Package 'iCellR'

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Type Package

Title iCellR; A Feature-rich Interactive R Package to Work with High-Throughput Single Cell Sequencing Technologies.

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Description

Single (i) Cell R package (iCellR) is an interactive R package to work with high-throughput single cell sequencing technologies (i.e scRNA-seq, scVDJ-seq and CITE-seq). iCellR is an R package that allows scientists unprecedented flexibility at every step of the analysis pipeline, including normalization, clustering, dimensionality reduction, imputation, visualization, and so on. iCellR allows users to design both unsupervised and supervised models to best suit their research. In addition, iCellR provides 2D and 3D interactive visualizations, differential expression analysis, filters based on cells, genes and clusters, data merging, normalizing for dropouts and filling them with imputation methods, batch differences, pathway analysis and tools to find marker genes for clusters and conditions, predict cell types and pseudotime analysis.

Keywords PCA, UMAP, Diffusion Map, tSNE, high throughput, single cell, differential gene expression, analysis, clustering, PseudoTime, dimension reduction, normalization, spike-in, scRNA-Seq, CITE-Seq, scVDJ-Seq, data visualization, interactive plots, 3D plots, cell type prediction, pathway analysis, batch correction, cell cycle, cell gating, ADT, filtering, Clustering methods: ward.D, ward.D2, single, complete, average, mcquitty, median, centroid, kmeans, distance calculation methods: euclidean, maximum, "manhattan, canberra, binary, minkowski, indexing methods: kl, ch, hartigan, ccc, scott, marriot, trcovw, tracew, fried-

2 R topics documented:

man, rubin, cindex, db, silhouette, duda, pseudot2, beale, ratkowsky, ball, ptbiserial, gap, frey, mcclain, gamma, gplus, tau, dunn, hubert, sdindex, dindex, sdbw, normalization methods: ranked.glsf, global.glsf, deseq, rpm, spike.in.

```
Depends R (>= 3.3.0), ggplot2, plotly
Imports Matrix,
Rtsne,
gridExtra,
ggpubr,
scatterplot3d,
RColorBrewer,
knitr,
NbClust,
reshape,
shiny,
```

License GPL-2

umap, pheatmap

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VignetteBuilder knitr

URL https://github.com/rezakj/iCellR

Suggests rmarkdown, Rmagic, phateR

R topics documented:

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Description

add.adt

This function takes a data frame of ADT values per cell and adds it to the iCellR object.

Add CITE-seq antibody-derived tags (ADT)

Usage

```
add.adt(x = NULL, adt.data = "data.frame")
```

Arguments

x An object of class iCellR.

adt.data A data frame containing ADT counts for cells.

Value

An object of class iCellR

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Examples

```
## Not run:
my.obj <- add.adt(my.obj, adt.data = adt.data)
## End(Not run)</pre>
```

add.vdj

Add V(D)J recombination data

Description

This function takes a data frame of VDJ information per cell and adds it to the iCellR object.

Usage

```
add.vdj(x = NULL, vdj.data = "data.frame")
```

Arguments

x An object of class iCellR.

adt.data A data frame containing VDJ information for cells.

Value

An object of class iCellR

Examples

```
## Not run:
my.obj <- add.adt(my.obj, adt.data = adt.data)
## End(Not run)</pre>
```

adt.rna.merge

Merge RNA and ADT data

Description

This function is to merge the RNA and ADT data to the main.data slot of the iCellR object.

Usage

```
adt.rna.merge(x = NULL, adt.data = "raw")
```

Arguments

An object of class iCellR.

adt.data Choose from raw or main (normalized) ADT data, default = "raw".

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Value

An object of class iCellR

Examples

```
## Not run:
my.obj <- adt.rna.merge(my.obj, adt.data = "raw")
## End(Not run)</pre>
```

CC

Calculate Cell cycle phase prediction

Description

This function takes an object of class iCellR and assignes cell cycle stage for the cells.

Usage

```
cc(object = NULL, s.genes = s.phase, g2m.genes = g2m.phase)
```

Arguments

object A data frame containing gene counts for cells.

s.genes Genes that are used as a marker for S phase.

g2m.genes Genes that are used as a marker for G2 and M phase.

Value

The data frame object

```
## Not run:
my.obj <- cc(my.obj, s.genes = s.phase, g2m.genes = g2m.phase)
## End(Not run)</pre>
```

6 cell.filter

cell.filter

Filter cells

Description

This function takes an object of class iCellR and filters the raw data based on the number of UMIs, genes per cell, percentage of mitochondrial genes per cell, genes, gene expression and cell ids.

Usage

```
cell.filter(x = NULL, min.mito = 0, max.mito = 1, min.genes = 0,
  max.genes = Inf, min.umis = 0, max.umis = Inf,
  filter.by.cell.id = "character", keep.cell.id = "character",
  filter.by.gene = "character", filter.by.gene.exp.min = 1)
```

Arguments

x	An object of class iCellR.
min.mito	Min rate for mitochondrial gene expression per cell, default = 0 .
max.mito	Max rate for mitochondrial gene expression per cell, default = 1.
min.genes	Min number genes per cell, default = 0 .
max.genes	Max number genes per cell, default = Inf.
min.umis	Min number UMIs per cell, default = 0 .
max.umis	Max number UMIs per cell, default = Inf.
filter.by.cell.id	
	A character vector of cell ids to be filtered out.
keep.cell.id	A character vector of cell ids to keep.
filter.by.gene	A character vector of gene names to be filtered by thier expression. If more then one gene is defined it would be OR not AND.
filter.by.gene.exp.min	
	Minimum gene expression to be filtered by the genes set in filter.by.gene, default = 1.

Value

An object of class iCellR.

```
## Not run:
cell.filter(my.obj,
   min.mito = 0,
   max.mito = 1,
   min.genes = 0,
   max.genes = Inf,
   min.umis = 0,
   max.umis = Inf)
## End(Not run)
```

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ting Cell gating

Description

This function takes an object of class iCellR and a 2D tSNE or UMAP plot and gates around cells to get their ids.

Usage

```
cell.gating(x = NULL, my.plot = NULL, plot.type = NULL)
```

Arguments

x An object of class iCellR.my.plot The plot to use for gating. Must be a 2D plot.plot.type Choose from UMAP and tSNE, default = NULL.

Value

An object of class iCellR.

Examples

```
## Not run:
cell.gating(my.obj, my.plot = PLOT, plot.type = "tsne")
## End(Not run)
```

cell.type.pred

Create heatmaps or dot plots for genes in clusters to find thier cell types using ImmGen data.

Description

This function takes an object of class iCellR and genes and provides a heatmap.

