

# Package ‘ProFAST’

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**Type** Package

**Title** Probabilistic Factor Analysis for Spatially-Aware Dimension Reduction

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**Description** Probabilistic factor analysis for spatially-aware dimension reduction across multi-section spatial transcriptomics data with millions of spatial locations.

More details can be referred to Wei Liu, et al. (2023) <[doi:10.1101/2023.07.11.548486](https://doi.org/10.1101/2023.07.11.548486)>.

**License** GPL-3

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AddAdj

*Calculate the adjacency matrix given a spatial coordinate matrix*

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### Description

Calculate the adjacency matrix given a spatial coordinate matrix with 2-dimension or 3-dimension or more.

### Usage

```
AddAdj(
  pos,
  type = "fixed_distance",
  platform = c("Others", "Visium", "ST"),
  neighbors = 6,
  ...
)
```

## Arguments

pos	a matrix object, with columns representing the spatial coordinates that can be any dimension, i.e., 2, 3 and >3.
type	an optional string, specify which type of neighbors' definition. Here we provide two definition: one is "fixed_distance", the other is "fixed_number".
platform	a string, specify the platform of the provided data, default as "Others". There are more platforms to be chosen, including "Visium", "ST" and "Others" ("Others" represents the other SRT platforms except for 'Visium' and 'ST') The platform helps to calculate the adjacency matrix by defining the neighborhoods when type="fixed_distance" is chosen.
neighbors	an optional positive integer, specify how many neighbors used in calculation, default as 6.
...	Other arguments passed to <a href="#">getAdj_auto</a> .

## Details

When the type = "fixed\_distance", then the spots within the Euclidean distance cutoffs from one spot are regarded as the neighbors of this spot. When the type = "fixed\_number", the K-nearest spots are regarded as the neighbors of each spot.

## Value

return a sparse matrix, representing the adjacency matrix.

## References

None

## See Also

None

## Examples

```
data(CosMx_subset)
pos <- as.matrix(CosMx_subset@meta.data[,c("x", "y")])
Adj_sp <- AddAdj(pos)
```

AddParSettingFAST      *Add FAST model settings for a PRECASTObj object*

### Description

Add FAST model settings for a PRECASTObj object

### Usage

```
AddParSettingFAST(PRECASTObj, ...)
```

### Arguments

PRECASTObj	a PRECASTObj object created by <a href="#">CreatePRECASTObj</a> .
...	other arguments to be passed to <a href="#">model_set_FAST</a> function.

### Value

Return a revised PRECASTObj object with slot parameterList changed.

### References

None

coembedding\_umap      *Calculate UMAP projections for coembedding of cells and features*

### Description

Calculate UMAP projections for coembedding of cells and features

### Usage

```
coembedding_umap(
  seu,
  reduction,
  reduction.name,
  gene.set = NULL,
  slot = "data",
  assay = "RNA",
  seed = 1
)
```

## Arguments

seu	a Seurat object with coembedding in the reductions slot wiht component name reduction.
reduction	a string, specify the reduction component that denotes coembedding.
reduction.name	a string, specify the reduction name for the obtained UMAP projection.
gene.set	a string vector, specify the features (genes) in calculating the UMAP projection, default as all features.
slot	an optional string, specify the slot in the assay, default as ‘data’.
assay	an optional string, specify the assay name in the Seurat object when adding the UMAP projection.
seed	an optional integer, specify the random seed for reproducibility.

## Details

None

## Value

return a revised Seurat object by adding a new reduction component named ‘reduction.name’.

## References

None

## See Also

None

## Examples

```
data(pbmc3k_subset)
data(top5_signatures)

pbmc3k_subset <- coembedding_umap(
  pbmc3k_subset, reduction = "ncfm", reduction.name = "UMAPsig",
  gene.set = top5_signatures$gene
)
```

---

coembed_plot	<i>Coembedding dimensional reduction plot</i>
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**Description**

Graph output of a dimensional reduction technique on a 2D scatter plot where each point is a cell or feature and it's positioned based on the coembeddings determined by the reduction technique. By default, cells and their signature features are colored by their identity class (can be changed with the group\_by parameter).

