

Package ‘dyngen’

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Type Package

Title A Multi-Modal Simulator for Spearheading Single-Cell Omics Analyses

Version 1.0.5

Description A novel, multi-modal simulation engine for studying dynamic cellular processes at single-cell resolution. 'dyngen' is more flexible than current single-cell simulation engines. It allows better method development and benchmarking, thereby stimulating development and testing of novel computational methods. Cannoodt et al. (2021) <[doi:10.1038/s41467-021-24152-2](https://doi.org/10.1038/s41467-021-24152-2)>.

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URL <https://dyngen.dynverse.org>, <https://github.com/dynverse/dyngen>

BugReports <https://github.com/dynverse/dyngen/issues>

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R topics documented:

as_dyno	3
backbone	4
bblego	6
combine_models	8
dyngen	9
example_model	12
generate_cells	12
generate_dataset	14
generate_experiment	15
generate_feature_network	17
generate_gold_standard	18
generate_kinetics	19
generate_tf_network	21
get_timings	22
initialise_model	23
kinetics_noise_none	24
list_backbones	25
plot_backbone_modulenet	27
plot_backbone_statenet	28
plot_experiment_dimred	28
plot_feature_network	29
plot_gold_expression	30
plot_gold_mappings	30
plot_gold_simulations	31
plot_simulations	32
plot_simulation_expression	33
plot_summary	34
realcounts	34
realnets	35
rnorm_bounded	35
runif_subrange	36
simtime_from_backbone	36

Index

38

`as_dyno`*Convert simulation output to different formats.*

Description

For use with other packages compatible with `dyno`, `anndata`, `SingleCellExperiment`, or `Seurat`.

Usage

```
as_dyno(  
  model,  
  store_dimred = !is.null(model$simulations$dimred),  
  store_cellwise_grn = !is.null(model$experiment$cellwise_grn),  
  store_rna_velocity = !is.null(model$experiment$rna_velocity)  
)  
  
as_anndata(  
  model,  
  store_dimred = !is.null(model$simulations$dimred),  
  store_cellwise_grn = !is.null(model$experiment$cellwise_grn),  
  store_rna_velocity = !is.null(model$experiment$rna_velocity)  
)  
  
as_sce(  
  model,  
  store_dimred = !is.null(model$simulations$dimred),  
  store_cellwise_grn = !is.null(model$experiment$cellwise_grn),  
  store_rna_velocity = !is.null(model$experiment$rna_velocity)  
)  
  
as_seurat(  
  model,  
  store_dimred = !is.null(model$simulations$dimred),  
  store_cellwise_grn = !is.null(model$experiment$cellwise_grn),  
  store_rna_velocity = !is.null(model$experiment$rna_velocity)  
)  
  
as_list(  
  model,  
  store_dimred = !is.null(model$simulations$dimred),  
  store_cellwise_grn = !is.null(model$experiment$cellwise_grn),  
  store_rna_velocity = !is.null(model$experiment$rna_velocity)  
)  
  
wrap_dataset(  
  model,  
  format = c("list", "dyno", "sce", "seurat", "anndata", "none"),
```

```

store_dimred = !is.null(model$simulations$dimred),
store_cellwise_grn = !is.null(model$experiment$cellwise_grn),
store_rna_velocity = !is.null(model$experiment$rna_velocity)
)

```

Arguments

model A dyngen output model for which the experiment has been emulated with `generate_experiment()`.

store_dimred Whether or not to store the dimensionality reduction constructed on the true counts.

store_cellwise_grn Whether or not to also store cellwise GRN information.

store_rna_velocity Whether or not to store the log propensity ratios.

format Which output format to use, must be one of 'dyno' (requires dynwrap), 'sce' (requires SingleCellExperiment), 'seurat' (requires Seurat), 'anndata' (requires anndata), 'list' or 'none'.

Value

A dataset object.

Examples

```

data("example_model")
dataset <- wrap_dataset(example_model, format = "list")

dataset <- wrap_dataset(example_model, format = "dyno")
dataset <- wrap_dataset(example_model, format = "sce")
dataset <- wrap_dataset(example_model, format = "seurat")
dataset <- wrap_dataset(example_model, format = "anndata")
dataset <- wrap_dataset(example_model, format = "none")

```

backbone

Backbone of the simulation model

Description

A module is a group of genes which, to some extent, shows the same expression behaviour. Several modules are connected together such that one or more genes from one module will regulate the expression of another module. By creating chains of modules, a dynamic behaviour in gene regulation can be created.

Usage

```
backbone(module_info, module_network, expression_patterns)
```

Arguments

- `module_info` A tibble containing meta information on the modules themselves.
- `module_id` (character): the name of the module
 - `basal` (numeric): basal expression level of genes in this module, must be between [0, 1]
 - `burn` (logical): whether or not outgoing edges of this module will be active during the burn in phase
 - `independence` (numeric): the independence factor between regulators of this module, must be between [0, 1]
- `module_network` A tibble describing which modules regulate which other modules.
- `from` (character): the regulating module
 - `to` (character): the target module
 - `effect` (integer): 1L if the regulating module upregulates the target module, -1L if it downregulates
 - `strength` (numeric): the strength of the interaction
 - `hill` (numeric): hill coefficient, larger than 1 for positive cooperativity, between 0 and 1 for negative cooperativity
- `expression_patterns`
- A tibble describing the expected expression pattern changes when a cell is simulated by dyngen. Each row represents one transition between two cell states.
- `from` (character): name of a cell state
 - `to` (character): name of a cell state
 - `module_progression` (character): differences in module expression between the two states. Example: "+4, -1 | +9 | -4" means the expression of module 4 will go up at the same time as module 1 goes down; afterwards module 9 expression will go up, and afterwards module 4 expression will go down again.
 - `start` (logical): Whether or not this from cell state is the start of the trajectory
 - `burn` (logical): Whether these cell states are part of the burn in phase. Cells will not get sampled from these cell states.
 - `time` (numeric): The duration of an transition.

Value

A dyngen backbone.

