Package 'scGate'

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Type Package

Title Marker-Based Cell Type Purification for Single-Cell Sequencing Data

Version 1.6.2

Description

A common bioinformatics task in single-cell data analysis is to purify a cell type or cell population of interest from heterogeneous datasets. 'scGate' automatizes marker-based purification of specific cell populations, without requiring training data or reference gene expression profiles. Briefly, 'scGate' takes as input: i) a gene expression matrix stored in a 'Seurat' object and ii) a "gating model" (GM), consisting of a set of marker genes that define the cell population of interest. The GM can be as simple as a single marker gene, or a combination of positive and negative markers. More complex GMs can be constructed in a hierarchical fashion, akin to gating strategies employed in flow cytometry. 'scGate' evaluates the strength of signature marker expression in each cell using the rank-based method 'UCell', and then performs knearest neighbor (kNN) smoothing by calculating the mean 'UCell' score across neighboring cells. kNN-smoothing aims at compensating for the large degree of sparsity in scRNA-seq data. Finally, a universal threshold over kNN-smoothed signature scores is applied in binary decision trees generated from the user-provided gating model, to annotate cells as either "pure" or "impure", with respect to the cell population of interest. See the related publication Andreatta et al. (2022) <doi:10.1093/bioinformatics/btac141>.

biocViews

Depends R (>= 4.3.0)

Imports Seurat (>= 4.0.0), UCell (>= 2.6.0), dplyr, stats, utils, methods, patchwork, ggridges, reshape2, ggplot2, BiocParallel

Suggests ggparty, partykit, knitr, rmarkdown

VignetteBuilder knitr

URL https://github.com/carmonalab/scGate

BugReports https://github.com/carmonalab/scGate/issues

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combine_scGate_multiclass

Combine scGate annotations

Description

If a single-cell dataset has precomputed results for multiple scGate models, combined them in multi-class annotation

```
combine_scGate_multiclass(
  obj,
  prefix = "is.pure_",
  scGate_classes = NULL,
  min_cells = 1,
  multi.asNA = FALSE,
  out_column = "scGate_multi"
)
```

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Arguments

obj	Seurat object with scGate results for multiple models stored as metadata	
prefix	Prefix in metadata column names for scGate result models	
scGate_classes	Vector of scGate model names. If NULL, use all columns that start with "prefix" above.	
min_cells	Minimum number of cells for a cell label to be considered	
multi.asNA	How to label cells that are "Pure" for multiple annotations: "Multi" (FALSE) or NA (TRUE) $$	
out_column	The name of the metadata column where to store the multi-class cell labels	

Value

A Seurat object with multi-class annotations based on the combination of multiple models. A new column (by default "scGate_multi") is added to the metadata of the Seurat object.

Examples

```
# Define gating models
model.B <- gating_model(name = "Bcell", signature = c("MS4A1"))
model.T <- gating_model(name = "Tcell", signature = c("CD2","CD3D","CD3E"))
# Apply scGate with these models
data(query.seurat)
query.seurat <- scGate(query.seurat, model=model.T,
    reduction="pca", output.col.name = "is.pure_Tcell")
query.seurat <- scGate(query.seurat, model=model.B,
    reduction="pca", output.col.name = "is.pure_Bcell")
query.seurat <- combine_scGate_multiclass(query.seurat, scGate_class=c("Tcell","Bcell"))
table(query.seurat$scGate_multi)</pre>
```

gating_model

Model creation and editing

Description

Generate an scGate model from scratch or edit an existing one

```
gating_model(
  model = NULL,
  level = 1,
  name,
  signature,
  positive = TRUE,
```

```
negative = FALSE,
remove = FALSE
)
```

Arguments

model scGate model to be modified. When is NULL (default) a new model will be initialized. level integer. It refers to the hierarchical level of the model tree in which the signature will be added (level=1 by default) Arbitrary signature name (i.e. Immune, Tcell, NK etc). name character vector indicating gene symbols to be included in the signature (e.g. signature CD3D). If a minus sign is placed to the end of a gene name (e.g. "CD3D-"), this gene will be used as negative in UCell computing. See UCell documentation for details positive Logical indicating if the signature must be used as a positive signature in those model level. Default is TRUE. Same as 'positive' but negated (negative=TRUE equals to positive=FALSE) negative

Whether to remove the given signature from the model

Value

remove

A scGate model that can be used by scGate to filter target cell types.

