```

Value

A building block of type 'pk_component'

See Also

[pk_model\(\)](#) for the creation of PK models

Other elimination components: [pk_elimination_linear_nl\(\)](#), [pk_elimination_linear\(\)](#)

pk_model	<i>Create a PK model</i>
----------	--------------------------

Description

This function creates the basis for a pharmacokinetic model.

Usage

```
pk_model()
```

Details

The function creates the fundament for a pharmacokinetic model to which different building blocks can be added. The following building blocks are relevant for this model type:

- Parameters: [prm_log_normal](#), [prm_logit_normal](#), [prm_no_var](#), [prm_normal](#)
- Observations: [obs_additive](#), [obs_combined](#), [obs_proportional](#)
- Algebraic relationships: [algebraic](#)
- PK components: [pk_absorption_fo](#), [pk_absorption_fo_lag](#), [pk_absorption_fo_transit](#), [pk_absorption_fo_zo](#), [pk_absorption_zo](#), [pk_absorption_zo_lag](#), [pk_distribution_1cmp](#), [pk_distribution_2cmp](#), [pk_distribution_3cmp](#), [pk_elimination_linear](#), [pk_elimination_linear_nl](#), [pk_elimination_nl](#), [pk_model](#)
- Input variables: [input_variable](#), [dataset](#)

Value

A `pk_model`

<code>prm_logit_normal</code>	<i>Parameter with logit-normal distribution</i>
-------------------------------	---

Description

This building block declares a parameter model for a parameter that follows the normal distribution on the logit-scale.

Usage

```
prm_logit_normal(name, mean_logit = 0, var_logit = 1)
```

Arguments

<code>name</code>	Parameter name
<code>mean_logit</code>	Mean on the logit scale
<code>var_logit</code>	Variance on the logit scale

Details

Parameter models specify type, name, and values for a parameter. The parameter model type is selected through the function name. The parameter name and values are provided as function arguments.

Parameter names:

Every parameter must have a valid name. A parameter name can contain letters, numbers as well as the underscore character. The name needs to start with a letter.

Adding a parameter with an already existing name will replace the definition of the parameter. For example, the parameter “base” will have a log-normal distribution in the following snippet:

```
m <- model() +
  prn_normal("base") +
  prn_log_normal("base")
```

Parameter values:

The parameter values that a parameter model expects vary by type. For example, `prn_normal()` requires the mean and the variance, whereas for `prn_log_normal()` median and variance on the log scale need to be provided. The argument name should indicate what parameter value is expected.

MU-referencing:

`assemblerr` can include mu-referencing statements for parameter distributions that support it. The functionality can be activated by setting the option `prn.use_mu_referencing` to `TRUE` as shown in the following snippet:

```
m <- model() +
  prn_normal("base") +
  prn_log_normal("slp") +
  obs_additive(response~base+slp*time)

render(
  model = m,
  options = assemblerr_options(prn.use_mu_referencing = TRUE)
)
```

Value

A building block of type ‘parameter’

See Also

Other parameter models: [prn_log_normal\(\)](#), [prn_no_var\(\)](#), [prn_normal\(\)](#)

Examples

```
# EMAX dose-response model with emax (log-normal) and ed50 (no variability) parameters
m2 <- model() +
  input_variable("dose") +
  prn_log_normal("emax", 10, 0.3) +
```

```

prn_no_var("ed50", 5) +
obs_proportional(effect~emax*dose/(ed50+dose))

# a log-normal parameter that is directly observed
m <- model() +
  prn_log_normal("wt") +
  obs_additive(~wt)

```

prn_log_normal	<i>Parameter with log-normal distribution</i>
----------------	---

Description

This building block declares a parameter model for a parameter that follows the normal distribution on the log scale.

Usage

```
prn_log_normal(name, median = 1, var_log = 0.1)
```

Arguments

name	Parameter name
median	Median (on the normal scale)
var_log	Variance on the log scale

Details

Parameter models specify type, name, and values for a parameter. The parameter model type is selected through the function name. The parameter name and values are provided as function arguments.

Parameter names:

Every parameter must have a valid name. A parameter name can contain letters, numbers as well as the underscore character. The name needs to start with a letter.

Adding a parameter with an already existing name will replace the definition of the parameter. For example, the parameter “base” will have a log-normal distribution in the following snippet:

```

m <- model() +
  prn_normal("base") +
  prn_log_normal("base")

```

Parameter values:

The parameter values that a parameter model expects vary by type. For example, `prn_normal()` requires the mean and the variance, whereas for `prn_log_normal()` median and variance on the log scale need to be provided. The argument name should indicate what parameter value is expected.

MU-referencing:

assemblerr can include mu-referencing statements for parameter distributions that support it. The functionality can be activated by setting the option `prm.use_mu_referencing` to `TRUE` as shown in the following snippet:

```
m <- model() +
  prm_normal("base") +
  prm_log_normal("slp") +
  obs_additive(response~base+slp*time)

render(
  model = m,
  options = assemblerr_options(prm.use_mu_referencing = TRUE)
)
```

Value

A building block of type 'parameter'

See Also

Other parameter models: [prm_logit_normal\(\)](#), [prm_no_var\(\)](#), [prm_normal\(\)](#)

Examples

```
# EMAX dose-response model with emax (log-normal) and ed50 (no variability) parameters
m2 <- model() +
  input_variable("dose") +
  prm_log_normal("emax", 10, 0.3) +
  prm_no_var("ed50", 5) +
  obs_proportional(effect~emax*dose/(ed50+dose))

# a log-normal parameter that is directly observed
m <- model() +
  prm_log_normal("wt") +
  obs_additive(~wt)
```

```
prm_normal
```

```
Parameter with normal distribution
```

Description

This building block declares a parameter model for a parameter that follows the normal distribution.

Usage

```
prm_normal(name, mean = 1, var = 0.1)
```

Arguments

name	Parameter name
mean	Mean
var	Variance

Details

Parameter models specify type, name, and values for a parameter. The parameter model type is selected through the function name. The parameter name and values are provided as function arguments.

Parameter names:

Every parameter must have a valid name. A parameter name can contain letters, numbers as well as the underscore character. The name needs to start with a letter.

Adding a parameter with an already existing name will replace the definition of the parameter. For example, the parameter “base” will have a log-normal distribution in the following snippet:

```
m <- model() +
  prn_normal("base") +
  prn_log_normal("base")
```

Parameter values:

The parameter values that a parameter model expects vary by type. For example, `prn_normal()` requires the mean and the variance, whereas for `prn_log_normal()` median and variance on the log scale need to be provided. The argument name should indicate what parameter value is expected.

MU-referencing:

`assemblerr` can include mu-referencing statements for parameter distributions that support it. The functionality can be activated by setting the option `prn.use_mu_referencing` to `TRUE` as shown in the following snippet:

```
m <- model() +
  prn_normal("base") +
  prn_log_normal("slp") +
  obs_additive(response~base+slp*time)

render(
  model = m,
  options = assemblerr_options(prn.use_mu_referencing = TRUE)
)
```

Value

A building block of type ‘parameter’

See Also

Other parameter models: [prn_log_normal\(\)](#), [prn_logit_normal\(\)](#), [prn_no_var\(\)](#)

Examples

```
# EMAX dose-response model with emax (log-normal) and ed50 (no variability) parameters
m2 <- model() +
  input_variable("dose") +
  prn_log_normal("emax", 10, 0.3) +
  prn_no_var("ed50", 5) +
  obs_proportional(effect~emax*dose/(ed50+dose))

# a log-normal parameter that is directly observed
m <- model() +
  prn_log_normal("wt") +
  obs_additive(~wt)
```

prn_no_var

Parameter without variability

Description

This building block declares a parameter model for a parameter that does not vary between subjects.

Usage

```
prn_no_var(name, value = 1)
```

Arguments

name	Parameter name
value	Parameter value

Details

Parameter models specify type, name, and values for a parameter. The parameter model type is selected through the function name. The parameter name and values are provided as function arguments.

Parameter names:

Every parameter must have a valid name. A parameter name can contain letters, numbers as well as the underscore character. The name needs to start with a letter.

Adding a parameter with an already existing name will replace the definition of the parameter. For example, the parameter “base” will have a log-normal distribution in the following snippet:

```
m <- model() +
  prn_normal("base") +
  prn_log_normal("base")
```

Parameter values:

The parameter values that a parameter model expects vary by type. For example, `prn_normal()` requires the mean and the variance, whereas for `prn_log_normal()` median and variance on the log scale need to be provided. The argument name should indicate what parameter value is expected.

MU-referencing:

`assemblerr` can include mu-referencing statements for parameter distributions that support it. The functionality can be activated by setting the option `prn.use_mu_referencing` to `TRUE` as shown in the following snippet:

```
m <- model() +
  prn_normal("base") +
  prn_log_normal("slp") +
  obs_additive(response~base+slp*time)

render(
  model = m,
  options = assemblerr_options(prn.use_mu_referencing = TRUE)
)
```

Value

A building block of type 'parameter'

See Also

Other parameter models: [prn_log_normal\(\)](#), [prn_logit_normal\(\)](#), [prn_normal\(\)](#)

Examples

```
# EMAX dose-response model with emax (log-normal) and ed50 (no variability) parameters
m2 <- model() +
  input_variable("dose") +
  prn_log_normal("emax", 10, 0.3) +
  prn_no_var("ed50", 5) +
  obs_proportional(effect~emax*dose/(ed50+dose))

# a log-normal parameter that is directly observed
m <- model() +
  prn_log_normal("wt") +
  obs_additive(~wt)
```

render	<i>Generate model code</i>
--------	----------------------------

Description

This function generates the code for a model object, prints it to the console or writes it to a file.

Usage

```
render(
  model,
  filename = NULL,
  target_tool = "nonmem",
  tasks = tsk_estimation(),
  options = assemblerr_options()
)
```

Arguments

model	A model object
filename	Name of the model file to create or NULL
target_tool	Name of the target tool (currently only 'nonmem')
tasks	A task specification
options	List of options for model generation

Details

The generated code will be written to the file specified by filename= or printed to the console if the filename is set to NULL. Only 'nonmem' is currently supported as a target_tool= option. The tasks= argument allows the specification of model tasks and the options= argument customizes the generated code.

Task specification:

Tasks are building blocks that allow to specify what a model should "do". Like other model building blocks, they can be combined using the + operator. For example, the following adds an estimation task and an xpose4 output task to the generated code:

```
render(m, tasks = tsk_estimation() +
      tsk_output_xpose4())
```

The default argument (tasks=tsk_estimation()) adds an FOCE estimation task to the code.

Rendering options:

The options= argument allows to modify the rendering process and, hence, the generated code. Options are provided as a list and the assemblerr_options() function helps to generate list with the proper formatting.

The following code block renders the model m with automatic mu-referencing for the model parameters

```
render(m, options = assemblerr_options(prm.use_mu_referencing = TRUE))
```


Value

The model code as a character vector

Examples

```
m <- model() +
  input_variable("dose") +
  prm_log_normal("emax") +
  prm_log_normal("ed50") +
  obs_additive(eff~emax*dose/(ed50+dose))
# render to console
render(m)

# render to file
## Not run:
setwd(tempdir())
render(m, "run1.mod")

## End(Not run)

# render to console with estimation & output task
render(m, tasks = tsk_estimation() + tsk_output_xpose4())
```

tsk_estimation

Task estimation

Description

This function defines an estimation task allowing to specify the estimation algorithm, estimation options, and whether standard errors should be obtained.

Usage

```
tsk_estimation(algorithm = "foce", se = FALSE, target_options = list())
```

Arguments

algorithm	The estimation algorithm to use for the task ("foce", "foce-inter", "foce-no-inter", "fo", "imp", "saem")
se	Whether to calculate parameter uncertainties
target_options	List of additional options that should be passed to NONMEM

Details

Tasks:

Tasks are building blocks that allow to specify what a model should “do”. Like other model building blocks, they can be combined using the + operator. However, they should not be added to a model but rather provided via the `tasks=` argument to the render function, e.g.,