See Also

[dyngen](#) on how to run a dyngen simulation

Examples

```
library(tibble)
backbone <- backbone(
  module_info = tribble(
    ~module_id, ~basal, ~burn, ~independence,
```

```

    "M1",      1,      TRUE,  1,
    "M2",      0,      FALSE, 1,
    "M3",      0,      FALSE, 1
  ),
  module_network = tribble(
    ~from, ~to, ~effect, ~strength, ~hill,
    "M1",  "M2", 1L,      1,        2,
    "M2",  "M3", 1L,      1,        2
  ),
  expression_patterns = tribble(
    ~from, ~to, ~module_progression, ~start, ~burn, ~time,
    "s0",  "s1", "+M1",      TRUE,  TRUE,  30,
    "s1",  "s2", "+M2,+M3", FALSE, FALSE,  80
  )
)

```

 bblego

Design your own custom backbone easily

Description

You can use the bblego functions in order to create custom backbones using various components. Please note that the bblego functions currently only allow you to create tree-like backbones.

Usage

```
bblego(..., .list = NULL)
```

```

bblego_linear(
  from,
  to,
  type = sample(c("simple", "doublerep1", "doublerep2"), 1),
  num_modules = sample(4:6, 1),
  burn = FALSE
)

```

```

bblego_branching(
  from,
  to,
  type = "simple",
  num_steps = 3,
  num_modules = 2 + length(to) * (3 + num_steps),
  burn = FALSE
)

```

```

bblego_start(
  to,
  type = sample(c("simple", "doublerep1", "doublerep2"), 1),

```

```

    num_modules = sample(4:6, 1)
  )

bblego_end(
  from,
  type = sample(c("simple", "doublerep1", "doublerep2"), 1),
  num_modules = sample(4:6, 1)
)

```

Arguments

<code>..., .list</code>	bblego components, either as separate args or as a list.
<code>from</code>	The begin state of this component.
<code>to</code>	The end state of this component.
<code>type</code>	Some components have alternative module regulatory networks. bblego_start(), bblego_linear(), bblego_end(): <ul style="list-style-type: none"> "simple": a sequence of modules in which every module upregulates the next module. "doublerep1": a sequence of modules in which every module downregulates the next module, and each module has positive basal expression. "doublerep2": a sequence of modules in which every module upregulates the next module, but downregulates the one after that. "flipflop": a sequence of modules in which every module upregulates the next module. In addition, the last module upregulates itself and strongly downregulates the first module. bblego_branching(): <ul style="list-style-type: none"> "simple": a set of n modules (with $n = \text{length}(\text{to})$) which all downregulate one another and upregulate themselves. This causes a branching to occur in the trajectory.
<code>num_modules</code>	The number of modules this component is allowed to use. Various components might require a minimum number of components in order to work properly.
<code>burn</code>	Whether or not these components are part of the warm-up simulation.
<code>num_steps</code>	The number of branching steps to reduce the odds of double positive cells occurring.

Details

A backbone always needs to start with a single `bblego_start()` state and needs to end with one or more `bblego_end()` states. The order of the mentioned states needs to be such that a state is never specified in the first argument (except for `bblego_start()`) before having been specified as the second argument.

Value

A dyngen backbone.

Examples

```
backbone <- bblego(
  bblego_start("A", type = "simple", num_modules = 2),
  bblego_linear("A", "B", type = "simple", num_modules = 3),
  bblego_branching("B", c("C", "D"), type = "simple", num_steps = 3),
  bblego_end("C", type = "flipflop", num_modules = 4),
  bblego_end("D", type = "doublerep1", num_modules = 7)
)
```

 combine_models

Combine multiple dyngen models

Description

Assume the given models have the exact same feature ids and ran up until the `generate_cells()` step. In addition, the user is expected to run `generate_experiment()` on the combined models.

Usage

```
combine_models(models, duplicate_gold_standard = TRUE)
```

Arguments

models	A named list of models. The names of the list will be used to prefix the different cellular states in the combined model.
duplicate_gold_standard	Whether or not the gold standards of the models are different and should be duplicated and prefixed.

Details

See the [vignette on simulating batch effects](#) on how to use this function.

Value

A combined dyngen model.

Examples

```
data("example_model")
model_ab <- combine_models(list("left" = example_model, "right" = example_model))

# show a dimensionality reduction
plot_simulations(model_ab)
plot_gold_mappings(model_ab, do_facet = FALSE)
```

dyngen *dyngen: A multi-modal simulator for spearheading single-cell omics analyses*

Description

A toolkit for generating synthetic single cell data.

Step 1, initialise dyngen model

- `initialise_model()`: Define and store settings for all following steps. See each of the sections below for more information.
- Use a predefined backbone:
 - `list_backbones()`
 - `backbone_bifurcating()`
 - `backbone_bifurcating_converging()`
 - `backbone_bifurcating_cycle()`
 - `backbone_bifurcating_loop()`
 - `backbone_branching()`
 - `backbone_binary_tree()`
 - `backbone_consecutive_bifurcating()`
 - `backbone_trifurcating()`
 - `backbone_converging()`
 - `backbone_cycle()`
 - `backbone_cycle_simple()`
 - `backbone_linear()`
 - `backbone_linear_simple()`
 - `backbone_disconnected()`
- Create a custom backbone:
 - `backbone()`
 - `bblego()`
 - `bblego_linear()`
 - `bblego_branching()`
 - `bblego_start()`
 - `bblego_end()`
- Visualise the backbone:
 - `plot_backbone_modulenet()`
 - `plot_backbone_statenet()`

Step 2, generate TF network

- `generate_tf_network()`: Generate a transcription factor network from the backbone
- `tf_network_default()`: Parameters for configuring this step

Step 3, add more genes to the gene network

- `generate_feature_network()`: Generate a target network
- `feature_network_default()`: Parameters for configuring this step
- `plot_feature_network()`: Visualise the gene network

Step 4, generate gene kinetics

- `generate_kinetics()`: Generate the gene kinetics
- `kinetics_default()`, `kinetics_random_distributions()`: Parameters for configuring this step