Examples

genes.blacklist.default

Blocklist of genes for dimensionality reduction

Description

A list of signatures, for mouse and human. These include cell cycling, heat-shock genes, mitochondrial genes, and other genes classes, that may confound the identification of cell types. These are used internally by scGate and excluded from the calculation of dimensional reductions (PCA).

Format

A list of signatures

get_scGateDB 5

|--|

Description

Download, update or load local version of the scGate model database. These are stored in a GitHub repository, from where you can download specific versions of the database.

Usage

```
get_scGateDB(
  destination = tempdir(),
  force_update = FALSE,
  version = "latest",
  branch = c("master", "dev"),
  verbose = FALSE,
  repo_url = "https://github.com/carmonalab/scGate_models"
)
```

Arguments

destination	Destination path for storing the DB. The default is tempdir(); if you wish to edit locally the models and link them to the current project, set this parameter to a new directory name, e.g. scGateDB
force_update	Whether to update an existing database.
version	Specify the version of the scGate_models database (e.g. 'v0.1'). By default downloads the latest available version.
branch	branch of the scGate model repository, either 'master' (default) or 'dev' for the latest models
verbose	display progress messages
repo_url	URL path to scGate model repository database

Details

Models for scGate are dataframes where each line is a signature for a given filtering level. A database of models can be downloaded using the function get_scGateDB. You may directly use the models from the database, or edit one of these models to generate your own custom gating model.

Value

A list of models, organized according to the folder structure of the database. See the examples below.

See Also

```
scGate load_scGate_model
```

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Examples

```
scGate.model.db <- get_scGateDB()
# To see a specific model, browse the list of models:
scGate.model.db$human$generic$Myeloid</pre>
```

get_testing_data

Download sample data

Description

Helper function to obtain some sample data

Usage

```
get_testing_data(version = "hsa.latest", destination = tempdir())
```

Arguments

version Which sample dataset destination Save to this directory

Value

A list of datasets that can be used to test scGate

Examples

```
testing.datasets <- get_testing_data(version = 'hsa.latest')</pre>
```

load_scGate_model

Load a single scGate model

Description

Loads a custom scGate model into R. For the format of these models, have a look or edit one of the default models obtained with $get_scGateDB$

```
load_scGate_model(model_file, master.table = "master_table.tsv")
```

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Arguments

model_file scGate model file, in .tsv format.

master.table File name of the master table (in repo_path folder) that contains cell type signa-

tures.

Value

A scGate model in dataframe format, which can given as input to the scGate function.

See Also

```
scGate get_scGateDB
```

Examples

```
dir <- tempdir() # this may also be set to your working directory
models <- get_scGateDB(destination=dir)
# Original or edited model
model.path <- paste0(dir,"/scGate_models-master/human/generic/Bcell_scGate_Model.tsv")
master.path <- paste0(dir,"/scGate_models-master/human/generic/master_table.tsv")
my.model <- load_scGate_model(model.path, master.path)
my.model</pre>
```

performance.metrics

Performance metrics

Description

Evaluate model performance for binary tasks

Usage

```
performance.metrics(actual, pred, return_contingency = FALSE)
```

Arguments

actual Logical or numeric binary vector giving the actual cell labels.

pred Logical or numeric binary vector giving the predicted cell labels.

return_contingency

Logical indicating if contingency table must be returned.

Value

Prediction performance metrics (Precision, Recall, MCC) between actual and predicted cell type labels.

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Examples

```
results <- performance.metrics(actual= sample(c(1,0),20,replace=TRUE), pred = sample(c(1,0),20,replace=TRUE,prob = c(0.65,0.35)))
```

plot_levels

Plot scGate filtering results by level

Description

Fast plotting of gating results over each model level.

Usage

```
plot_levels(obj, pure.col = "green", impure.col = "gray")
```

Arguments

obj Gated Seurat object output of scGate filtering function

pure.col Color code for pure category

impure.col Color code for impure category

Value

UMAP plots with 'Pure'/'Impure' labels for each level of the scGate model

Examples

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plot_tree

Plot model tree

Description

View scGate model as a decision tree (require ggparty package)

Usage

```
plot_tree(model, box.size = 8, edge.text.size = 4)
```

Arguments

```
model A scGate model to be visualized box.size Box size edge.text.size Edge text size
```

Value

A plot of the model as a decision tree. At each level, green boxes indicate the 'positive' (accepted) cell types, red boxed indicate the 'negative' cell types (filtered out). The final Pure population is the bottom right subset in the tree.