Usage

```
cell.type.pred(immgen.data = "rna", gene = "NULL",
  top.cell.types = 50, plot.type = "heatmap", heat.colors = c("blue",
  "white", "red"))
```

Arguments

```
immgen.data Choose from "rna", "uli.rna" or "mca", default = "rna" gene A set of gene names to used to predict cell type. top.cell.types Top cell types sorted by cumulative expression, default = 25. plot.type Choose from "heatmap" od "point.plot", default = "heatmap" heat.colors Colors for heatmap, default = c("blue", "white", "red").
```

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Value

An object of class iCellR

Examples

```
## Not run:
imm.gen(immgen.data = "uli.rna", gene = MyGenes, plot.type = "heatmap")
imm.gen(immgen.data = "rna", gene = MyGenes, plot.type = "point.plot")
## End(Not run)
```

change.clust

Change the cluster number or re-name them

Description

This function re-names the clusters in the best.clust slot of the iCellR object.

Usage

```
change.clust(x = NULL, change.clust = 0, to.clust = 0,
  clust.reset = F)
```

Arguments

x An object of class iCellR.

to.clust The new name for the cluster.
clust.reset Reset to the original clustering.

Value

An object of class iCellR.

```
## Not run:
my.obj <- change.clust(my.obj, change.clust = 3, to.clust = 1)
my.obj <- change.clust(my.obj, change.clust = 2, to.clust = "B Cell")
my.obj <- change.clust(my.obj, clust.reset = T)
## End(Not run)</pre>
```

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clono.plot

Make 2D and 3D scatter plots for clonotypes.

Description

This function takes an object of class iCellR and provides plots for clonotypes.

Usage

```
clono.plot(x = NULL, plot.data.type = "tsne", clono = 1,
  clust.dim = 2, cell.size = 1, cell.colors = c("red", "gray"),
  box.cell.col = "black", back.col = "white",
  cell.transparency = 0.5, interactive = TRUE, out.name = "plot")
```

Arguments

```
An object of class iCellR.
х
plot.data.type Choose from "tsne" and "pca", default = "tsne".
                   A clonotype name to be plotted, default = 1.
clono
clust.dim
                   2 for 2D plots and 3 for 3D plots, default = 2.
cell.size
                   A number for the size of the points in the plot, default = 1.
cell.colors
                   Colors for heat mapping the points in "scatterplot", default = c("gray","red").
back.col
                   A color for the plot background, default = "black".
cell.transparency
                   Color transparency for points, default = 0.5.
                   If set to TRUE an intractive HTML file will be created, default = TRUE.
interactive
out.name
                   If "interactive" is set to TRUE, the out put name for HTML, default = "plot".
```

Value

An object of class iCellR.

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clust.avg.exp

Create a data frame of mean expression of genes per cluster

Description

This function takes an object of class iCellR and creates an average gene expression for every cluster.

Usage

```
clust.avg.exp(x = NULL)
```

Arguments

Χ

An object of class iCellR.

Value

An object of class iCellR.

Examples

```
## Not run:
my.obj <- clust.avg.exp(my.obj)
## End(Not run)</pre>
```

clust.cond.info

Calculate cluster and conditions frequencies

Description

This function takes an object of class iCellR and calculates cluster and conditions frequencies.

Usage

```
clust.cond.info(x = NULL, plot.type = "pie", normalize.ncell = TRUE)
```

Arguments

```
x An object of class iCellR.
plot.type Choose from pie or bar, defult = pie.
normalize.ncell
```

If TRUE the values will be normalized to the number of cells by downsampling.

Value

An object of class iCellR.

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Examples

```
## Not run:
clust.cond.info(my.obj, plot.type = "pie", normalize.ncell = TRUE)
clust.cond.info(my.obj, plot.type = "bar")
## End(Not run)
```

clust.rm

Remove the cells that are in a cluster

Description

This function removes the cells from a designated cluster. Notice the cells will be removed from the main data (raw data would still have the original data).

Usage

```
clust.rm(x = NULL, clust.to.rm = "numeric")
```

Arguments

clust.to.rm

A data frame containing gene counts for cells. The name of the cluster to be removed.

Value

An object of class iCellR

Examples

```
## Not run:
my.obj <- clust.rm(my.obj, clust.to.rm = 5)</pre>
## End(Not run)
```

clust.stats.plot

QC on clusters (nGenes, UMIs and percent mito)

Description

This function takes an object of class iCellR and creates QC plot.

Usage

```
clust.stats.plot(x = NULL, plot.type = "box.mito",
  cell.color = "slategray3", cell.size = 1, cell.transparency = 0.5,
 box.color = "red", box.line.col = "green", back.col = "white",
 notch = F, interactive = TRUE, out.name = "plot")
```

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Arguments

	х	An object of class iCellR.
	plot.type	Choose from "box.umi", "box.mito", "box.gene", default = "box.mito".
	cell.color	Choose a color for points in the plot.
	cell.size	A number for the size of the points in the plot, default = 1 .
cell.transparency		ncy
		Color transparency for points in "scatterplot" and "boxplot", default = 0.5.
	box.color	A color for the boxes in the "boxplot", default = "red".
	box.line.col	A color for the lines around the "boxplot", default = "green".
	notch	Notch the box plots, $default = F$.
	interactive	If set to TRUE an interactive HTML file will be created, default = TRUE.
	out.name	If "interactive" is set to TRUE, the out put name for HTML, default = "plot".

Value

An object of class iCellR.

Examples

```
## Not run:
clust.stats.plot(my.obj, plot.type = "box.mito", interactive = F, out.name = "box.mito.clusters")
## End(Not run)
```

cluster.plot

Plot nGenes, UMIs and perecent mito

Description

This function takes an object of class iCellR and creates plot.

Usage

```
cluster.plot(x = NULL, cell.size = 1, plot.type = "tsne",
  cell.color = "black", back.col = "white", col.by = "clusters",
  cond.shape = F, cell.transparency = 0.5, clust.dim = 2,
  angle = 20, clonotype.max = 10, density = F, interactive = TRUE,
  static3D = F, out.name = "plot")
```

Arguments

Х	An object of class iCellR.
cell.size	A numeric value for the size of the cells, default = 1 .
plot.type	Choose between "tsne" and "pca", default = "tsne".
cell.color	Choose cell color if col.by = "monochrome", default = "black".
back.col	Choose background color, default = "black".

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col.by Choose between "clusters", "conditions", "cc" (cell cycle) or "monochrome",

default = "clusters".

cond. shape If TRUE the conditions will be shown in shapes.

cell.transparency

A numeric value between 0 to 1, default = 0.5.

clust.dim A numeric value for plot dimensions. Choose either 2 or 3, default = 2.

angle A number to rotate the non-interactive 3D plot.

density If TRUE the density plots for PCA/tSNE second dimension will be created,

default = FALSE.

interactive If TRUE an html interactive file will be made, default = TRUE.

static3D If TRUE a non-interactive 3D plot will be made.

out.name Output name for html file if interactive = TRUE, default = "plot".