**Usage**

```
coembed_plot(
  seu,
  reduction,
  gene_txtdata = NULL,
  cell_label = NULL,
  xy_name = reduction,
  dims = c(1, 2),
  cols = NULL,
  shape_cg = c(1, 5),
  pt_size = 1,
  pt_text_size = 5,
  base_size = 16,
  base_family = "serif",
  legend.point.size = 5,
  legend.key.size = 1.5,
  alpha = 0.3
)
```

**Arguments**

<code>seu</code>	a Seurat object with coembedding in the reductions slot with component name reduction.
<code>reduction</code>	a string, specify the reduction component that denotes coembedding.
<code>gene_txtdata</code>	a data.frame object with columns including 'gene' and 'label', specify the cell type/spatial domain and signature genes. Default as NULL, all features will be used in coembeddings.
<code>cell_label</code>	an optional character in columns of metadata, specify the group of cells/spots. Default as NULL, use Idents as the group.
<code>xy_name</code>	an optional character, specify the names of x and y-axis, default as the same as reduction.
<code>dims</code>	a positive integer vector with length 2, specify the two components for visualization.
<code>cols</code>	an optional string vector, specify the colors for cell group in visualization.

shape_cg	a positive integers with length 2, specify the shapes of cell/spot and feature in plot.
pt_size	an optional integer, specify the point size, default as 1.
pt_text_size	an optional integer, specify the point size of text, default as 5.
base_size	an optional integer, specify the basic size.
base_family	an optional character, specify the font.
legend.point.size	an optional integer, specify the point size of legend.
legend.key.size	an optional integer, specify the size of legend key.
alpha	an optional positive real, range from 0 to 1, specify the transparency of points.

## Details

None

## Value

return a ggplot object

## References

None

## See Also

[coembedding\\_umap](#)

## Examples

```
data(pbmc3k_subset)
data(top5_signatures)
coembed_plot(pbmc3k_subset, reduction = "UMAPsig",
            gene_txtdat = top5_signatures, pt_text_size = 3, alpha=0.3)
```

CosMx\_subset

*A Seurat object including spatial transcriptomics dataset from CosMx platform*

## Description

This data is a subset of SCLC CosMx spatial transcriptomics dataset.

## Usage

```
data(CosMx_subset)
```

## Format

A Seurat object, including count matrix, spatial coordinates, and manual annotation.

## Source

The data is from the CosMx SRT sequencing platform.

## References

None

## Examples

```
# Show some examples of how to use the dataset.
data(CosMx_subset)
library(Seurat)
CosMx_subset
```

**diagnostic.cor.eigs**     *Determine the dimension of low dimensional embedding*

## Description

This function estimate the dimension of low dimensional embedding for a given cell by gene expression matrix. For more details, see Franklin et al. (1995) and Crawford et al. (2010).

## Usage

```
diagnostic.cor.eigs(object, ...)

## Default S3 method:
diagnostic.cor.eigs(
  object,
  q_max = 50,
  plot = TRUE,
  n.sims = 10,
  parallel = TRUE,
  ncores = 10,
  seed = 1,
  ...
)

## S3 method for class 'Seurat'
diagnostic.cor.eigs(
  object,
  assay = NULL,
  slot = "data",
```

```
nfeatures = 2000,
q_max = 50,
seed = 1,
...
)
```

## Arguments

object	A Seurat or matrix object
...	Other arguments passed to <a href="#">diagnostic.cor.eigs.default</a> .
q_max	the upper bound of low dimensional embedding. Default is 50.
plot	a indicator of whether plot eigen values.
n.sims	number of simulaton times. Default is 10.
parallel	a indicator of whether use parallel analysis.
ncores	the number of cores used in parallel analysis. Default is 10.
seed	a postive integer, specify the random seed for reproducibility
assay	an optional string, specify the name of assay in the Seurat object to be used.
slot	an optional string, specify the name of slot.
nfeatures	an optional integer, specify the number of features to select as top variable features. Default is 2000.

## Value

A data.frame with attribute ‘q\_est’ and ‘plot’, which is the estimated dimension of low dimensional embedding. In addition, this data.frame containing the following components:

- q - The index of eigen values.
- eig\_value - The eigen values on observed data.
- eig\_sim - The mean value of eigen values of n.sims simulated data.
- q\_est - The selected dimension in attr(obj, ‘q\_est’).
- plot - The plot saved in attr(obj, ‘plot’).