Examples

```
library(ggparty)
models <- get_scGateDB()
plot_tree(models$human$generic$Tcell)</pre>
```

plot_UCell_scores

Plot UCell scores by level

Description

Show distribution of UCell scores for each level of a given scGate model

```
plot_UCell_scores(
  obj,
  model,
  overlay = 5,
  pos.thr = 0.2,
  neg.thr = 0.2,
  ncol = NULL,
  combine = TRUE
)
```

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Arguments

obj	Gated Seurat object (output of scGate)
model	scGate model used to identify a target population in obj
overlay	Degree of overlay for ggridges
pos.thr	Threshold for positive signatures used in scGate model (set to NULL to disable)
neg.thr	Threshold for negative signatures used in scGate model (set to NULL to disable)
ncol	Number of columns in output object (passed to wrap_plots)
combine	Whether to combine plots into a single object, or to return a list of plots

Value

Returns a density plot of UCell scores for the signatures in the scGate model, for each level of the model

Either a plot combined by patchwork (combine=T) or a list of plots (combine=F)

Examples

query.seurat	Toy dataset to test the package	

Description

A downsampled version (300 cells) of the single-cell dataset by Zilionis et al. (2019) <doi:10.1016/j.immuni.2019.03.009>, with precalculated PCA and UMAP reductions.

Format

A Seurat object

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scGate

Filter single-cell data by cell type

Description

Apply scGate to filter specific cell types in a query dataset

Usage

```
scGate(
 data,
 model,
 pos.thr = 0.2,
 neg.thr = 0.2,
 assay = NULL,
  slot = "data",
 ncores = 1,
 BPPARAM = NULL,
  seed = 123,
  keep.ranks = FALSE,
  reduction = c("calculate", "pca", "umap", "harmony"),
 min.cells = 30,
 nfeatures = 2000,
 pca.dim = 30,
 param_decay = 0.25,
 maxRank = 1500,
  output.col.name = "is.pure",
 k.param = 30,
  smooth.decay = 0.1,
  smooth.up.only = FALSE,
  genes.blacklist = "default",
  return.CellOntology = TRUE,
 multi.asNA = FALSE,
 additional.signatures = NULL,
  save.levels = FALSE,
 verbose = FALSE,
 progressbar = T
)
```

Arguments

data	Seurat object containing a query data set - filtering will be applied to this object
model	A single scGate model, or a list of scGate models. See Details for this format
pos.thr	Minimum UCell score value for positive signatures
neg.thr	Maximum UCell score value for negative signatures
assay	Seurat assay to use

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slot Data slot in Seurat object to calculate UCell scores

ncores Number of processors for parallel processing

BPPARAM A [BiocParallel::bpparam()] object that tells scGate how to parallelize. If pro-

vided, it overrides the 'ncores' parameter.

seed Integer seed for random number generator

keep.ranks Store UCell rankings in Seurat object. This will speed up calculations if the

same object is applied again with new signatures.

reduction Dimensionality reduction to use for knn smoothing. By default, calculates a new

reduction based on the given assay; otherwise you may specify a precalculated dimensionality reduction (e.g. in the case of an integrated dataset after batch-

effect correction)

min.cells Minimum number of cells to cluster or define cell types

nfeatures Number of variable genes for dimensionality reduction

pca.dim Number of principal components for dimensionality reduction

param_decay Controls decrease in parameter complexity at each iteration, between 0 and 1.

param_decay == 0 gives no decay, increasingly higher param_decay gives in-

creasingly stronger decay

maxRank Maximum number of genes that UCell will rank per cell

output.col.name

Column name with 'pure/impure' annotation

k.param Number of nearest neighbors for knn smoothing

 $smooth.\,decay \qquad Decay \;parameter \;for \;knn \;weights: \;(1-decay)^{\wedge}n$

smooth.up.only If TRUE, only let smoothing increase signature scores

genes.blacklist

Genes blacklisted from variable features. The default loads the list of genes in scGate::genes.blacklist.default; you may deactivate blacklisting by

setting genes.blacklist=NULL

return.CellOntology

If TRUE Cell ontology name and id are returned as additional metadata columns

when running multiple models.