Value

An object of class iCellR.

Examples

```
## Not run:
tsne.plot(my.obj)
## End(Not run)
```

data.aggregation Merge multiple data frames and add the condition names to their cell

ids

Description

This function takes data frame and merges them while also adding condition names to cell ids..

Usage

```
data.aggregation(samples = NULL, condition.names = NULL)
```

Arguments

samples A character vector of data.frame object names.

condition.names

A character vector of data.frame condition names.

Value

An object of class iCellR

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Examples

data.scale

Scale data

Description

This function takes an object of class iCellR and scales the normalized data.

Usage

```
data.scale(x = NULL)
```

Arguments

Х

An object of class iCellR.

Value

An object of class iCellR.

Examples

```
## Not run:
my.obj <- data.scale(my.obj)
## End(Not run)</pre>
```

diff.exp

Differential expression (DE) analysis

Description

This function takes an object of class iCellR and performs differential expression (DE) analysis for clusters and conditions.

Usage

```
diff.exp(x = NULL, de.by = "clusters", cond.1 = "array",
  cond.2 = "array", base.cond = 0)
```

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Arguments

x	An object of class iCellR.
de.by	Choose from "clusters", "conditions", "clustBase.condComp" or "condBase.clustComp".
cond.1	First condition to do DE analysis on.
cond.2	Second condition to do DE analysis on.
base.cond	A base condition or cluster if de.by is either cond.clust or clust.cond

Value

An object of class iCellR

Examples

```
## Not run:
diff.res <- diff.exp(my.obj, de.by = "clusters", cond.1 = c(1,4), cond.2 = c(2))
diff.res <- diff.exp(my.obj, de.by = "conditions", cond.1 = c("WT"), cond.2 = c("KO"))
diff.res <- diff.exp(my.obj, de.by = "clustBase.condComp", cond.1 = c("WT"), cond.2 = c("KO"), base.cond = 1)
diff.res <- diff.exp(my.obj, de.by = "condBase.clustComp", cond.1 = c(1), cond.2 = c(2), base.cond = "WT")
## End(Not run)</pre>
```

down.sample

Down sample conditions

Description

This function takes an object of class iCellR and down samples the condition to have equal number of cells in each condition.

Usage

```
down.sample(x = NULL)
```

Arguments

Х

An object of class iCellR.

Value

An object of class iCellR.

```
## Not run:
my.obj <- down.sample(my.obj)
## End(Not run)</pre>
```

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find.dim.	genes
-----------	-------

Find model genes from PCA data

Description

This function takes an object of class iCellR finds the model genes to run a second round of PCA.

Usage

```
find.dim.genes(x = NULL, dims = 1:10, top.pos = 15, top.neg = 5)
```

Arguments

X	An object of class iCellR.
dims	PC dimentions to be used.
top.pos	Number of top positive marker genes to be taken from each PC, default = 15.
top.neg	Number of top negative marker genes to be taken from each PC, default = 5.

Value

An object of class iCellR.

Examples

```
## Not run:
my.obj <- run.pca(my.obj, clust.method = "gene.model", gene.list = "my_model_genes.txt")
## End(Not run)</pre>
```

findMarkers

Find marker genes for each cluster

Description

This function takes an object of class iCellR and performs differential expression (DE) analysis to find marker genes for each cluster.

Usage

```
findMarkers(x = NULL, fold.change = 2, padjval = 0.1, Inf.FCs = F,
  uniq = F, positive = TRUE)
```

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Arguments

x An object of class iCellR.

fold. change A number that designates the minimum fold change for out put, default = 2.

padjval Minimum adjusted p value for out put, default = 0.1.

Inf.FCs If set to FALSE the infinite fold changes would be filtered from out put, default

= FALSE.

uniq If set to TRUE only genes that are a marker for only one cluster would be in the

out put, default = TRUE.

positive If set to FALSE both the up regulated (positive) and down regulated (negative)

markers would be in the out put, default = FALSE.

Value

An object of class iCellR

Examples

```
## Not run:
marker.genes <- findMarkers(my.obj,fold.change = 2,padjval = 0.1,uniq = T)
## End(Not run)</pre>
```

gate.to.clust

Assign cluster number to cell ids

Description

This function takes an object of class iCellR and assigns cluster number to a vector of cell ids.

Usage

```
gate.to.clust(x = NULL, my.gate = NULL, to.clust = 0)
```

Arguments

x An object of class iCellR.

my.gate A vector of cell ids.

to.clust A cluster id to be assigned to the provided cell ids.

Value

An object of class iCellR.

```
## Not run:
my.obj <- gate.to.clust(my.obj, my.gate = readLines("ids.txt"), to.clust = 1)
## End(Not run)</pre>
```

18 gene.plot

gene.plot	Make scatter, box and bar plots for genes
-----------	---

Description

This function takes an object of class iCellR and provides plots for genes.

Usage

```
gene.plot(x = NULL, gene = "NULL", cond.shape = F,
  data.type = "main", box.to.test = 0, box.pval = "sig.signs",
  plot.data.type = "tsne", scaleValue = F, min.scale = -2.5,
  max.scale = 2.5, clust.dim = 2, col.by = "clusters",
  plot.type = "scatterplot", cell.size = 1, cell.colors = c("gray",
  "red"), box.cell.col = "black", box.color = "red",
  box.line.col = "green", back.col = "white",
  cell.transparency = 0.5, interactive = TRUE, out.name = "plot")
```

Arguments

X	An object of class iCellR.
gene	A gene name to be plotted.
cond.shape	If TRUE the conditions will be shown in shapes.
box.to.test	A cluster number so that all the boxes in the box plot would be compared to. If set to "0" the cluster with the highest avrage would be choosen, default = 0 .
box.pval	Choose from "sig.values" and "sig.signs". If set to "sig.signs" p values would be replaced with signs ("na", "*", "**", "***"), default = "sig.signs".
plot.data.type	Choose from "tsne" and "pca", default = "tsne".
clust.dim	2 for 2D plots and 3 for 3D plots, default = 2.
col.by	Choose from "clusters" and "conditions", default = "clusters".
plot.type	Choose from "scatterplot", "boxplot" and "barplot", default = "scatterplot".
cell.size	A number for the size of the points in the plot, default = 1 .
cell.colors	Colors for heat mapping the points in "scatterplot", default = $c("gray","red")$.
box.cell.col	A color for the points in the box plot, default = "black".
box.color	A color for the boxes in the "boxplot", default = "red".
box.line.col	A color for the lines around the "boxplot", default = "green".
back.col	A color for the plot background, default = "black".
cell.transparer	ncy
	Color transparency for points in "scatterplot" and "boxplot", default = 0.5 .
interactive	If set to TRUE an interactive HTML file will be created, default = TRUE.
out.name	If "interactive" is set to TRUE, the out put name for HTML, default = "plot".