## References

1. Franklin, S. B., Gibson, D. J., Robertson, P. A., Pohlmann, J. T., & Fralish, J. S. (1995). Parallel analysis: a method for determining significant principal components. *Journal of Vegetation Science*, 6(1), 99-106.
2. Crawford, A. V., Green, S. B., Levy, R., Lo, W. J., Scott, L., Svetina, D., & Thompson, M. S. (2010). Evaluation of parallel analysis methods for determining the number of factors. *Educational and Psychological Measurement*, 70(6), 885-901.

## Examples

```
n <- 100
p <- 50
d <- 15
object <- matrix(rnorm(n*d), n, d) %*% matrix(rnorm(d*p), d, p)
diagnostic.cor.eigs(object, n.sims=2)
```

FAST

*Run FAST model for a PRECASTObj object*

## Description

Run FAST model for a PRECASTObj object

## Usage

```
FAST(PRECASTObj, q = 15, fit.model = c("poisson", "gaussian"))
```

## Arguments

- |            |  |
|------------|--|
| PRECASTObj | a PRECASTObj object created by <a href="#">CreatePRECASTObj</a> .  |
| q          | an optional integer, specify the number of low-dimensional embeddings to extract in FAST   |
| fit.model  | an optional string, specify the version of FAST to be fitted. The Gaussian version models the log-count matrices while the Poisson versions models the count matrices; default as poisson. |

## Value

Return a revised PRECASTObj object with slot PRECASTObj@resList added by a FAST component.

## References

None

---

FAST\_run(Varitional) ICM-EM algorithm for implementing FAST model

---

## Description

(Varitional) ICM-EM algorithm for implementing FAST model

## Usage

```
FAST_run(
  XList,
  AdjList,
  q = 15,
  fit.model = c("gaussian", "poisson"),
  AList = NULL,
  maxIter = 25,
  epsLogLik = 1e-05,
  verbose = TRUE,
  seed = 1,
  error_heter = TRUE,
  Psi_diag = FALSE,
  Vint_zero = FALSE
)
```

## Arguments

XList	an M-length list consisting of multiple matrices with class dgCMatrix or matrix that specifies the count/log-count gene expression matrix for each data batch used for FAST model.
AdjList	an M-length list of sparse matrices with class dgCMatrix, specify the adjacency matrix used for intrinsic CAR model in FAST. We provide this interface for those users who would like to define the adjacency matrix by themselves.
q	an optional integer, specify the number of low-dimensional embeddings to extract in FAST. Larger q means more information extracted.
fit.model	an optional string, specify the version of FAST to be fitted. The Gaussian version models the log-count matrices while the Poisson versions models the count matrices; default as gaussian due to faster computation.
AList	an optional list with each component being a vector whose length is equal to the rows of component in XList, specify the normalization factor in FAST. The default is NULL that means the normalization factor equal to 1.
maxIter	the maximum iteration of ICM-EM algorithm. The default is 30.
epsLogLik	an optional positive value, tolerance of relative variation rate of the observed pseudo loglikelihood value, default as '1e-5'.
verbose	a logical value, whether output the information in iteration.
seed	a positive integer, the random seed to be set in initialization.

<code>error_heter</code>	a logical value, whether use the heterogenous error for FAST model, default as TRUE. If <code>error.heter=FALSE</code> , then the homogenous error is used.
<code>Psi_diag</code>	a logical value, whether set the conditional covariance matrix of the intrinsic CAR to diagonal, default as FALSE.
<code>Vint_zero</code>	an optional logical value, specify whether the intial value of intrinsic CAR component is set to zero; default as FALSE.

**Details**

None

**Value**

return a list including the following components: (1) hV: an M-length list consisting of spatial embeddings in FAST; (2) nu: the estimated intercept vector; (3) Psi: the estimated covariance matrix; (4) W: the estimated shared loading matrix; (5) Lam: the estimated covariance matrix of error term; (6): ELBO: the ELBO value when algorithm convergence; (7) ELBO\_seq: the ELBO values for all itrations.