Step 5, simulate the gold standard

- `generate_gold_standard()`: Simulate the gold standard backbone, used for mapping to cell states afterwards
- `gold_standard_default()`: Parameters for configuring this step
- `plot_gold_mappings()`: Visualise the mapping of the simulations to the gold standard
- `plot_gold_simulations()`: Visualise the gold standard simulations using the dimred
- `plot_gold_expression()`: Visualise the expression of the gold standard over simulation time

Step 6, simulate the cells

- `generate_cells()`: Simulate the cells based on its GRN
- `simulation_default()`: Parameters for configuring this step
- `simulation_type_wild_type()`, `simulation_type_knockdown()`: Used for configuring the type of simulation
- `kinetics_noise_none()`, `kinetics_noise_simple()`: Different kinetics randomisers to apply to each simulation
- `plot_simulations()`: Visualise the simulations using the dimred
- `plot_simulation_expression()`: Visualise the expression of the simulations over simulation time

Step 7, simulate cell and transcripting sampling

- `generate_experiment()`: Sample cells and transcripts from experiment
- `list_experiment_samplers()`, `experiment_snapshot()`, `experiment_synchronised()`: Parameters for configuring this step
- `simtime_from_backbone()`: Determine the simulation time from the backbone
- `plot_experiment_dimred()`: Plot a dimensionality reduction of the final dataset

Step 8, convert to dataset

- `as_dyno()`, `wrap_dataset()`: Convert a dyngen model to a dyno dataset
- `as_anndata()`: Convert a dyngen model to an anndata dataset
- `as_sce()`: Convert a dyngen model to a SingleCellExperiment dataset
- `as_seurat()`: Convert a dyngen model to a Seurat dataset

One-shot function

- `generate_dataset()`: Run through steps 2 to 8 with a single function
- `plot_summary()`: Plot a summary of all dyngen simulation steps

Data objects

- `example_model`: A (very) small toy dyngen model, used for documentation and testing purposes
- `realcounts`: A set of real single-cell expression datasets, to be used as reference datasets
- `realnets`: A set of real gene regulatory networks, to be sampled in step 3

Varia functions

- `dyngen`: This help page
- `get_timings()`: Extract execution timings for each of the dyngen steps
- `combine_models()`: Combine multiple dyngen models
- `rnorm_bounded()`: A bounded version of `rnorm()`
- `runif_subrange()`: A subrange version of `runif()`

Examples

```

model <- initialise_model(
  backbone = backbone_bifurcating()
)

model <- model %>%
  generate_tf_network() %>%
  generate_feature_network() %>%
  generate_kinetics() %>%
  generate_gold_standard() %>%
  generate_cells() %>%
  generate_experiment()

dataset <- wrap_dataset(model, format = "dyno")
# format can also be set to "sce", "seurat", "anndata" or "list"

# library(dynplot)
# plot_dimred(dataset)

```

example_model	<i>A (very!) small toy dyngen model</i>
---------------	-----------------------------------------

Description

Used for showcasing examples of functions.

Usage

```
example_model
```

Format

An object of class list (inherits from dyngen::init) of length 19.

generate_cells	<i>Simulate the cells</i>
----------------	---------------------------

Description

[generate_cells\(\)](#) runs simulations in order to determine the gold standard of the simulations. [simulation_default\(\)](#) is used to configure parameters pertaining this process.

Usage

```
generate_cells(model)

simulation_default(
  burn_time = NULL,
  total_time = NULL,
  ssa_algorithm = ssa_etl(tau = 30/3600),
  census_interval = 4,
  experiment_params = bind_rows(simulation_type_wild_type(num_simulations = 32),
    simulation_type_knockdown(num_simulations = 0)),
  store_reaction_firings = FALSE,
  store_reaction_propensities = FALSE,
  compute_cellwise_grn = FALSE,
  compute_dimred = TRUE,
  compute_rna_velocity = FALSE,
  kinetics_noise_function = kinetics_noise_simple(mean = 1, sd = 0.005)
)

simulation_type_wild_type(
  num_simulations,
  seed = sample.int(10 * num_simulations, num_simulations)
```

```

)

simulation_type_knockdown(
  num_simulations,
  timepoint = runif(num_simulations),
  genes = "*",
  num_genes = sample(1:5, num_simulations, replace = TRUE, prob = 0.25^(1:5)),
  multiplier = runif(num_simulations, 0, 1),
  seed = sample.int(10 * num_simulations, num_simulations)
)

```

Arguments

model	A dyngen intermediary model for which the gold standard been generated with generate_gold_standard() .
burn_time	The burn in time of the system, used to determine an initial state vector. If NULL, the burn time will be inferred from the backbone.
total_time	The total simulation time of the system. If NULL, the simulation time will be inferred from the backbone.
ssa_algorithm	Which SSA algorithm to use for simulating the cells with GillespieSSA2::ssa()
census_interval	A granularity parameter for the outputted simulation.
experiment_params	A tibble generated by rbinding multiple calls of simulation_type_wild_type() and simulation_type_knockdown() .
store_reaction_firings	Whether or not to store the number of reaction firings.
store_reaction_propensities	Whether or not to store the propensity values of the reactions.
compute_cellwise_grn	Whether or not to compute the cellwise GRN activation values.
compute_dimred	Whether to perform a dimensionality reduction after simulation.
compute_rna_velocity	Whether or not to compute the propensity ratios after simulation.
kinetics_noise_function	A function that will generate noise to the kinetics of each simulation. It takes the <code>feature_info</code> and <code>feature_network</code> as input parameters, modifies them, and returns them as a list. See kinetics_noise_none() and kinetics_noise_simple() .
num_simulations	The number of simulations to run.
seed	A set of seeds for each of the simulations.
timepoint	The relative time point of the knockdown
genes	Which genes to sample from. "*" for all genes.
num_genes	The number of genes to knockdown.
multiplier	The strength of the knockdown. Use 0 for a full knockout, $0 < x < 1$ for a knock-down, and > 1 for an overexpression.