Value

An object of class iCellR.

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Examples

```
## Not run:
cluster.plot(my.obj,
            cell.size = 1,
            plot.type = "tsne",
            cell.color = "black",
            back.col = "white",
            col.by = "clusters",
            cell.transparency = 0.5,
            clust.dim = 3,
            interactive = T,
            density = F,
            out.name = "tSNE_3D_clusters")
cluster.plot(my.obj, cell.size = 1,
            plot.type = "tsne",
            col.by = "clusters",
            cell.transparency = 0.5,
            clust.dim = 2,
            interactive = T,
            density = F,
            out.name = "tSNE_2D_clusters")
cluster.plot(my.obj,
           cell.size = 2,
           plot.type = "tsne",
           clust.dim = 2,
           interactive = F)
cluster.plot(my.obj,
           cell.size = 1,
           plot.type = "tsne",
           col.by = "clusters",
           clust.dim = 3,
           interactive = F,
           angle = 45)
cluster.plot(my.obj,
          cell.size = 1,
          plot.type = "pca",
          cell.color = "black",
          back.col = "white",
          col.by = "conditions";
          cell.transparency = 0.5,
          clust.dim = 3,
          interactive = T,
          density = F,
          out.name = "PCA_3D_conditions")
## End(Not run)
```

gene.stats Mai

Make statistical information for each gene across all the cells (SD, mean, expression, etc.)

20 gg.cor

Description

This function takes an object of class iCellR and provides some statistical information for the genes.

Usage

```
gene.stats(x = NULL, which.data = "raw.data", each.cond = F)
```

Arguments

x An object of class iCellR.

which.data Choose from "raw.data" or "main.data", default = "raw.data".

each.cond If TRUE each condition will be calculated, default = F.

Value

An object of class iCellR.

Examples

```
## Not run:
my.obj <- gene.stats(my.obj, which.data = "main.data")
## End(Not run)</pre>
```

gg.cor

Normalize ADT data. This function takes data frame and Normalizes ADT data.

Description

Normalize ADT data. This function takes data frame and Normalizes ADT data.

Usage

```
gg.cor(x = NULL, data.type = "imputed", gene1 = NULL, gene2 = NULL,
conds = NULL, cell.size = 1, cell.transparency = 0.5,
interactive = TRUE, out.name = "plot")
```

Arguments

out.name

x An object of class iCellR.

data.type Choose from imputed and main, default = "imputed".

gene1 First gene name.

gene2 Second gene name.

conds Filter only one condition (only one), default is all conditions.

cell.size A numeric value for the size of the cells, default = 1.

cell.transparency
A numeric value between 0 to 1, default = 0.5.

interactive If TRUE an html interactive file will be made, default = TRUE.

Output name for html file if interactive = TRUE, default = "plot".

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Value

An object of class iCellR

Examples

```
## Not run:
gg.cor(my.obj, interactive = F, gene1 = "NKG7",gene2 = "GNLY", conds=c("WT"))
## End(Not run)
```

heatmap.gg.plot

Create heatmaps for genes in clusters or conditions.

Description

This function takes an object of class iCellR and genes and provides a heatmap.

Usage

```
heatmap.gg.plot(x = NULL, gene = "NULL", data.type = "main",
  cluster.by = "clusters", min.scale = -2.5, max.scale = 2.5,
  interactive = T, cex.col = 10, cex.row = 10, no.key = FALSE,
  out.name = "plot", heat.colors = c("blue", "white", "red"))
```

Arguments

x A data frame containing gene counts for cells.

gene A set of gene names to be heatmapped.

cluster.by Choose from "clusters" or "conditions", default = "clusters".

interactive If TRUE an html interactive file will be made, default = TRUE.

no.key If you want a color legend key, default = FALSE.

 $\label{eq:outname} \text{Output name for html file if interactive} = TRUE, \ default = "plot".$

 $\label{eq:colors} \textbf{Colors for heatmap, default} = \textbf{c}("blue" , "white", "red").$

Value

An object of class iCellR

```
## Not run:
heatmap.gg.plot(my.obj, gene = MyGenes, interactive = T, out.name = "plot", cluster.by = "clusters")
## End(Not run)
```

22 make.gene.model

load10x

Load 10X data as data.frame

Description

This function takes 10X data files barcodes.tsv, genes.tsv and matrix.mtx and converts them to proper matrix file for iCellR.

Usage

```
load10x(dir.10x = NULL, gene.name = 2)
```

Arguments

```
\mbox{dir.10x} \qquad \mbox{A directory that includes the $10$X barcodes.tsv, genes.tsv and matrix.mtx files.} \\ \mbox{gene.name.column}
```

Gene names or ids column number, default = 2.

Value

The data frame object

Examples

```
## Not run:
load10x("/hg19", gene.name = "geneSymbol")
## End(Not run)
```

make.gene.model

Make a gene model for clustering

Description

This function takes an object of class iCellR and provides a gene list for clustering based on the parameters set in the model.

Usage

```
make.gene.model(x = NULL, dispersion.limit = 1.5,
  base.mean.rank = 500, non.sig.col = "darkgray",
  right.sig.col = "chartreuse3", left.sig.col = "cadetblue3",
  disp.line.col = "black", rank.line.col = "red", cell.size = 1.75,
  cell.transparency = 0.5, no.mito.model = T, no.cell.cycle = T,
  mark.mito = T, interactive = TRUE, out.name = "plot")
```

make.gene.model 23

Arguments

```
An object of class iCellR.
dispersion.limit
                   A number for taking the genes that have dispersion above this number, default
base.mean.rank A number taking the top genes ranked by base mean, default = 500.
                  Color for the genes not used for the model, default = "darkgray".
non.sig.col
right.sig.col
                  Color for the genes above the dispersion limit, default = "chartreuse3".
left.sig.col
                  Color for the genes above the rank limit, default = "cadetblue3".
                  Color of the line for dispersion limit, default = "black".
disp.line.col
rank.line.col
                  Color of the line for rank limit, default = "red".
                  A number for the size of the points in the plot, default = 1.75.
cell.size
cell.transparency
                  Color transparency for the points in the plot, default = 0.5.
                  If set to TRUE, mitochondrial genes would be excluded from the gene list made
no.mito.model
                  for clustering, default = TRUE.
                  Mark mitochondrial genes in the plot, default = TRUE.
mark.mito
                  If set to TRUE an interactive HTML file will be created, default = TRUE.
interactive
out.name
                  If "interactive" is set to TRUE, the out put name for HTML, default = "plot".
```

Value

An object of class iCellR.

```
## Not run:
make.gene.model(my.obj,
               dispersion.limit = 1.5,
               base.mean.rank = 500,
               no.mito.model = T,
               mark.mito = T,
               interactive = T,
               out.name = "gene.model")
make.gene.model(my.obj,
             dispersion.limit = 1.5,
             base.mean.rank = 500,
             no.mito.model = T,
             mark.mito = T,
             interactive = F
             out.name = "gene.model")
## End(Not run)
```

24 norm.adt

make.obj

Create an object of class iCellR.