**References**

None

**See Also**

[FAST\\_structure](#), [FAST](#), [model\\_set\\_FAST](#)

**FAST\_single**

*Fit FAST model for single-section SRT data*

**Description**

Fit FAST model for single-section SRT data.

**Usage**

```
FAST_single(
  seu,
  Adj_sp,
  q = 15,
  fit.model = c("poisson", "gaussian"),
  slot = "data",
  assay = NULL,
  reduction.name = "fast",
  verbose = TRUE,
  ...
)
```

## Arguments

seu	a Seurat object.
Adj_sp	a sparse matrix, specify the adjacency matrix among spots.
q	an optional integer, specify the number of low-dimensional embeddings to extract in FAST. Larger q means more information extracted.
fit.model	an optional string, specify the version of FAST to be fitted. The Gaussian version models the log-count matrices while the Poisson versions models the count matrices; default as poisson model.
slot	an optional string, specify the slot in Seurat object as the input of FAST model, default as ‘data’.
assay	an optional string, specify the assay in Seurat object, default as ‘NULL’ that means the default assay in Seurat object.
reduction.name	an optional string, specify the reduction name for the fast embedding, default as ‘fast’.
verbose	a logical value, whether output the information in iteration.
...	other arguments passed to <a href="#">FAST_run</a> .

## Value

return a list including the parameters set in the arguments.

## See Also

[FAST\\_run](#)

FAST_structure	(Varitional) ICM-EM algorithm for implementing FAST model with structurized parameters
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## Description

(Varitional) ICM-EM algorithm for implementing FAST model with structurized parameters

## Usage

```
FAST_structure(
  XList,
  AdjList,
  q = 15,
  fit.model = c("poisson", "gaussian"),
  parameterList = NULL
)
```

## Arguments

XList	an M-length list consisting of multiple matrices with class dgCMatrix or matrix that specify the count/log-count gene expression matrix for each data batch used for FAST model.
AdjList	an M-length list of sparse matrices with class dgCMatrix, specify the adjacency matrix used for intrinsic CAR model in FAST. We provide this interface for those users who would like to define the adjacency matrix by themselves.
q	an optional integer, specify the number of low-dimensional embeddings to extract in FAST
fit.model	an optional string, specify the version of FAST to be fitted. The Gaussian version models the log-count matrices while the Poisson versions models the count matrices; default as gaussian due to faster computation.
parameterList	an optional list, specify other parameters in FAST model; see <a href="#">model_set_FAST</a> for other parameters. The default is NULL that means the default parameters produced by <a href="#">model_set_FAST</a> is used.

## Details

None

## Value

return a list including the following components: (1) hV: an M-length list consisting of spatial embeddings in FAST; (2) nu: the estimated intercept vector; (3) Psi: the estimated covariance matrix; (4) W: the estimated shared loading matrix; (5) Lam: the estimated covariance matrix of error term; (6): ELBO: the ELBO value when algorithm convergence; (7) ELBO\_seq: the ELBO values for all iterations.

## References

None

## See Also

[FAST\\_run](#), [FAST](#), [model\\_set\\_FAST](#)

**find.signature.genes**   *Find the signature genes for each group of cell/spots*

## Description

Find the signature genes for each group of cell/spots based on coembedding distance and expression ratio.

**Usage**

```
find.signature.genes(  
  seu,  
  distce.assay = "distce",  
  ident = NULL,  
  expr.prop.cutoff = 0.1,  
  assay = NULL,  
  genes.use = NULL  
)
```

**Arguments**

seu	a Seurat object with coembedding in the reductions slot with component name reduction.
distce.assay	an optional character, specify the assay name that contains distance matrix between cells/spots and features, default as 'distce' (distance of coembeddings).
ident	an optional character in columns of metadata, specify the group of cells/spots. Default as NULL, use Idents as the group.
expr.prop.cutoff	an optional positive real ranging from 0 to 1, specify cutoff of expression proportion of features, default as 0.1.
assay	an optional character, specify the assay in seu, default as NULL, representing the default assay in seu.
genes.use	an optional string vector, specify genes as the signature candidates.