Value

A dyngen model.

See Also

[dyngen](#) on how to run a complete dyngen simulation

Examples

```
library(dplyr)
model <-
  initialise_model(
    backbone = backbone_bifurcating(),
    simulation = simulation_default(
      ssa_algorithm = ssa_etl(tau = .1),
      experiment_params = bind_rows(
        simulation_type_wild_type(num_simulations = 4),
        simulation_type_knockdown(num_simulations = 4)
      )
    )
  )

data("example_model")
model <- example_model %>% generate_cells()

plot_simulations(model)
plot_gold_mappings(model)
plot_simulation_expression(model)
```

generate_dataset

Generate a dataset

Description

This function contains the complete pipeline for generating a dataset with **dyngen**. In order to have more control over how the dataset is generated, run each of the steps in this function separately.

Usage

```
generate_dataset(
  model,
  format = c("list", "dyno", "sce", "seurat", "anndata", "none"),
  output_dir = NULL,
  make_plots = FALSE,
  store_dimred = model$simulation_params$compute_dimred,
  store_cellwise_grn = model$simulation_params$compute_cellwise_grn,
  store_rna_velocity = model$simulation_params$compute_rna_velocity
)
```

Arguments

model	A dyngen initial model created with <code>initialise_model()</code> .
format	Which output format to use, must be one of 'dyno' (requires dynwrap), 'sce' (requires SingleCellExperiment), 'seurat' (requires Seurat), 'anndata' (requires anndata), 'list' or 'none'.
output_dir	If not NULL, then the generated model and dynwrap dataset will be written to files in this directory.
make_plots	Whether or not to generate an overview of the dataset.
store_dimred	Whether or not to store the dimensionality reduction constructed on the true counts.
store_cellwise_grn	Whether or not to also store cellwise GRN information.
store_rna_velocity	Whether or not to store the log propensity ratios.

Value

A list containing a dyngen model (`li$model`) and a dynwrap dataset (`li$dataset`).

Examples

```

model <-
  initialise_model(
    backbone = backbone_bifurcating()
  )

# generate dataset and output as a list format
# please note other output formats exist: "dyno", "sce", "seurat", "anndata"
out <- generate_dataset(model, format = "list")

model <- out$model
dataset <- out$dataset

```

generate_experiment *Sample cells from the simulations*

Description

`generate_experiment()` samples cells along the different simulations. Two approaches are implemented: sampling from an unsynchronised population of single cells (snapshot) or sampling at multiple time points in a synchronised population (time series).

Usage

```

generate_experiment(model)

list_experiment_samplers()

experiment_snapshot(
  realcount = NULL,
  map_reference_cpm = TRUE,
  map_reference_ls = TRUE,
  weight_bw = 0.1
)

experiment_synchronised(
  realcount = NULL,
  map_reference_cpm = TRUE,
  map_reference_ls = TRUE,
  num_timepoints = 8,
  pct_between = 0.75
)

```

Arguments

model	A dyngen intermediary model for which the simulations have been run with generate_cells() .
realcount	The name of a dataset in realcounts . If NULL, a random dataset will be sampled from realcounts .
map_reference_cpm	Whether or not to try to match the CPM distribution to that of a reference dataset.
map_reference_ls	Whether or not to try to match the distribution of the library sizes to that of the reference dataset.
weight_bw	[snapshot] A bandwidth parameter for determining the distribution of cells along each edge in order to perform weighted sampling.
num_timepoints	[synchronised] The number of time points used in the experiment.
pct_between	[synchronised] The percentage of 'unused' simulation time.

Details

[experiment_snapshot\(\)](#) samples the cells using the length of each edge in the milestone network as weights. See Supplementary Figure 7A from the dyngen paper for an illustration of how these weights are computed.

[experiment_synchronised\(\)](#) samples the cells along the simulation timeline by binning it into `num_timepoints` groups separated by `num_timepoints-1` gaps. See Supplementary Figure 7B from the dyngen paper for an illustration of how the timepoint groups are computed.

Value

A dyngen model.

Examples

```

names(list_experiment_samplers())

model <-
  initialise_model(
    backbone = backbone_bifurcating(),
    experiment = experiment_synchronised()
  )

data("example_model")
model <- example_model %>% generate_experiment()

plot_experiment_dimred(model)

```

```
generate_feature_network
```

Generate a target network

Description

`generate_feature_network()` generates a network of target genes that are regulated by the previously generated TFs, and also a separate network of housekeeping genes (HKs). `feature_network_default()` is used to configure parameters pertaining this process.

Usage

```

generate_feature_network(model)

feature_network_default(
  realnet = NULL,
  damping = 0.01,
  target_resampling = Inf,
  max_in_degree = 5
)

```

Arguments

<code>model</code>	A dynngen intermediary model for which the transcription network has been generated with <code>generate_tf_network()</code> .
<code>realnet</code>	The name of a gene regulatory network (GRN) in <code>realnets</code> . If NULL, a random network will be sampled from <code>realnets</code> . Alternatively, a custom GRN can be used by passing a weighted sparse matrix.
<code>damping</code>	A damping factor used for the page rank algorithm used to subsample the realnet.
<code>target_resampling</code>	How many targets / HKs to sample from the realnet per iteration.
<code>max_in_degree</code>	The maximum in-degree of a target / HK.

Value

A dyngen model.

See Also

[dyngen](#) on how to run a complete dyngen simulation

Examples

```
model <-  
  initialise_model(  
    backbone = backbone_bifurcating(),  
    feature_network = feature_network_default(damping = 0.1)  
  )  
  
data("example_model")  
model <- example_model %>%  
  generate_tf_network() %>%  
  generate_feature_network()  
  
plot_feature_network(model)
```

generate_gold_standard

Simulate the gold standard

Description

[generate_gold_standard\(\)](#) runs simulations in order to determine the gold standard of the simulations. [gold_standard_default\(\)](#) is used to configure parameters pertaining this process.