Description

This function takes data frame and makes an object of class iCellR.

Usage

```
make.obj(x = NULL)
```

Arguments

Χ

A data frame containing gene counts for cells.

Value

An object of class iCellR

Examples

```
## Not run:
my.obj <- make.obj(my.data)
## End(Not run)</pre>
```

norm.adt

Normalize ADT data. This function takes data frame and Normalizes ADT data.

Description

Normalize ADT data. This function takes data frame and Normalizes ADT data.

Usage

```
norm.adt(x = NULL)
```

Arguments

x

An object of class iCellR.

Value

An object of class iCellR

norm.data 25

Examples

```
## Not run:
my.obj <- norm.adt(my.obj)
## End(Not run)</pre>
```

norm.data

Normalize data

Description

This function takes an object of class iCellR and normalized the data based on "global.glsf", "ranked.glsf" or "spike.in" methods.

Usage

```
norm.data(x = NULL, norm.method = "ranked.glsf", top.rank = 500,
    spike.in.factors = NULL, rpm.factor = 1000)
```

Arguments

An object of class iCellR.

Norm.method Choose a normalization method, there are three option currently. Choose from "global.glsf", "ranked.glsf", "ranked.deseq", "deseq", "rpm", "spike.in" or no.norm, default = "ranked.glsf".

top.rank If the method is set to "ranked.glsf", you need to set top number of genes sorted based on global base mean, default = 500.

spike.in.factors

A numeric vector of spike-in values with the same cell id order as the main data.

rpm.factor If the norm.method is set to "rpm" the library sizes would be diveded by this number, default = 1000 (higher numbers recomanded for bulk RNA-Seq).

Value

An object of class iCellR.

26 prep.vdj

opt.pcs.plot

Find optimal number of PCs for clustering

Description

This function takes an object of class iCelIR and finds optimal number of PCs for clustering.

Usage

```
opt.pcs.plot(x = NULL, pcs.in.plot = 50)
```

Arguments

x An object of class iCellR.

pcs.in.plot Number of PCs to show in plot, defult = 50.

Value

An object of class iCellR.

Examples

```
## Not run:
find.opt.pcs(my.obj)
## End(Not run)
```

prep.vdj

Add CITE-seq antibody-derived tags (ADT)

Description

This function takes a data frame of ADT values per cell and adds it to the iCellR object.

Usage

```
prep.vdj(vdj.data = "all_contig_annotations.csv", cond.name = "NULL")
```

Arguments

x An object of class iCellR.

adt.data A data frame containing ADT counts for cells.

Value

An object of class iCellR

pseudotime 27

Examples

```
## Not run:
my.obj <- add.adt(my.obj, adt.data = adt.data)
## End(Not run)</pre>
```

pseudotime

Pseudotime

Description

This function takes an object of class iCellR and marker genes for clusters and performs pseudotime analysis.

Usage

```
pseudotime(x = NULL, marker.genes = "NULL", dim = 1:10)
```

Arguments

```
x An object of class iCellR.marker.genes A list of marker genes for clusters.
```

Value

An object of class iCellR.

Examples

```
## Not run:
my.obj <- pseudotime(my.obj, marker.genes = MyGenes, dim = 1:10)
## End(Not run)</pre>
```

pseudotime.tree

Pseudotime Tree

Description

This function takes an object of class iCellR and marker genes for clusters and performs pseudotime for differentiation or time course analysis.

Usage

```
pseudotime.tree(x = NULL, marker.genes = "NULL",
   clust.names = "NULL", dist.method = "euclidean",
   clust.method = "complete", label.offset = 0.5, type = "classic",
   hang = 1, cex = 1)
```

28 qc.stats

Arguments

x	An object of class iCellR.
marker.genes	A list of marker genes for clusters.
clust.names	A list of names for clusters.
dist.method	Choose from "euclidean", "maximum", "manhattan", "canberra", "binary" or "minkowski", default = "euclidean".
clust.method	Choose from "ward.D", "ward.D2", "single", "complete", "average", "mcquitty", "median" or "centroid", default = "complete".
label.offset	Space between names and tree, default = 0.5 .
type	Choose from "classic", "jitter", "unrooted", "fan", "cladogram", "radial", default = "classic".
cex	Text size, $default = 1$.

Value

An object of class iCellR.

Examples

```
## Not run:
my.obj <- run.pca(my.obj, clust.method = "gene.model", gene.list = "my_model_genes.txt")
## End(Not run)

qc.stats

Calculate the number of UMIs and genes per cell and percentage of mitochondrial genes per cell and cell cycle genes.</pre>
```

Description

This function takes data frame and calculates the number of UMIs, genes per cell and percentage of mitochondrial genes per cell and cell cycle genes.

Usage

```
qc.stats(x = NULL, which.data = "raw.data", mito.genes = "default",
    s.phase.genes = s.phase, g2m.phase.genes = g2m.phase)
```

Arguments

Χ	A data frame containing gene counts for cells.	
which.data	Choose from raw data or main data, default = "raw.data".	
mito.genes	A character vector of mitochondrial genes names , default is the genes starting with mt .	
s.phase.genes	A character vector of gene names for S phase, default = s.phase.	
g2m.phase.genes		
	A character vector of gene names for G2 and M phase, default = $g2m.phase$.	

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Value

The data frame object

Examples

```
## Not run:
UMIs.genes.mit(my.data)
## End(Not run)
```

run.cca

Run PCA on the main data

Description

This function takes an object of class iCellR and runs PCA on the main data.

Usage

```
run.cca(x = NULL, top.vari.genes = 1000, cc.number = 30,
  dims.align = 1:20, normalize.data = TRUE, scale.data = TRUE,
  normalization.method = "LogNormalize", scale.factor = 10000,
  display.progress = TRUE)
```

Arguments

x An object of class iCellR.

clust.method Choose from "base.mean.rank" or "gene.model", default is "base.mean.rank".

top.rank A number taking the top genes ranked by base mean, default = 500.

plus.log.value A number to add to each value in the matrix before log transformasion to aviond Inf numbers, default = 0.1.

gene.list A list of genes to be used for PCA. If "clust.method" is set to "gene.model", default = "my_model_genes.txt".