**Details**

In each data.frame object of the returned value, the row.names are gene names, and these genes are sorted by decreasing order of 'distance'. User can define the signature genes as top n genes in distance and that the 'expr.prop' larger than a cutoff. We set the cutoff as 0.1.

**Value**

return a list with each component a data.frame object having two columns: 'distance' and 'expr.prop'.

**References**

None

**See Also**

None

## Examples

```
library(Seurat)
data(pbmc3k_subset)
pbmc3k_subset <- pdistance(pbmc3k_subset, reduction='ncfm')
df_list_rna <- find.signature.genes(pbmc3k_subset)
```

`get.top.signature.dat` *Obtain the top signature genes and related information*

## Description

Obtain the top signature genes and related information.

## Usage

```
get.top.signature.dat(df.list, ntop = 5, expr.prop.cutoff = 0.1)
```

## Arguments

- `df.list` a list that is obtained by the function `find.signature.genes`.
- `ntop` an optional positive integer, specify the how many top signature genes extracted, default as 5.
- `expr.prop.cutoff` an optional positive real ranging from 0 to 1, specify cutoff of expression proportion of features, default as 0.1.

## Details

Using this function, we obtain the top signature genes and organize them into a data.frame. The ‘row.names’ are gene names. The colname ‘distance’ means the distance between gene (i.e., VPREB3) and cells with the specific cell type (i.e., B cell), which is calculated based on the co-embedding of genes and cells in the coembedding space. The distance is smaller, the association between gene and the cell type is stronger. The colname ‘expr.prop’ represents the expression proportion of the gene (i.e., VPREB3) within the cell type (i.e., B cell). The colname ‘label’ means the cell types and colname ‘gene’ denotes the gene name. By the data.frame object, we know ‘VPREB3’ is the one of the top signature gene of B cell.

## Value

return a ‘data.frame’ object with four columns: ‘distance’, ‘expr.prop’, ‘label’ and ‘gene’.

## References

None

**See Also**

None

**Examples**

```
library(Seurat)
data(pbmc3k_subset)
pbmc3k_subset <- pdistance(pbmc3k_subset, reduction='ncfm')
df_list_rna <- find.signature.genes(pbmc3k_subset)
dat.sig <- get.top.signature.dat(df_list_rna, ntop=5)
head(dat.sig)
```

---

get\_r2\_mcfadden

*Calcuuate the the adjusted McFadden's pseudo R-square*

---

**Description**

Calcuuate the the adjusted McFadden's pseudo R-square between the embeddings and the labels

**Usage**

```
get_r2_mcfadden(embeds, y)
```

**Arguments**

embeds	a n-by-q matrix, specify the embedding matrix.
y	a n-length vector, specify the labels.

**Details**

None

**Value**

return the adjusted McFadden's pseudo R-square.

**References**

McFadden, D. (1987). Regression-based specification tests for the multinomial logit model. Journal of econometrics, 34(1-2), 63-82.

`IntegrateSRTData`      *Integrate multiple SRT data into a Seurat object*

## Description

Integrate multiple SRT data based on the PRECASTObj object by FAST and other model fitting.

## Usage

```
IntegrateSRTData(
  PRECASTObj,
  seulist_HK,
  Method = c("iSC-MEB", "HarmonyLouvain"),
  seuList_raw = NULL,
  covariates_use = NULL,
  Tm = NULL,
  subsample_rate = 1,
  verbose = TRUE
)
```

## Arguments

<code>PRECASTObj</code>	a PRECASTObj object created by <a href="#">CreatePRECASTObj</a> .
<code>seulist_HK</code>	a list with Seurat object as component including only the housekeeping genes.
<code>Method</code>	a string, specify the method to be used and two methods are supported: iSC-MEB and HarmonyLouvain. The default is iSC-MEB.
<code>seuList_raw</code>	an optional list with Seurat object, the raw data.
<code>covariates_use</code>	a string vector, the colnames in <code>PRECASTObj@seulist[[1]]@meta.data</code> , representing other biological covariates to be considered when removing batch effects. This is achieved by adding additional covariates for biological conditions in the regression, such as case or control. Default as 'NULL', denoting no other covariates to be considered.
<code>Tm</code>	an optional numeric vector with the length equal to <code>PRECASTObj@seulist</code> , the time point information if the data include the temporal information. Default as NULL that means there is no temporal information.
<code>subsample_rate</code>	a real ranging in (0,1], specify the rate of spot drawing for speeding up the computation when the number of spots is very large. Default is 1, meaning using all spots.
<code>verbose</code>	an optional logical value, default as TRUE.