Usage

```
generate_gold_standard(model)  
  
gold_standard_default(  
  tau = 30/3600,  
  census_interval = 10/60,  
  simulate_targets = FALSE  
)
```

Arguments

model	A dyngen intermediary model for which the kinetics of the feature network has been generated with generate_kinetics() .
tau	The time step of the ODE algorithm used to generate the gold standard.

`census_interval`
A granularity parameter of the gold standard time steps. Should be larger than or equal to `tau`.

`simulate_targets`
Also simulate the targets during the gold standard simulation

Value

A dyngen model.

See Also

[dyngen](#) on how to run a complete dyngen simulation

Examples

```
model <-  
  initialise_model(  
    backbone = backbone_bifurcating(),  
    gold_standard = gold_standard_default(tau = .01, census_interval = 1)  
  )  
  
data("example_model")  
model <- example_model %>% generate_gold_standard()  
  
plot_gold_simulations(model)  
plot_gold_mappings(model)  
plot_gold_expression(model)
```

`generate_kinetics` *Determine the kinetics of the feature network*

Description

`generate_kinetics()` samples the kinetics of genes in the feature network for which the kinetics have not yet been defined. `kinetics_default()` is used to configure parameters pertaining this process. `kinetics_random_distributions()` will do the same, but the distributions are also randomised.

Usage

```
generate_kinetics(model)  
  
kinetics_default()  
  
kinetics_random_distributions()
```

Arguments

`model` A dyngen intermediary model for which the feature network has been generated with `generate_feature_network()`.

Details

To write different kinetics settings, you need to write three functions with interface `function(feature_info, feature_network)`. Described below are the default kinetics samplers.

`sampler_tfs()` mutates the `feature_info` data frame by adding the following columns:

- `transcription_rate`: the rate at which pre-mRNAs are transcribed, in pre-mRNA / hour. Default distribution: $U(1, 2)$.
- `translation_rate`: the rate at which mRNAs are translated into proteins, in protein / mRNA / hour. Default distribution: $U(100, 150)$.
- `mrna_half-life`: the half-life of (pre-)mRNA molecules, in hours. Default distribution: $U(2.5, 5)$.
- `protein_half-life`: the half-life of proteins, in hours. Default distribution: $U(5, 10)$.
- `splicing_rate`: the rate at which pre-mRNAs are spliced into mRNAs, in reactions / hour. Default value: $\log(2) / (10/60)$, which corresponds to a half-life of 10 minutes.
- `independence`: the degree to which all regulators need to be bound for transcription to occur (0), or whether transcription can occur if only one of the regulators is bound (1).

`sampler_nontfs()` samples the `transcription_rate`, `translation_rate`, `mrna_half-life` and `protein_half-life` from a supplementary file of Schwannhäusser et al., 2011, doi.org/10.1038/nature10098. `splicing_rate` is by default the same as in `sampler_tfs()`. `independence` is sampled from $U(0, 1)$.

`sampler_interactions()` mutates the `feature_network` data frame by adding the following columns.

- `effect`: the effect of the interaction; upregulating = +1, downregulating = -1. By default, sampled from -1, 1 with probabilities .25, .75.
- `strength`: the strength of the interaction. Default distribution: $10^U(0, 2)$.
- `hill`: the hill coefficient. Default distribution: $N(2, 2)$ with a minimum of 1 and a maximum of 10.

Value

A dyngen model.

See Also

[dyngen](#) on how to run a complete dyngen simulation

Examples

```
model <-
  initialise_model(
    backbone = backbone_bifurcating(),
    kinetics_params = kinetics_default()
  )

data("example_model")
model <- example_model %>%
  generate_kinetics()
```

generate_tf_network *Generate a transcription factor network from the backbone*

Description

[generate_tf_network\(\)](#) generates the transcription factors (TFs) that drive the dynamic process a cell undergoes. [tf_network_default\(\)](#) is used to configure parameters pertaining this process.

Usage

```
generate_tf_network(model)

tf_network_default(
  min_tfs_per_module = 1L,
  sample_num_regulators = function() 2,
  weighted_sampling = FALSE
)
```

Arguments

model A dyngen initial model created with [initialise_model\(\)](#).

min_tfs_per_module The number of TFs to generate per module in the backbone.

sample_num_regulators A function to generate the number of TFs per module each TF will be regulated by.

weighted_sampling When determining what TFs another TF is regulated by, whether to perform weighted sampling (by rank) or not.

Value

A dyngen model.

See Also

[dyngen](#) on how to run a complete dyngen simulation

Examples

```
model <-  
  initialise_model(  
    backbone = backbone_bifurcating()  
  )  
model <- model %>%  
  generate_tf_network()  
  
plot_feature_network(model)
```

get_timings

Return the timings of each of the dyngen steps

Description

Return the timings of each of the dyngen steps

Usage

```
get_timings(model)
```

Arguments

model A dyngen object

Value

A data frame with columns "group", "task", "time_elapsed".

Examples

```
data("example_model")  
timings <- get_timings(example_model)
```

initialise_model	<i>Initial settings for simulating a dyngen dataset</i>
------------------	---------------------------------------------------------

Description

Initial settings for simulating a dyngen dataset

Usage

```
initialise_model(
  backbone,
  num_cells = 1000,
  num_tfs = nrow(backbone$module_info),
  num_targets = 100,
  num_hks = 50,
  distance_metric = c("pearson", "spearman", "cosine", "euclidean", "manhattan"),
  tf_network_params = tf_network_default(),
  feature_network_params = feature_network_default(),
  kinetics_params = kinetics_default(),
  gold_standard_params = gold_standard_default(),
  simulation_params = simulation_default(),
  experiment_params = experiment_snapshot(),
  verbose = TRUE,
  download_cache_dir = getOption("dyngen_download_cache_dir"),
  num_cores = getOption("Ncpus") %||% 1L,
  id = NULL
)
```