Value

An object of class iCellR.

```
## Not run:
my.obj <- run.pca(my.obj, clust.method = "gene.model", gene.list = "my_model_genes.txt")
## End(Not run)</pre>
```

30 run.clustering

run.clustering

Clustering the data

Description

This function takes an object of class iCellR and finds optimal number of clusters and clusters the data.

Usage

```
run.clustering(x = NULL, clust.method = "kmeans",
  dist.method = "euclidean", index.method = "silhouette",
  max.clust = 25, min.clust = 2, dims = 1:10)
```

Arguments

x An object of class iCellR.

clust.method the cluster analysis method to be used. This should be one of: "ward.D",

"ward.D2", "single", "complete", "average", "mcquitty", "median", "centroid",

'kmeans".

dist.method the distance measure to be used to compute the dissimilarity matrix. This must

be one of: "euclidean", "maximum", "manhattan", "canberra", "binary", "minkowski" or "NULL". By default, distance="euclidean". If the distance is "NULL", the dissimilarity matrix (diss) should be given by the user. If distance is not

"NULL", the dissimilarity matrix should be "NULL".

index.method the index to be calculated. This should be one of: "kl", "ch", "hartigan", "ccc",

"scott", "marriot", "trcovw", "tracew", "friedman", "rubin", "cindex", "db", "silhouette", "duda", "pseudot2", "beale", "ratkowsky", "ball", "ptbiserial", "gap", "frey", "mcclain", "gamma", "gplus", "tau", "dunn", "hubert", "sdindex", "dindex", "sdbw", "all" (all indices except GAP, Gamma, Gplus and Tau), "alllong"

(all indices with Gap, Gamma, Gplus and Tau included).

max.clust maximal number of clusters, between 2 and (number of objects - 1), greater or

equal to min.nc.

min.clust minimum number of clusters, default = 2.

dims PCA dimentions to be use for clustering, default = 1:10.

Value

An object of class iCellR.

run.diffusion.map 31

run.diffusion.map	Run diffusion map on PCA data (PHATE - Potential of Heat-Diffusion
	for Affinity-Based Transition Embedding)

Description

This function takes an object of class iCellR and runs diffusion map on PCA data.

Usage

```
run.diffusion.map(x = NULL, dims = 1:10, method = "phate",
 ndim = 3, k = 5, alpha = 40, n.landmark = 2000, gamma = 1,
 t = "auto", knn.dist.method = "euclidean", init = NULL,
 mds.method = "metric", mds.dist.method = "euclidean", t.max = 100,
 npca = 100, plot.optimal.t = FALSE, verbose = 1, n.jobs = 1,
 seed = NULL, potential.method = NULL, use.alpha = NULL,
 n.svd = NULL, pca.method = NULL, g.kernel = NULL, diff.op = NULL,
 landmark.transitions = NULL, diff.op.t = NULL, dist.method = NULL)
```

Arguments

x	An object of class iCellR.
dims	PC dimentions to be used for UMAP analysis.
ndim	int, optional, default: 2 number of dimensions in which the data will be embedded
k	int, optional, default: 5 number of nearest neighbors on which to build kernel
alpha	int, optional, default: $40\ \mathrm{sets}$ decay rate of kernel tails. If NULL, alpha decaying kernel is not used
n.landmark	int, optional, default: 2000 number of landmarks to use in fast PHATE
gamma	float, optional, default: 1 Informational distance constant between -1 and 1. gamma=1 gives the PHATE log potential, gamma=0 gives a square root potential.
t	int, optional, default: 'auto' power to which the diffusion operator is powered sets the level of diffusion

knn.dist.method

string, optional, default: 'euclidean'. recommended values: 'euclidean', 'cosine', 'precomputed' Any metric from scipy.spatial.distance can be used distance metric for building kNN graph. If 'precomputed', data should be an n_samples x n_samples distance or affinity matrix. Distance matrices are assumed to have zeros down the diagonal, while affinity matrices are assumed to have non-zero values down the diagonal. This is detected automatically using data[0,0]. You can override this detection with knn.dist.method='precomputed_distance'

or knn.dist.method='precomputed_affinity'.

init phate object, optional object to use for initialization. Avoids recomputing inter-

mediate steps if parameters are the same.

string, optional, default: 'metric' choose from 'classic', 'metric', and 'nonmds.method

metric' which MDS algorithm is used for dimensionality reduction

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mds.dist.method

string, optional, default: 'euclidean' recommended values: 'euclidean' and 'co-

sine'

t.max int, optional, default: 100. Maximum value of t to test for automatic t selection.

npca int, optional, default: 100 Number of principal components to use for calcu-

lating neighborhoods. For extremely large datasets, using $n_pca < 20$ allows

neighborhoods to be calculated in log(n_samples) time.

plot.optimal.t boolean, optional, default: FALSE If TRUE, produce a plot showing the Von

Neumann Entropy curve for automatic t selection.

verbose int or boolean, optional (default : 1) If TRUE or > 0, print verbose updates.

n. jobs int, optional (default: 1) The number of jobs to use for the computation. If -1 all

CPUs are used. If 1 is given, no parallel computing code is used at all, which is useful for debugging. For n_j obs below -1, (n.cpus + 1 + n.jobs) are used. Thus

for $n_{jobs} = -2$, all CPUs but one are used

seed int or NULL, random state (default: NULL)

potential.method

Deprecated. For log potential, use gamma=1. For sqrt potential, use gamma=0.

use.alpha Deprecated To disable alpha decay, use alpha=NULL

n.svd Deprecated.
pca.method Deprecated.
g.kernel Deprecated.
diff.op Deprecated.

landmark.transitions

Deprecated.

diff.op.t Deprecated.
dist.method Deprecated.

Value

An object of class iCellR.

Examples

```
## Not run:
my.obj <- run.diffusion.map(my.obj, dims = 1:10, method = "phate")
## End(Not run)</pre>
```

run.impute

Run MAGIC on Main Data.

Description

This function takes an object of class iCellR and runs MAGIC on main data. Markov Affinity-based Graph Imputation of Cells (MAGIC) is an algorithm for denoising and transcript recover of single cells applied to single-cell RNA sequencing data, as described in van Dijk et al, 2018.

