

# Package ‘scCAN’

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**Type** Package

**Title** Single-Cell Clustering using Autoencoder and Network Fusion

**Version** 1.0.4

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**Description** A single-cell Clustering method using 'Autoencoder' and Network fusion ('sc-CAN') for segregating the cells from the high-dimensional 'scRNA-Seq' data. The software automatically determines the optimal number of clusters and then partitions the cells in a way such that the results are robust to noise and dropouts. 'sc-CAN' is fast and it supports Windows, Linux, and Mac OS.

**License** LGPL

**Encoding** UTF-8

**LazyData** true

**LazyDataCompression** xz

**Depends** R (>= 3.5.0), scDHA, FNN, purrr

**Imports** stats

**RoxygenNote** 7.1.2

**Suggests** knitr

**VignetteBuilder** knitr

**NeedsCompilation** no

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**Repository** CRAN

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adjustedRandIndex      *adjustedRandIndex*

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### Description

The function to calculate adjusted Rand index value with the inputs of true clusters and predicted clusters

### Usage

```
adjustedRandIndex(x, y)
```

### Arguments

x	A vector that contain predicted cluster assignment.
y	A vector that contain true cluster assignment.

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calculate\_celltype\_prob  
*calculate\_celltype\_prob*

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### Description

Calculate clusters and cell types similarity based on the markers.

### Usage

```
calculate_celltype_prob(clt_marker_list, marker_database_list, type = "jacc")
```

### Arguments

clt_marker_list	
marker_database_list	A list of markers for all cluster.
type	A parameter to select the method to measure cluster and cell type similarity <ul style="list-style-type: none"> <li>• jacc - Jaccard index.</li> <li>• ac - Accuracy.</li> <li>• fl - F1 score.</li> </ul>

**Value**

A confusion matrix between clusters and cell types. Each cell represents a probability of a cluster belongs to a cell type.

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curate_markers	<i>curate_markers</i>
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**Description**

Filter genes that have low p-value and fold-change.

**Usage**

```
curate_markers(
  whole_list,
  gene_names,
  wilcox_threshold = 0.001,
  logfc_threshold = 1.5
)
```

**Arguments**

<code>whole_list</code>	A list of markers for all clusters.
<code>gene_names</code>	All the gene names of the expression matrix.
<code>wilcox_threshold</code>	A threshold for p-value <code>wilcox_threshold = 0.001</code> by default.
<code>logfc_threshold</code>	A threshold for fold-change <code>logfc_threshold = 1.5</code> by default.

**Value**

A list of markers that are strong expressed for discovered clusters.

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find_markers	<i>find_markers</i>
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**Description**

Perform cluster-wise Wilcox test and fold-change for each gene.

**Usage**

```
find_markers(input_data_matrix, cluster_labels, identity = 1, threads = 8)
```

**Arguments**

<code>input_data_matrix</code>	An expression matrix in which rows are genes and columns are cells.
<code>cluster_labels</code>	A vector of cluster labels obtained from clustering methods.
<code>identity</code>	A parameter to select specific cluster identity = 1 by default.
<code>threads</code>	A parameter to control number of cores used for analysis threads = 1 by default.

**Value**

A list that contains p-value and fold-change ratio for all genes of each cluster.

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`find_specific_marker` *find\_specific\_marker*

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**Description**

Calculate cluster and cell type similarity based on the markers.

**Usage**

```
find_specific_marker(gene_name, f_list, type = "jacc")
```

**Arguments**

<code>gene_name</code>	A list of markers belong to the cluster.
<code>f_list</code>	A list of markers belongs to a reference cell type.
<code>type</code>	A parameter to select the method to measure cluster and cell type similarity <ul style="list-style-type: none"> <li>• jacc - Jaccard index.</li> <li>• ac - Accuracy.</li> <li>• f1 - F1 score.</li> </ul>

**Value**

A vector of probabilities of a cluster belongs to cell types.

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get_cluster_markers	<i>get_cluster_markers</i>
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**Description**

Find markers for each cluster

**Usage**

```
get_cluster_markers(input_data_matrix, labels_vector, threads = 1)
```

**Arguments**

`input_data_matrix` An expression matrix in which rows are genes and columns are cells.

`labels_vector` A vector of cluster labels obtained from clustering methods.

`threads` A parameter to control number of cores used for analysis threads = 1 by default.

**Value**

A list that contains markers for each cluster.

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scCAN	<i>scCAN</i>
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**Description**

This is the main function to perform sc-RNA seq data clustering. scCAN is fully unsupervised scRNA-seq clustering framework that uses deep neural network and network fusion-based clustering algorithm. First, scCAN applies a non-negative autoencoder to filter scRNA-seq data. Second, the filtered data is passed to stacked Bayesian autoencoder to get multiple low-dimensional representations of input data. Subsequently, scCAN converts these compressed data into networks and unify those networks to a single graph. Then, scCAN uses a spectral clustering algorithm to obtain final clusters assignment.

**Usage**

```
scCAN(  
  data,  
  sparse = FALSE,  
  n.neighbors = 30,  
  alpha = 0.5,  
  n.iters = 10,  
  ncores = 10,
```

```

    r.seed = 1,
    subsamp = T,
    k = 2:15,
    samp.size = 5000
  )

```

### Arguments

<code>data</code>	Gene expression matrix, with rows represent samples and columns represent genes.
<code>sparse</code>	Boolean variable indicating whether data is a sparse matrix. The input must be a non negative sparse matrix.
<code>n.neighbors</code>	Number of neighboring cells that are used to calculate the edge's weight. The number of neighbors are set <code>n.neighbors = 30</code> by default.
<code>alpha</code>	A hyper parameter that control the weight of graph. This values is set to <code>alpha = 0.5</code> by default.
<code>n.iters</code>	A hyper-parameter to set the number of network fusion iterations. It is set to <code>n.iters = 10</code> by default.
<code>ncores</code>	Number of processor cores to use.
<code>r.seed</code>	A parameter to set a seed for reproducibility. This values is set to <code>r.seed = 1</code> by default.
<code>subsamp</code>	Enable subsampling process for big data. This values is set to <code>subsamp = T</code> by default.
<code>k</code>	A vector to search for optimal number of cluster.
<code>samp.size</code>	A parameter to control number of sub-sampled cells.

### Value

List with the following keys:

- `cluster` - A numeric vector containing cluster assignment for each sample.
- `k` - The optimal number of cluster.
- `latent` - The latent data generated from autoencoders.

### References

1. Duc Tran, Hung Nguyen, Bang Tran, Carlo La Vecchia, Hung N. Luu, Tin Nguyen (2021). Fast and precise single-cell data analysis using a hierarchical autoencoder. *Nature Communications*, 12, 1029. doi: 10.1038/s41467-021-21312-2

### Examples

```

## Not run:
# Load the package and the example data (SCE dataset)
library(scCAN)
#Load example data
data("SCE")

```

```
#Get data matrix and label
data <- t(SCE$data); label <- as.character(SCE$cell_type1)

#Generate clustering result, the input matrix has rows as samples and columns as genes
result <- scCAN(data, r.seed = 1)

#Get the clustering result
cluster <- result$cluster

#Calculate adjusted Rand Index
ari <- round(scCAN::adjustedRandIndex(cluster,label), 2)
print(paste0("ARI = ", ari))

## End(Not run)
```

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SCE

*SCE*

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**Description**

SCE dataset includes scRNA-seq data and cell type information.

**Usage**

SCE

**Format**

An object of class list of length 2.

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