Arguments

backbone	The gene module configuration that determines the type of dynamic process being simulated. See list_backbones() for a full list of different backbones available in this package.
num_cells	The number of cells to sample.
num_tfs	The number of transcription factors (TFs) to generate. TFs are the main drivers of the changes that occur in a cell. TFs are regulated only by other TFs.
num_targets	The number of target genes to generate. Target genes are regulated by TFs and sometimes by other target genes.
num_hks	The number of housekeeping genes (HKs) to generate. HKs are typically highly expressed, and are not regulated by the TFs or targets.
distance_metric	The distance metric to be used to calculate the distance between cells. See dynutils::calculate_distance() for a list of possible distance metrics.
tf_network_params	Settings for generating the TF network with generate_tf_network() , see tf_network_default() .

feature_network_params	Settings for generating the feature network with generate_feature_network() , see feature_network_default() .
kinetics_params	Settings for determining the kinetics of the feature network with generate_kinetics() , see kinetics_default() .
gold_standard_params	Settings pertaining simulating the gold standard with generate_gold_standard() , see gold_standard_default() .
simulation_params	Settings pertaining the simulation itself with generate_cells() , see simulation_default() .
experiment_params	Settings related to how the experiment is simulated with generate_experiment() , see experiment_snapshot() or experiment_synchronised() .
verbose	Whether or not to print messages during the simulation.
download_cache_dir	If not NULL, temporary downloaded files will be cached in this directory.
num_cores	Parallellisation parameter for various steps in the pipeline.
id	An identifier for the model.

Value

A dyngen model.

See Also

[dyngen](#) on how to run a complete dyngen simulation

Examples

```
model <- initialise_model(
  backbone = backbone_bifurcating(),
  num_cells = 555,
  verbose = FALSE,
  download_cache_dir = "~/cache/dyngen"
)
```

kinetics_noise_none *Add small noise to the kinetics of each simulation*

Description

Add small noise to the kinetics of each simulation

Usage

```
kinetics_noise_none()

kinetics_noise_simple(mean = 1, sd = 0.005)
```

Arguments

mean	The mean level of noise (should be 1)
sd	The sd of the noise (should be a relatively small value)

Value

A list of noise generators for the kinetics.

list_backbones	<i>List of all predefined backbone models</i>
----------------	-----------------------------------------------

Description

A module is a group of genes which, to some extent, shows the same expression behaviour. Several modules are connected together such that one or more genes from one module will regulate the expression of another module. By creating chains of modules, a dynamic behaviour in gene regulation can be created.

Usage

```
list_backbones()

backbone_bifurcating()

backbone_bifurcating_converging()

backbone_bifurcating_cycle()

backbone_bifurcating_loop()

backbone_branching(
  num_modifications = rbinom(1, size = 6, 0.25) + 1,
  min_degree = 3,
  max_degree = sample(min_degree:5, 1)
)

backbone_binary_tree(num_modifications = rbinom(1, size = 6, 0.25) + 1)

backbone_consecutive_bifurcating()
```

```

backbone_trifurcating()

backbone_converging()

backbone_cycle()

backbone_cycle_simple()

backbone_linear()

backbone_linear_simple()

backbone_disconnected(
  left_backbone = list_backbones() %>% keep(., names(.) != "disconnected") %>%
    sample(1) %>% first(),
  right_backbone = list_backbones() %>% keep(., names(.) != "disconnected") %>%
    sample(1) %>% first(),
  num_common_modules = 10
)

```

Arguments

num_modifications	The number of branch points in the generated backbone.
min_degree	The minimum degree of each node in the backbone.
max_degree	The maximum degree of each node in the backbone.
left_backbone	A backbone (other than a disconnected backbone), see list_backbones() .
right_backbone	A backbone (other than a disconnected backbone), see list_backbones() .
num_common_modules	The number of modules which are regulated by either backbone.

Value

A list of all the available backbone generators.

See Also

[dyngen](#) on how to run a dyngen simulation

Examples

```

names(list_backbones())

bb <- backbone_bifurcating()
bb <- backbone_bifurcating_converging()
bb <- backbone_bifurcating_cycle()
bb <- backbone_bifurcating_loop()
bb <- backbone_binary_tree()
bb <- backbone_branching()

```

```
bb <- backbone_consecutive_bifurcating()
bb <- backbone_converging()
bb <- backbone_cycle()
bb <- backbone_cycle_simple()
bb <- backbone_disconnected()
bb <- backbone_linear()
bb <- backbone_linear_simple()
bb <- backbone_trifurcating()

model <- initialise_model(
  backbone = bb
)
```

plot_backbone_modulenet

Visualise the backbone of a model

Description

Visualise the backbone of a model

Usage

```
plot_backbone_modulenet(model)
```

Arguments

model A dynngen initial model created with `initialise_model()`.

Value

A ggplot2 object.

Examples

```
data("example_model")
plot_backbone_modulenet(example_model)
```

plot_backbone_statenet

Visualise the backbone state network of a model

Description

Visualise the backbone state network of a model

Usage

```
plot_backbone_statenet(model, detailed = FALSE)
```

Arguments

model	A dyngen initial model created with <code>initialise_model()</code> .
detailed	Whether or not to also plot the substates of transitions.

Value

A ggplot2 object.

Examples

```
data("example_model")  
plot_backbone_statenet(example_model)
```

plot_experiment_dimred

Plot a dimensionality reduction of the final dataset

Description

Plot a dimensionality reduction of the final dataset

Usage

```
plot_experiment_dimred(model, mapping = aes_string("comp_1", "comp_2"))
```

Arguments

model	A dyngen intermediary model for which the simulations have been run with <code>generate_experiment()</code> .
mapping	Which components to plot.

Value

A ggplot2 object.

Examples

```
data("example_model")
plot_experiment_dimred(example_model)
```

plot_feature_network *Visualise the feature network of a model*

Description

Visualise the feature network of a model

Usage

```
plot_feature_network(  
  model,  
  show_tfs = TRUE,  
  show_targets = TRUE,  
  show_hks = FALSE  
)
```

Arguments

model	A dyngen intermediary model for which the feature network has been generated with generate_feature_network() .
show_tfs	Whether or not to show the transcription factors.
show_targets	Whether or not to show the targets.
show_hks	Whether or not to show the housekeeping genes.