## Details

If `seuList_raw` is not equal NULL or `PRECASTObj@seuList` is not NULL, this function will remove the unwanted variations for all genes in `seuList_raw` object. Otherwise, only the the unwanted variation of genes in `PRECASTObj@seulist` will be removed. The former requires a big memory to

be run, while the latter not. To speed up the computation when the number of spots is very large, we also provide a subsampling schema controlled by the argument `subsample_rate`. When the total number of spots is larger than 80,000, this function will automatically draw 50,000 spots to calculate the parameters in the spatial linear model for removing unwanted variations.

### Value

Return a Seurat object by integrating all SRT data batches into a SRT data, where the column "batch" in the meta.data represents the batch ID, and the column "cluster" represents the clusters. The embeddings are put in `seu@reductions` slot and `Idents(seu)` is set to cluster label. Note that only the normalized expression is valid in the data slot while count is invalid.

`iscmеб_run`

*Fit an iSC-MEB model using specified multi-section embeddings*

### Description

Integrate multiple SRT data based on the PRECASTObj by FAST and iSC-MEB model fitting.

### Usage

```
iscmеб_run(
  VList,
  AdjList,
  K,
  beta_grid = seq(0, 5, by = 0.2),
  maxIter = 25,
  epsLogLik = 1e-05,
  verbose = TRUE,
  int.model = "EEE",
  init.start = 1,
  Sigma_equal = FALSE,
  Sigma_diag = TRUE,
  seed = 1
)
```

### Arguments

<code>VList</code>	a M-length list of embeddings. The i-th element is a $n_i \times q$ matrix, where $n_i$ is the number of spots of sample i, and $q$ is the number of embeddings. We provide this interface for those users who would like to define the embeddings by themselves.
<code>AdjList</code>	an M-length list of sparse matrices with class <code>dgCMatrix</code> , specify the adjacency matrix used for intrinsic CAR model in FAST. We provide this interface for those users who would like to define the adjacency matrix by themselves.
<code>K</code>	an integer, specify the number of clusters.

<code>beta_grid</code>	an optional vector of positive value, the candidate set of the smoothing parameter to be searched by the grid-search optimization approach, default as a sequence starts from 0, ends with 5, increase by 0.2.
<code>maxIter</code>	the maximum iteration of ICM-EM algorithm. The default is 25.
<code>epsLogLik</code>	a string, the species, one of 'Human' and 'Mouse'.
<code>verbose</code>	an optional integer, specify the number of housekeeping genes to be selected.
<code>int.model</code>	an optional string, specify which Gaussian mixture model is used in evaluating the initial values for iSC.MEB, default as "EEE"; and see <a href="#">Mclust</a> for more models' names.
<code>init.start</code>	an optional number of times to calculate the initial value (1 by default). When <code>init.start</code> is larger than 1, initial value will be determined by log likelihood of mclust results.
<code>Sigma_equal</code>	an optional logical value, specify whether Sigma matrices are equal, default as FALSE.
<code>Sigma_diag</code>	an optional logical value, specify whether Sigma matrices are diagonal matrices, default as TRUE.
<code>seed</code>	an optional integer, the random seed in fitting iSC-MEB model.

**Value**

returns a `iSCMEBResObj` object which contains all model results.

`model_set_FAST`      *Set parameters for FAST model*

**Description**

Prepare parameters setup for FAST model fitting.

**Usage**

```
model_set_FAST(
  maxIter = 30,
  epsLogLik = 1e-05,
  error_heter = TRUE,
  Psi_diag = FALSE,
  verbose = TRUE,
  seed = 1
)
```

**Arguments**

<code>maxIter</code>	the maximum iteration of ICM-EM algorithm. The default is 30.
<code>epsLogLik</code>	an optional positive value, tolerance of relative variation rate of the observed pseudo loglikelihood value, default as '1e-5'.

error_heter	a logical value, whether use the heterogenous error for FAST model, default as TRUE. If error.heter=FALSE, then the homogenous error is used.
Psi_diag	a logical value, whether set the conditional covariance matrices of intrinsic CAR to diagonal, default as FALSE
verbose	a logical value, whether output the information in iteration.
seed	a positive integer, the random seed to be set in initialization.