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Usage

```
run.impute(x = NULL, genes = "all_genes", k = 10, alpha = 15,
    t = "auto", npca = 100, init = NULL, t.max = 20,
    knn.dist.method = "euclidean", verbose = 1, n.jobs = 1,
    seed = NULL)
```

Arguments

x	An object of class iCellR.	
genes	character or integer vector, default: NULL vector of column names or column indices for which to return smoothed data If 'all_genes' or NULL, the entire smoothed matrix is returned	
k	int, optional, default: 10 number of nearest neighbors on which to build kernel	
alpha	int, optional, default: 15 sets decay rate of kernel tails. If NULL, alpha decaying kernel is not used	
t	int, optional, default: 'auto' power to which the diffusion operator is powered sets the level of diffusion. If 'auto', t is selected according to the Procrustes disparity of the diffused data.'	
npca	number of PCA components that should be used; default: 100.	
init	magic object, optional object to use for initialization. Avoids recomputing intermediate steps if parameters are the same.	
t.max	int, optional, default: 20 Maximum value of t to test for automatic t selection.	
knn.dist.method		
	string, optional, default: 'euclidean'. recommended values: 'euclidean', 'co-sine' Any metric from 'scipy.spatial.distance' can be used distance metric for building kNN graph.	
verbose	'int' or 'boolean', optional (default : 1) If 'TRUE' or '> 0', print verbose updates.	
n.jobs	'int', optional (default: 1) The number of jobs to use for the computation. If -1 all CPUs are used. If 1 is given, no parallel computing code is used at all, which is useful for debugging. For n_jobs below -1, (n.cpus + 1 + n.jobs) are used. Thus for n_jobs = -2, all CPUs but one are used	
seed	int or 'NULL', random state (default: 'NULL')	

Value

An object of class iCellR.

```
## Not run:
my.obj <- run.impute(my.obj)
## End(Not run)</pre>
```

34 run.pc.tsne

run.pc.tsne	Run tSNE on PCA Data. Barnes-Hut implementation of t-Distributed
	Stochastic Neighbor Embedding

Description

This function takes an object of class iCellR and runs tSNE on PCA data. Wrapper for the C++ implementation of Barnes-Hut t-Distributed Stochastic Neighbor Embedding. t-SNE is a method for constructing a low dimensional embedding of high-dimensional data, distances or similarities. Exact t-SNE can be computed by setting theta=0.0.

Usage

```
run.pc.tsne(x = NULL, dims = 1:10, my.seed = 0, initial_dims = 50,
  perplexity = 30, theta = 0.5, check_duplicates = FALSE,
  pca = TRUE, max_iter = 1000, verbose = FALSE,
  is_distance = FALSE, Y_init = NULL, pca_center = TRUE,
  pca_scale = FALSE, stop_lying_iter = ifelse(is.null(Y_init), 250L,
  0L), mom_switch_iter = ifelse(is.null(Y_init), 250L, 0L),
  momentum = 0.5, final_momentum = 0.8, eta = 200,
  exaggeration_factor = 12)
```

Arguments

pca_center
pca_scale

An object of class iCellR. х dims PC dimentions to be used for tSNE analysis. initial_dims integer; the number of dimensions that should be retained in the initial PCA step (default: 50) perplexity numeric; Perplexity parameter numeric; Speed/accuracy trade-off (increase for less accuracy), set to 0.0 for theta exact TSNE (default: 0.5) check_duplicates logical; Checks whether duplicates are present. It is best to make sure there are no duplicates present and set this option to FALSE, especially for large datasets (default: TRUE) logical; Whether an initial PCA step should be performed (default: TRUE) рса integer; Number of iterations (default: 1000) max_iter logical; Whether progress updates should be printed (default: FALSE) verbose is_distance logical; Indicate whether X is a distance matrix (experimental, default: FALSE) Y_init matrix; Initial locations of the objects. If NULL, random initialization will be used (default: NULL). Note that when using this, the initial stage with exagger-

stop_lying_iter integer; Iteration after which the perplexities are no longer exaggerated (default: 250, except when Y_init is used, then 0)

ated perplexity values and a larger momentum term will be skipped. logical; Should data be centered before pca is applied? (default: TRUE)

logical; Should data be scaled before pca is applied? (default: FALSE)

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```
mom_switch_iter
```

integer; Iteration after which the final momentum is used (default: 250, except

when Y_init is used, then 0)

momentum numeric; Momentum used in the first part of the optimization (default: 0.5) final_momentum numeric; Momentum used in the final part of the optimization (default: 0.8)

eta numeric; Learning rate (default: 200.0)

exaggeration_factor

numeric; Exaggeration factor used to multiply the P matrix in the first part of

the optimization (default: 12.0)

Value

An object of class iCellR.

Examples

```
## Not run:
my.obj <- run.pc.tsne(my.obj, dims = 1:10)
## End(Not run)</pre>
```

run.pca

Run PCA on the main data

Description

This function takes an object of class iCellR and runs PCA on the main data.

Usage

```
run.pca(x = NULL, data.type = "main", method = "base.mean.rank",
  top.rank = 500, plus.log.value = 0.1, batch.norm = F,
  gene.list = "character")
```

Arguments

x An object of class iCellR.

method Choose from "base.mean.rank" or "gene.model", default is "base.mean.rank". If

gene.model is chosen you need to provide gene.list.

top.rank A number taking the top genes ranked by base mean, default = 500.

plus.log.value A number to add to each value in the matrix before log transformasion to aviond

Inf numbers, default = 0.1.

batch.norm If TRUE the data will be normalized based on the genes in gene.list or top ranked

genes.

gene.list A charactor vector of genes to be used for PCA. If "clust.method" is set to

"gene.model", default = "my_model_genes.txt".

Value

An object of class iCellR.

run.tsne

Examples

```
## Not run:
my.obj <- run.pca(my.obj, clust.method = "gene.model", gene.list = "my_model_genes.txt")
## End(Not run)

run.tsne

Run tSNE on the Main Data. Barnes-Hut implementation of t-
Distributed Stochastic Neighbor Embedding</pre>
```

Description

This function takes an object of class iCellR and runs tSNE on PCA data. Wrapper for the C++ implementation of Barnes-Hut t-Distributed Stochastic Neighbor Embedding. t-SNE is a method for constructing a low dimensional embedding of high-dimensional data, distances or similarities. Exact t-SNE can be computed by setting theta=0.0.

Usage