```
render(m, tasks = tsk_estimation() +
       tsk_output_xpose4())
```

Estimation tasks:

Estimation tasks provide details on the parameter estimation process, in terms of estimation algorithm, estimation options and whether standard errors should be obtained.

Algorithm:

The algorithm argument allows to select the estimation algorithm among the following options:

foce	First-order conditional estimation with interaction detection
foce-inter	First-order conditional estimation with interaction
foce-no-inter	First-order conditional estimation without interaction
fo	First-order estimation
imp	Importance sampling
saem	Stochastic approximation expectation maximization

The default algorithm “foce” detects whether the observation model includes an epsilon-eta interaction and includes the INTERACTION option accordingly. The `foce-inter` option forces the use of the INTERACTION argument independent of the residual error model, `foce-no-inter` enforces no interaction.

Each algorithm includes a set of default options that the package authors consider sensible defaults (for example `MAXEVAL=999999` for FOCE). These defaults can be overwritten using the `target_options=` argument which is described below.

Standard errors:

The `se=` argument allows to request the calculation of parameter standard errors. When standard errors are requested (`se=TRUE`) it will result in the inclusion of the \$COVARIANCE record in the generated control stream.

Target options:

The `target_options=` argument provides a mechanism to specify additional estimation options for the selected algorithm. The options should be provided as a list, e.g.,

```
tsk_estimation(algorithm = "foce", target_options = list(mcceta=100))
```

The provided options are passed verbatim to the target tool and not checked by `assemblerr` for correctness.

The `target_options=` argument

Multiple estimation tasks:

A sequence of estimation tasks can be specified in `assemblerr` by combining multiple estimations, for example

```
render(m, tasks = tsk_estimation("foce") + tsk_estimation("imp"))
```

will create model code that contains an FOCE as well as an importance sampling estimation step.

Value

A building block of type 'estimation_task'

See Also

Other tasks: [tsk_output\(\)](#)

Examples

```
m <- model() +
  input_variable("dose") +
  prn_log_normal("emax", median = 10, var_log = 0.09) +
  prn_log_normal("ed50", median = 50, var_log = 0.09) +
  algebraic(effect~emax*dose/(ed50 + dose)) +
  obs_proportional(~effect, var_prop = 1)

# add estimation task using importance sampling, covariance step
# and user-defined ISAMPLE option
render(
  model = m,
  tasks = tsk_estimation(
    algorithm = "imp",
    se = TRUE,
    target_options = list(isample=1000)
  )
)
```

tsk_output

Task output

Description

These functions define output tasks that include the selected variables in the output of the generated model.

Usage

```
tsk_output(filename = "sdtab", variables)
```

```
tsk_output_xpose4()
```

Arguments

filename	The filename for the output file
variables	The model variables that be included in the output

Details

Tasks:

Tasks are building blocks that allow to specify what a model should “do”. Like other model building blocks, they can be combined using the + operator. However, they should not be added to a model but rather provided via the `tasks=` argument to the render function, e.g.,

```
render(m, tasks = tsk_estimation() +
       tsk_output_xpose4())
```

Output tasks:

For NONMEM, an output task defines the \$TABLE records by specifying the `filename=` as well as the `variables=` to include.

The variables can be specified by providing a character vector of variable names (e.g., `variables = c('c1', 'v')`) or by using a set of variable selection helpers (e.g., `variables = vars_prms()`). The latter is shorter if many variables are to be selected and allows the specification of tasks independent from the model. The details of the variable selection language can be found on the help pages for [model-variable-selection](#).

xpose4 output task:

The `tsk_output_xpose4()` function includes \$TABLE records that follow the output conventions of the model diagnostic package `xpose4`. It is a shortcut for the following two output tasks:

```
xpose4_output <- tsk_output("sdtab", variables = any_of(c("id", "time"))) | vars_nm_std() +
  tsk_output("patab", variables = vars_prms() | vars_eta())
```

Value

A building block of type 'output_task'

See Also

Other tasks: [tsk_estimation\(\)](#)

Examples

```
m <- model() +
  input_variable("dose") +
  prm_log_normal("emax", median = 10, var_log = 0.09) +
  prm_log_normal("ed50", median = 50, var_log = 0.09) +
  algebraic(effect~emax*dose/(ed50 + dose)) +
  obs_proportional(~effect, var_prop = 1)
# output model parameters to file 'prms'
render(m, tasks = tsk_output("prms", variables = vars_prms()))
# output variables required by xpose4
render(m, tasks = tsk_output_xpose4())
```

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