**Value**

return a Seurat object with new reduction (named reduction.name) added to the ‘reductions’ slot.

**Examples**

```
model_set_FAST(maxIter = 30, epsLogLik = 1e-5,
               error_heter=TRUE, Psi_diag=FALSE, verbose=TRUE, seed=2023)
```

**Description**

Cell-feature coembedding for scRNA-seq data based on FAST model.

**Usage**

```
NCFM(
  object,
  assay = NULL,
  slot = "data",
  nfeatures = 2000,
  q = 10,
  reduction.name = "ncfm",
  weighted = FALSE,
  var.features = NULL
)
```

**Arguments**

object	a Seurat object.
assay	an optional string, specify the name of assay in the Seurat object to be used, ‘NULL’ means default assay in seu.
slot	an optional string, specify the name of slot.
nfeatures	an optional integer, specify the number of features to select as top variable features. Default is 2000.

<code>q</code>	an optional positive integer, specify the dimension of low dimensional embeddings to compute and store. Default is 10.
<code>reduction.name</code>	an optional string, specify the dimensional reduction name, ‘ncfm’ by default.
<code>weighted</code>	an optional logical value, specify whether use weighted method.
<code>var.features</code>	an optional string vector, specify the variable features used to calculate cell embedding.

## Examples

```
data(pbmc3k_subset)
pbmc3k_subset <- NCFM(pbmc3k_subset)
```

NCFM\_fast

*Cell-feature coembedding for SRT data*

## Description

Run cell-feature coembedding for SRT data based on FAST model.

## Usage

```
NCFM_fast(
  object,
  Adj_sp,
  assay = NULL,
  slot = "data",
  nfeatures = 2000,
  q = 10,
  reduction.name = "fast",
  var.features = NULL,
  ...
)
```

## Arguments

<code>object</code>	a Seurat object.
<code>Adj_sp</code>	a sparse matrix, specify the adjacency matrix among spots.
<code>assay</code>	an optional string, the name of assay used.
<code>slot</code>	an optional string, the name of slot used.
<code>nfeatures</code>	an optional postive integer, the number of features to select as top variable features. Default is 2000.
<code>q</code>	an optional positive integer, specify the dimension of low dimensional embeddings to compute and store. Default is 10.
<code>reduction.name</code>	an optional string, dimensional reduction name, ‘fast’ by default.
<code>var.features</code>	an optional string vector, specify the variable features, used to calculate cell embedding.
<code>...</code>	Other argument passed to the <a href="#">FAST_run</a> .

## Examples

```
data(CosMx_subset)
pos <- as.matrix(CosMx_subset@meta.data[,c("x", "y")])
Adj_sp <- AddAdj(pos)
# Here, we set maxIter = 3 for fast computation and demonstration.
CosMx_subset <- NCFM_fast(CosMx_subset, Adj_sp = Adj_sp, maxIter=3)
```

---

pbmc3k\_subset

*A Seurat object including scRNA-seq PBMC dataset*

---

## Description

This data is a subset of PBMC3k scRNA-seq data in SeuratData package.

## Usage

```
data(pbmc3k_subset)
```

## Format

A Seurat object, including count matrix, and manual annotation.

## Source

The data is from the scRNA-seq sequencing platform.

## References

None

## Examples

```
# Show examples of how to use the dataset.
data(pbmc3k_subset)
library(Seurat)
pbmc3k_subset
```

<code>pdistance</code>	<i>Calculate the cell-feature distance matrix</i>
------------------------	---

**Description**

Calculate the cell-feature distance matrix based on coembeddings.

**Usage**

```
pdistance(object, reduction = "fast", assay.name = "distce", eta = 1e-10)
```

**Arguments**

<code>object</code>	a Seurat object.
<code>reduction</code>	a opstional string, dimensional reduction name, ‘fast’ by default.
<code>assay.name</code>	a opstional string, specify the new generated assay name, ‘distce’ by default.
<code>eta</code>	an optional postive real, a quantity to avoid numerical errors. 1e-10 by default.