```
run.tsne(x = NULL, clust.method = "base.mean.rank", top.rank = 500,
  gene.list = "character", initial_dims = 50, perplexity = 30,
  theta = 0.5, check_duplicates = TRUE, pca = TRUE,
  max_iter = 1000, verbose = FALSE, is_distance = FALSE,
  Y_init = NULL, pca_center = TRUE, pca_scale = FALSE,
  stop_lying_iter = ifelse(is.null(Y_init), 250L, 0L),
  mom_switch_iter = ifelse(is.null(Y_init), 250L, 0L), momentum = 0.5,
  final_momentum = 0.8, eta = 200, exaggeration_factor = 12)
```

Arguments

is_distance

_		
X	An object of class iCellR.	
clust.method	Choose from "base.mean.rank" or "gene.model", defult is "base.mean.rank".	
top.rank	A number taking the top genes ranked by base mean, defult = 500 .	
gene.list	A list of genes to be used for tSNE analysis. If "clust.method" is set to "gene.model" defult = " $my_model_genes.txt$ ".	
initial_dims	integer; the number of dimensions that should be retained in the initial PCA step (default: 50)	
perplexity	numeric; Perplexity parameter	
theta	numeric; Speed/accuracy trade-off (increase for less accuracy), set to 0.0 for exact TSNE (default: 0.5)	
check_duplicates		
	logical; Checks whether duplicates are present. It is best to make sure there are no duplicates present and set this option to FALSE, especially for large datasets (default: TRUE)	
рса	logical; Whether an initial PCA step should be performed (default: TRUE)	
max_iter	integer; Number of iterations (default: 1000)	
verbose	logical; Whether progress updates should be printed (default: FALSE)	

logical; Indicate whether X is a distance matrix (experimental, default: FALSE)

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Y_init matrix; Initial locations of the objects. If NULL, random initialization will be

used (default: NULL). Note that when using this, the initial stage with exagger-

ated perplexity values and a larger momentum term will be skipped.

pca_center logical; Should data be centered before pca is applied? (default: TRUE)

pca_scale logical; Should data be scaled before pca is applied? (default: FALSE)

stop_lying_iter

integer; Iteration after which the perplexities are no longer exaggerated (default:

250, except when Y_init is used, then 0)

mom_switch_iter

integer; Iteration after which the final momentum is used (default: 250, except

when Y_init is used, then 0)

momentum numeric; Momentum used in the first part of the optimization (default: 0.5)

final_momentum numeric; Momentum used in the final part of the optimization (default: 0.8)

eta numeric; Learning rate (default: 200.0)

exaggeration_factor

numeric; Exaggeration factor used to multiply the P matrix in the first part of

the optimization (default: 12.0)

Value

An object of class iCellR.

Examples

```
## Not run:
my.obj <- run.tsne(my.obj, clust.method = "gene.model", gene.list = "my_model_genes.txt")
## End(Not run)</pre>
```

run.umap

Run UMAP on PCA Data (Computes a manifold approximation and projection)

Description

This function takes an object of class iCellR and runs UMAP on PCA data.

Usage

```
run.umap(x = NULL, dims = 1:10, method = "naive")
```

Arguments

x An object of class iCellR.

dims PC dimentions to be used for UMAP analysis.

method Character, implementation. Available methods are 'naive' (an implementation

written in pure R) and 'umap-learn' (requires python package 'umap-learn').

Choose from "naive" and "umap-learn", default = "naive".

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Value

An object of class iCellR.

Examples

```
## Not run:
my.obj <- run.umap(my.obj, dims = 1:10)
## End(Not run)</pre>
```

stats.plot

Plot nGenes, UMIs and percent mito

Description

This function takes an object of class iCellR and creates QC plot.

Usage

```
stats.plot(x = NULL, plot.type = "box.umi",
  cell.color = "slategray3", cell.size = 1, cell.transparency = 0.5,
  box.color = "red", box.line.col = "green", back.col = "white",
  interactive = TRUE, out.name = "plot")
```

Arguments

```
An object of class iCellR.
Х
plot.type
                   Choose from "box.umi", "box.mito", "box.gene", "box.s.phase", "box.g2m.phase", "all.in.one",
                   "point.mito.umi", "point.gene.umi".
cell.color
                   Choose a color for points in the plot.
                   A number for the size of the points in the plot, default = 1.
cell.size
cell.transparency
                   Color transparency for points in "scatterplot" and "boxplot", default = 0.5.
                   A color for the boxes in the "boxplot", default = "red".
box.color
box.line.col
                   A color for the lines around the "boxplot", default = "green".
                   If set to TRUE an interactive HTML file will be created, default = TRUE.
interactive
out.name
                   If "interactive" is set to TRUE, the out put name for HTML, default = "plot".
```

Value

An object of class iCellR.

top.markers 39

Examples

top.markers

Choose top marker genes

Description

This function takes the marker genes info if chooses marker gene names for plots.

Usage

```
top.markers(x = NULL, topde = 10, min.base.mean = 0.2,
  filt.ambig = T, cluster = 0)
```

Arguments

x An object of class iCellR.

topde Number of top differentially expressed genes to be choosen from each cluster,

default = 10.

min.base.mean Minimum base mean of the genes to be chosen, default = 0.5.

Value

A set of gene names

```
## Not run:
MyGenes <- top.markers(marker.genes, topde = 10, min.base.mean = 0.8)
## End(Not run)</pre>
```

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vdj.stats

Add CITE-seq antibody-derived tags (ADT)

Description

This function takes a data frame of ADT values per cell and adds it to the iCellR object.

Usage

```
vdj.stats(vdj.data = "VDJ_analysis_ready.tsv")
```

Arguments

x An object of class iCellR.adt.data A data frame containing ADT counts for cells.

Value

An object of class iCellR

Examples

```
## Not run:
my.obj <- add.adt(my.obj, adt.data = adt.data)
## End(Not run)</pre>
```

volcano.ma.plot

Create MA and Volcano plots.

Description

This function takes the result of differential expression (DE) analysis and provides MA and volcano plots.

Usage

```
volcano.ma.plot(x = NULL, sig.value = "padj", sig.line = 0.1,
  plot.type = "volcano", x.limit = 2, y.limit = 2, limit.force = F,
  scale.ax = T, dot.size = 1.75, dot.transparency = 0.5,
  dot.col = c("#E64B35", "#3182bd", "#636363"), interactive = TRUE,
  out.name = "plot")
```

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Arguments

X	A data frame containing differential expression (DE) analysis results.
sig.value	Choose from "pval" or "padj", default = "padj".
sig.line	A number to draw the line for the significant genes based on sig.value type, $default = 0.1$.
plot.type	Choose from "ma" or "volcano", default = "volcano".
x.limit	A number to set a limit for the x axis.
y.limit	A number to set a limit for the y axis.
limit.force	If set to TRUE the x.limit and y.limit will be forced, default = FALSE.
scale.ax	If set to TRUE the y axis will be scaled to include all the points, default = TRUE.
dot.size	A number for the size of the points in the plot, default = 1.75 .
dot.transparen	су
	Color transparency for points in "scatterplot" and "boxplot", default = 0.5.
dot.col	A set of three colors for the points in the volcano plot, default = $c("#E64B35","#3182bd","#636363")$.
interactive	If set to TRUE an interactive HTML file will be created, default = TRUE.
out.name	If "interactive" is set to TRUE, the out put name for HTML, default = "plot".

Value

Plots

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