multi.asNA How to label cells that are "Pure" for multiple annotations: "Multi" (FALSE) or

NA (TRUE)

additional.signatures

A list of additional signatures, not included in the model, to be evaluated (e.g. a cycling signature). The scores for this list of signatures will be returned but not

used for filtering.

save.levels Whether to save in metadata the filtering output for each gating model level

verbose Verbose output

progressbar Whether to show a progressbar or not

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Details

Models for scGate are data frames where each line is a signature for a given filtering level. A database of models can be downloaded using the function get_scGateDB. You may directly use the models from the database, or edit one of these models to generate your own custom gating model.

Multiple models can also be evaluated at once, by running scGate with a list of models. Gating for each individual model is returned as metadata, with a consensus annotation stored in scGate_multi metadata field. This allows using scGate as a multi-class classifier, where only cells that are "Pure" for a single model are assigned a label, cells that are "Pure" for more than one gating model are labeled as "Multi", all others cells are annotated as NA.

Value

A new metadata column is.pure is added to the query Seurat object, indicating which cells passed the scGate filter. The active.ident is also set to this variable.

See Also

```
load_scGate_model get_scGateDB plot_tree
```

Examples

```
### Test using a small toy set
data(query.seurat)
# Define basic gating model for B cells
my_scGate_model <- gating_model(name = "Bcell", signature = c("MS4A1"))</pre>
query.seurat <- scGate(query.seurat, model = my_scGate_model, reduction="pca")</pre>
table(query.seurat$is.pure)
### Test with larger datasets
library(Seurat)
testing.datasets <- get_testing_data(version = 'hsa.latest')</pre>
seurat_object <- testing.datasets[["JerbyArnon"]]</pre>
# Download pre-defined models
models <- get_scGateDB()</pre>
seurat_object <- scGate(seurat_object, model=models$human$generic$PanBcell)</pre>
DimPlot(seurat_object)
seurat_object_filtered <- subset(seurat_object, subset=is.pure=="Pure")</pre>
### Run multiple models at once
models <- get_scGateDB()</pre>
model.list <- list("Bcell" = models$human$generic$Bcell,</pre>
                    "Tcell" = models$human$generic$Tcell)
seurat_object <- scGate(seurat_object, model=model.list)</pre>
DimPlot(seurat_object, group.by = "scGate_multi")
```

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test_my_model

Test your model

Description

Wrapper for fast model testing on 3 sampled datasets

Usage

```
test_my_model(
 model,
  testing.version = "hsa.latest",
  custom.dataset = NULL,
  target = NULL,
  plot = TRUE
)
```

Arguments

model scGate model in data.frame format testing.version

> Character indicating the version of testing tatasets to be used. By default "hsalatest" will be used. It will be ignored if a custom dataset is provided (in Seurat format).

custom.dataset Seurat object to be used as a testing dataset. For testing purposes, metadata seurat object must contain a column named 'cell_type' to be used as a gold standard. Also a set of positive targets must be provided in the target variable.

target

Positive target cell types. If default testing version is used this variable must be a character indicating one of the available target models ('immune','Lymphoid','Myeloid','Tcell','Bcell','C 'NK', 'MoMacDC', 'Plasma_cell', 'PanBcell'). If a custom dataset is provided in Seurat format, this variable must be a vector of positive cell types in your data. The last case also require that such labels were named as in your cell_type

meta.data column.

plot Whether to return plots to device

Value

Returns performance metrics for the benchmarking datasets, and optionally plots of the predicted cell type labels in reduced dimensionality space.

Examples

```
scGate.model.db <- get_scGateDB()</pre>
# Browse the list of models and select one:
model.panBcell <- scGate.model.db$human$generic$PanBcell</pre>
# Test the model with available testing datasets
```

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```
panBcell.performance <- test_my_model(model.panBcell, target = "PanBcell")
model.Myeloid <- scGate.model.db$human$generic$Myeloid
myeloid.performance <- test_my_model(model.Myeloid, target = "Myeloid")</pre>
```

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