**Details**

This function calculate the distance matrix between cells/spots and features, and then put the distance matrix in a new generated assay. This distance matrix will be used in the siganture gene identification.

**Examples**

```
data(pbmc3k_subset)
pbmc3k_subset <- NCFM(pbmc3k_subset)
pbmc3k_subset <- pdistance(pbmc3k_subset, "ncfm")
```

<code>RunHarmonyLouvain</code>	<i>Embedding alignment and clustering based on the embeddings from FAST</i>
--------------------------------	---

**Description**

Embedding alignment and clustering using the Harmony and Louvain based on the ebmeddings from FAST as well as determining the number of clusters.

**Usage**

```
RunHarmonyLouvain(PRECASTObj, resolution = 0.5)
```

**Arguments**

- `PRECASTObj` a PRECASTObj object created by [CreatePRECASTObj](#).  
`resolution` an optional real, the value of the resolution parameter, use a value above (below) 1.0 if you want to obtain a larger (smaller) number of communities.

**Value**

Return a revised PRECASTObj object with slot `PRECASTObj@resList` added by a Harmony component (including the aligned embeddings and embeddings of batch effects) and a Louvain component (including the clusters).

RuniSCMEB

*Fit an iSC-MEB model using the embeddings from FAST***Description**

Fit an iSC-MEB model using the embeddings from FAST and the number of clusters obtained by Louvain.

**Usage**

```
RuniSCMEB(PRECASTObj, ...)
```

**Arguments**

- `PRECASTObj` a PRECASTObj object created by [CreatePRECASTObj](#).  
`...` other arguments passed to [iscmeb\\_run](#).

**Value**

Return a revised PRECASTObj object with an added component iSCMEB in the slot `PRECASTObj@resList` (including the aligned embeddings, clusters and posterior probability matrix of clusters).

SelectHKgenes

*Select housekeeping genes***Description**

Select housekeeping genes for preparation of removing unwanted variations in expression matrices

**Usage**

```
SelectHKgenes(seuList, species = c("Human", "Mouse"), HK.number = 200)
```

**Arguments**

<code>seuList</code>	an M-length list consisting of Seurat object, include the information of expression matrix and spatial coordinates (named <code>row</code> and <code>col</code> ) in the slot <code>meta.data</code> .
<code>species</code>	a string, the species, one of 'Human' and 'Mouse'.
<code>HK.number</code>	an optional integer, specify the number of housekeeping genes to be selected.

**Value**

Return a string vector of the selected gene names.

<code>top5_signatures</code>	<i>A data.frame object including top five signature genes in scRNA-seq PBMC dataset</i>
------------------------------	---

**Description**

This data is a data.frame object that includes top five signature genes in scRNA-seq PBMC dataset

**Usage**

```
data(top5_signatures)
```

**Format**

A data.frame object, including signature genes, distance, and manual annotation.

**Source**

None

**References**

None

**Examples**

```
# Show examples of how to use the dataset.
data(top5_signatures)
head(top5_signatures)
```

---

transferGeneNames	<i>Transfer gene names from one format to the other format</i>
-------------------	--

---

## Description

Transfer gene names from one format to the other format for two species: human and mouse.

## Usage

```
transferGeneNames(  
  genelist,  
  now_name = "ensembl",  
  to_name = "symbol",  
  species = c("Human", "Mouse")  
)
```

## Arguments

genelist	a string vector, the gene list to be transferred.
now_name	a string, the current format of gene names, one of 'ensembl', 'symbol'.
to_name	a string, the format of gene names to transfer, one of 'ensembl', 'symbol'.
species	a string, the species, one of 'Human' and 'Mouse'.

## Value

Return a string vector of transferred gene names. The gene names not matched in the database will not change.

## Examples

```
## Not run:  
geneNames <- c("ENSG00000171885", "ENSG00000115756")  
transferGeneNames(geneNames, now_name = "ensembl", to_name="symbol", species="Human")  
  
## End(Not run)
```

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