Value

A ggplot2 object.

Examples

```
data("example_model")
plot_feature_network(example_model)
```

plot_gold_expression *Visualise the expression of the gold standard over simulation time*

Description

Visualise the expression of the gold standard over simulation time

Usage

```
plot_gold_expression(  
  model,  
  what = c("mol_premrna", "mol_mrna", "mol_protein"),  
  label_changing = TRUE  
)
```

Arguments

model A dyngen intermediary model for which the simulations have been run with [generate_gold_standard\(\)](#).

what Which molecule types to visualise.

label_changing Whether or not to add a label next to changing molecules.

Value

A ggplot2 object.

Examples

```
data("example_model")  
plot_gold_expression(example_model, what = "mol_mrna", label_changing = FALSE)
```

plot_gold_mappings *Visualise the mapping of the simulations to the gold standard*

Description

Visualise the mapping of the simulations to the gold standard

Usage

```
plot_gold_mappings(  
  model,  
  selected_simulations = NULL,  
  do_facet = TRUE,  
  mapping = aes_string("comp_1", "comp_2")  
)
```

Arguments

model	A dyngen intermediary model for which the simulations have been run with <code>generate_cells()</code> .
selected_simulations	Which simulation indices to visualise.
do_facet	Whether or not to facet according to simulation index.
mapping	Which components to plot.

Value

A ggplot2 object.

Examples

```
data("example_model")
plot_gold_mappings(example_model)
```

plot_gold_simulations *Visualise the simulations using the dimred*

Description

Visualise the simulations using the dimred

Usage

```
plot_gold_simulations(  
  model,  
  detailed = FALSE,  
  mapping = aes_string("comp_1", "comp_2"),  
  highlight = 0  
)
```

Arguments

model	A dyngen intermediary model for which the simulations have been run with <code>generate_cells()</code> .
detailed	Whether or not to colour according to each separate sub-edge in the gold standard.
mapping	Which components to plot.
highlight	Which simulation to highlight. If <code>highlight == 0</code> then the gold simulation will be highlighted.

Value

A ggplot2 object.

Examples

```
data("example_model")  
plot_gold_simulations(example_model)
```

plot_simulations	<i>Visualise the simulations using the dimred</i>
------------------	---------------------------------------------------

Description

Visualise the simulations using the dimred

Usage

```
plot_simulations(model, mapping = aes_string("comp_1", "comp_2"))
```

Arguments

model	A dyngen intermediary model for which the simulations have been run with generate_cells() .
mapping	Which components to plot.

Value

A ggplot2 object.

Examples

```
data("example_model")  
plot_simulations(example_model)
```

`plot_simulation_expression`*Visualise the expression of the simulations over simulation time*

Description

Visualise the expression of the simulations over simulation time

Usage

```
plot_simulation_expression(  
  model,  
  simulation_i = 1:4,  
  what = c("mol_premrna", "mol_mrna", "mol_protein"),  
  facet = c("simulation", "module_group", "module_id", "none"),  
  label_nonzero = FALSE  
)
```

Arguments

<code>model</code>	A dyngen intermediary model for which the simulations have been run with generate_cells() .
<code>simulation_i</code>	Which simulation to visualise.
<code>what</code>	Which molecule types to visualise.
<code>facet</code>	What to facet on.
<code>label_nonzero</code>	Plot labels for non-zero molecules.

Value

A ggplot2 object.

Examples

```
data("example_model")  
plot_simulation_expression(example_model)
```

plot_summary	<i>Plot a summary of all dyngen simulation steps.</i>
--------------	-------------------------------------------------------

Description

Plot a summary of all dyngen simulation steps.

Usage

```
plot_summary(model)
```

Arguments

model	A dyngen intermediary model for which the simulations have been run with <code>generate_experiment()</code> .
-------	---------------------------------------------------------------------------------------------------------------

Value

A ggplot2 object.

Examples

```
data("example_model")  
plot_summary(example_model)
```

realcounts	<i>A set of real single cell expression datasets</i>
------------	------------------------------------------------------

Description

Statistics are derived from these datasets in order to simulate single cell experiments.

Usage

```
realcounts
```

Format

An object of class `tbl_df` (inherits from `tbl`, `data.frame`) with 111 rows and 9 columns.

realnets	<i>A set of gold standard gene regulatory networks</i>
----------	--------------------------------------------------------

Description

These networks are subsampled in order to generate realistic feature and housekeeping networks.

Usage

```
realnets
```

Format

An object of class `tbl_df` (inherits from `tbl`, `data.frame`) with 32 rows and 2 columns.

<code>rnorm_bounded</code>	<i>A bounded version of <code>rnorm</code></i>
----------------------------	------------------------------------------------

Description

A bounded version of `rnorm`

Usage

```
rnorm_bounded(n, mean = 0, sd = 1, min = -Inf, max = Inf)
```

Arguments

<code>n</code>	number of observations. If <code>length(n) > 1</code> , the length is taken to be the number required.
<code>mean</code>	vector of means.
<code>sd</code>	vector of standard deviations.
<code>min</code>	lower limits of the distribution.
<code>max</code>	upper limits of the distribution.

Value

Generates values with `rnorm`, bounded by `[min, max]`

Examples

```
rnorm_bounded(10)
```

runif_subrange	<i>A subrange version of runif</i>
----------------	------------------------------------

Description

Will generate numbers from a random subrange within the given range. For example, if [min, max] is set to `\[0, 10\]`, this function could decide to generate numbers between 2 and 6.

Usage

```
runif_subrange(n, min, max)
```

Arguments

n	Number of observations
min	Lower limits of the distribution.
max	Upper limits of the distribution.

Value

Generates values with runif, bounded by a range drawn from `sort(runif(2, min, max))`.

