

Package ‘scCATCH’

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Type Package

Title Single Cell Cluster-Based Annotation Toolkit for Cellular Heterogeneity

Version 3.2.2

Depends R (>= 4.0.0)

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Description An automatic cluster-based annotation pipeline based on evidence-based score by matching the marker genes with known cell markers in tissue-specific cell taxonomy reference database for single-cell RNA-seq data. See Shao X, et al (2020) <[doi:10.1016/j.isci.2020.100882](https://doi.org/10.1016/j.isci.2020.100882)> for more details.

URL <https://github.com/ZJUFanLab/scCATCH>

License GPL (>= 3)

Encoding UTF-8

LazyData true

RoxygenNote 7.1.2

Suggests rmarkdown, knitr, testthat, prettydoc

VignetteBuilder knitr

Imports Matrix, methods, progress, stats, reshape2

NeedsCompilation no

Repository CRAN

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cellmatch	<i>cellmatch</i>
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Description

Marker genes of 'Human' and 'Mouse'.

Usage

```
cellmatch
```

Format

An object of class `data.frame` with 49560 rows and 11 columns.

Source

<https://github.com/ZJUFanLab/scCATCH/tree/master/data>

createscCATCH	<i>scCATCH object</i>
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Description

create scCATCH object using single-cell count data and cluster information.

Usage

```
createscCATCH(data, cluster)
```

Arguments

- | | |
|----------------------|--|
| <code>data</code> | A matrix or dgCMatrix containing normalized single-cell RNA-seq data, each column representing a cell, each row representing a gene. See demo_data . |
| <code>cluster</code> | A character containing the cluster information for each cell. The length of it must be equal to the ncol of the data. |

Value

scCATCH object

demo_data	<i>Demo data of single-cell RNA-seq data</i>
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Description

Demo data of single-cell RNA-seq data

Usage

```
demo_data()
```

Details

data used in [createscCATCH](#) must be a matrix object, each column representing a cell, each row representing a gene.

Value

A demo data matrix.

Examples

```
data_demo <- demo_data()
```

demo_geneinfo	<i>Demo data of geneinfo</i>
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Description

Demo data of geneinfo

Usage

```
demo_geneinfo()
```

Details

geneinfo used in [rev_gene](#) must be a data.frame object with three columns, namely 'symbol', 'synonyms', 'species'.

Value

A demo geneinfo data.frame.

Examples

```
geneinfo_demo <- demo_geneinfo()
```

demo_marker	<i>Demo data of markers</i>
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Description

Demo data of markers

Usage

```
demo_marker()
```

Details

markers used in [findmarkergene](#) must be a `data.frame` object with eleven columns.

Value

A demo marker `data.frame`.

Examples

```
markers_demo <- demo_marker()
```

findcelltype	<i>Evidence-based score and annotation for each cluster</i>
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Description

Evidence-based score and annotation for each cluster.

Usage

```
findcelltype(object, verbose = TRUE)
```

Arguments

- | | |
|---------|--|
| object | scCATCH object generated from findmarkergene . |
| verbose | Show progress messages. |

Value

scCATCH object containing the results of predicted cell types for each cluster.

<code>findmarkergene</code>	<i>Find potential marker genes for each cluster</i>
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Description

Identify potential marker genes for each cluster.

Usage

```
findmarkergene(  
  object,  
  species = NULL,  
  cluster = "All",  
  if_use_custom_marker = FALSE,  
  marker = NULL,  
  cancer = "Normal",  
  tissue = NULL,  
  use_method = "1",  
  comp_cluster = NULL,  
  cell_min_pct = 0.25,  
  logfc = 0.25,  
  pvalue = 0.05,  
  verbose = TRUE  
)
```

Arguments

<code>object</code>	scCATCH object generated from createscCATCH .
<code>species</code>	The specie of cells. The species must be defined. 'Human' or 'Mouse'. When <code>if_use_custom_marker</code> is set TRUE, no need to define the species.
<code>cluster</code>	Select which clusters for potential marker genes identification. e.g. '1', '2', etc. The default is 'All' to find potential makrer genes for each cluster.
<code>if_use_custom_marker</code>	Whether to use custom markers data.frame.
<code>marker</code>	A data.frame containing marker genes. See demo_marker . Default is to use the system <code>cellmatch</code> data.frame.
<code>cancer</code>	If the sample is from cancer tissue, then the cancer type may be defined. When <code>if_use_custom_marker</code> is set TRUE, no need to define the cancer.
<code>tissue</code>	Tissue origin of cells must be defined. Select one or more related tissue types. When <code>if_use_custom_marker</code> is set TRUE, no need to define the tissue.
<code>use_method</code>	'1' is to compare with other every cluster. '2' is to compare with other clusters together.
<code>comp_cluster</code>	Number of clusters to compare. Default is to compare all other cluster for each cluster. Set it between 1 and length of unique clusters. More marker genes will be obtained for smaller <code>comp_cluster</code> .

<code>cell_min_pct</code>	Include the gene detected in at least this many cells in each cluster.
<code>logfc</code>	Include the gene with at least this fold change of average gene expression compared to every other clusters.
<code>pvalue</code>	Include the significantly highly expressed gene with this cutoff of p value from wilcox test compared to every other clusters.
<code>verbose</code>	Show progress messages.

Details

Details of available tissues see <https://github.com/ZJUFanLab/scCATCH/wiki>

Value

scCATCH object

`geneinfo`

geneinfo

Description

Gene symbols of 'Human' and 'Mouse' updated on Jan. 2, 2022 for revising genes.

Usage

`geneinfo`

Format

An object of class `data.frame` with 240502 rows and 3 columns.

Source

<https://www.ncbi.nlm.nih.gov/gene>

rev_gene	<i>Pre-processing step: revising gene symbols</i>
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Description

Revise genes according to NCBI Gene symbols updated in June 19, 2022 for count matrix, user-custom cell marker data.frame.

Usage

```
rev_gene(data = NULL, data_type = NULL, species = NULL, geneinfo = NULL)
```

Arguments

data	A matrix or dgCMatrix containing count or normalized data, each column representing a spot or a cell, each row representing a gene; Or a data.frame containing cell markers, use demo_marker .
data_type	A character to define the type of data, select 'data' for the data matrix, 'marker' for the data.frame containing cell markers.
species	Species of the data.'Human' or 'Mouse'.
geneinfo	A data.frame of the system data containing gene symbols of 'Human' and 'Mouse' updated on Jan. 1, 2022.

Value

A new matrix or data.frame.

scCATCH	<i>Definition of 'scCATCH' class</i>
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Description

An S4 class containing the data, meta, and results of inferred cell types.

Slots

data	A list containing normalized data. See demo_data .
meta	A data frame containing the meta data.
para	A list containing the parameters.
markergene	A data frame containing the identified markers for each cluster.
celltype	A data frame containing the cell types for each cluster.
marker	A data frame containing the known markers. See demo_marker .

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