Examples

```
runif_subrange(20, 0, 10)
```

simtime_from_backbone	<i>Determine simulation time from backbone</i>
-----------------------	------------------------------------------------

Description

Determine simulation time from backbone

Usage

```
simtime_from_backbone(backbone, burn = FALSE)
```

Arguments

backbone	A valid dyngen backbone object
burn	Whether or not to compute the simtime for only the burn phase

Value

An estimation of the required simulation time

Examples

```
backbone <- backbone_linear()

simtime_from_backbone(backbone)

model <- initialise_model(
  backbone = backbone,
  simulation_params = simulation_default(
    burn_time = simtime_from_backbone(backbone, burn = TRUE),
    total_time = simtime_from_backbone(backbone, burn = FALSE)
  )
)
```

Index

- * **datasets**
 - example_model, 12
 - realcounts, 34
 - realnets, 35
- as_anndata (as_dyno), 3
- as_anndata(), 11
- as_dyno, 3
- as_dyno(), 11
- as_list (as_dyno), 3
- as_sce (as_dyno), 3
- as_sce(), 11
- as_seurat (as_dyno), 3
- as_seurat(), 11

- backbone, 4
- backbone(), 9
- backbone_bifurcating (list_backbones), 25
- backbone_bifurcating(), 9
- backbone_bifurcating_converging (list_backbones), 25
- backbone_bifurcating_converging(), 9
- backbone_bifurcating_cycle (list_backbones), 25
- backbone_bifurcating_cycle(), 9
- backbone_bifurcating_loop (list_backbones), 25
- backbone_bifurcating_loop(), 9
- backbone_binary_tree (list_backbones), 25
- backbone_binary_tree(), 9
- backbone_branching (list_backbones), 25
- backbone_branching(), 9
- backbone_consecutive_bifurcating (list_backbones), 25
- backbone_consecutive_bifurcating(), 9
- backbone_converging (list_backbones), 25
- backbone_converging(), 9
- backbone_cycle (list_backbones), 25
- backbone_cycle(), 9
- backbone_cycle_simple (list_backbones), 25
- backbone_cycle_simple(), 9
- backbone_disconnected (list_backbones), 25
- backbone_disconnected(), 9
- backbone_linear (list_backbones), 25
- backbone_linear(), 9
- backbone_linear_simple (list_backbones), 25
- backbone_linear_simple(), 9
- backbone_trifurcating (list_backbones), 25
- backbone_trifurcating(), 9
- bblego, 6
- bblego(), 9
- bblego_branching (bblego), 6
- bblego_branching(), 9
- bblego_end (bblego), 6
- bblego_end(), 9
- bblego_linear (bblego), 6
- bblego_linear(), 9
- bblego_start (bblego), 6
- bblego_start(), 9

- combine_models, 8
- combine_models(), 11

- dyngen, 5, 9, 11, 14, 18–20, 22, 24, 26
- dynutils::calculate_distance(), 23

- example_model, 11, 12
- experiment_snapshot (generate_experiment), 15
- experiment_snapshot(), 10, 16, 24
- experiment_synchronised (generate_experiment), 15
- experiment_synchronised(), 10, 16, 24

- feature_network_default
 - (generate_feature_network), 17
- feature_network_default(), 10, 17, 24
- generate_cells, 12
- generate_cells(), 10, 12, 16, 24, 31–33
- generate_dataset, 14
- generate_dataset(), 11
- generate_experiment, 15
- generate_experiment(), 4, 10, 15, 24, 28, 34
- generate_feature_network, 17
- generate_feature_network(), 10, 17, 20, 24, 29
- generate_gold_standard, 18
- generate_gold_standard(), 10, 13, 18, 24, 30
- generate_kinetics, 19
- generate_kinetics(), 10, 18, 19, 24
- generate_tf_network, 21
- generate_tf_network(), 9, 17, 21, 23
- get_timings, 22
- get_timings(), 11
- GillespieSSA2::ssa(), 13
- gold_standard_default
 - (generate_gold_standard), 18
- gold_standard_default(), 10, 18, 24
- initialise_model, 23
- initialise_model(), 9, 15, 21, 27, 28
- kinetics_default (generate_kinetics), 19
- kinetics_default(), 10, 19, 24
- kinetics_noise_none, 24
- kinetics_noise_none(), 10, 13
- kinetics_noise_simple
 - (kinetics_noise_none), 24
- kinetics_noise_simple(), 10, 13
- kinetics_random_distributions
 - (generate_kinetics), 19
- kinetics_random_distributions(), 10, 19
- list_backbones, 25
- list_backbones(), 9, 23, 26
- list_experiment_samplers
 - (generate_experiment), 15
- list_experiment_samplers(), 10
- plot_backbone_modulenet, 27
- plot_backbone_modulenet(), 9
- plot_backbone_statenet, 28
- plot_backbone_statenet(), 9
- plot_experiment_dimred, 28
- plot_experiment_dimred(), 10
- plot_feature_network, 29
- plot_feature_network(), 10
- plot_gold_expression, 30
- plot_gold_expression(), 10
- plot_gold_mappings, 30
- plot_gold_mappings(), 10
- plot_gold_simulations, 31
- plot_gold_simulations(), 10
- plot_simulation_expression, 33
- plot_simulation_expression(), 10
- plot_simulations, 32
- plot_simulations(), 10
- plot_summary, 34
- plot_summary(), 11
- realcounts, 11, 16, 34
- realnets, 11, 17, 35
- rnorm(), 11
- rnorm_bounded, 35
- rnorm_bounded(), 11
- runif(), 11
- runif_subrange, 36
- runif_subrange(), 11
- simtime_from_backbone, 36
- simtime_from_backbone(), 10
- simulation_default (generate_cells), 12
- simulation_default(), 10, 12, 24
- simulation_type_knockdown
 - (generate_cells), 12
- simulation_type_knockdown(), 10, 13
- simulation_type_wild_type
 - (generate_cells), 12
- simulation_type_wild_type(), 10, 13
- tf_network_default
 - (generate_tf_network), 21
- tf_network_default(), 9, 21, 23
- wrap_dataset (as_dyno), 3
- wrap_dataset(), 11