Package 'WormTensor'

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Type Package

Title A Clustering Method for Time-Series Whole-Brain Activity Data of 'C. elegans'

Version 0.1.1

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Description A toolkit to detect clusters from distance matrices.

The distance matrices are assumed to be calculated between the cells of multiple animals ('Caenorhabditis elegans') from input time-series matrices. Some functions for generating distance matrices, performing clustering, evaluating the clustering, and visualizing the results of clustering and evaluation are available. We're also providing the download function to retrieve the calculated distance matrices from

'figshare' <https://figshare.com>.

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Encoding UTF-8

URL https://github.com/rikenbit/WormTensor

BugReports https://github.com/rikenbit/WormTensor/issues

Suggests knitr, rmarkdown, testthat (>= 3.0.0)

Config/testthat/edition 3

Depends R (>= 4.1.0)

Imports rTensor, usedist, dtwclust, clusterSim, clValid, aricode, cluster, ggplot2, Rtsne, uwot, factoextra, ggrepel, cowplot, methods

RoxygenNote 7.2.0

VignetteBuilder knitr

NeedsCompilation no

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AllGenerics_component Components for WormTensor object

Description

These are generic methods for WormTensor

Usage

```
worm_membership(object, k)
worm_clustering(
  object,
  algorithm = c("MCMI", "OINDSCAL", "CSPA"),
  num.iter = 30,
  thr = 1e-10,
  verbose = FALSE
)
worm_evaluate(object, labels = NULL)
worm_visualize(
  object,
  out.dir = tempdir(),
  algorithm = c("tSNE", "UMAP"),
  seed = 1234,
  tsne.dims = 2,
  tsne.perplexity = 15,
  tsne.verbose = FALSE,
  tsne.max_iter = 1000,
  umap.n_neighbors = 15,
  umap.n_components = 2,
  silhouette.summary = FALSE
)
```

as_worm_tensor 3

Arguments

object WormTensor object

k Assumed number of clusters
algorithm Dimensional reduction methods

num.iter The upper limit of iterations (Default value is 30)

thr The lower limit of relative change in estimates (Default value is 1E-10)

verbose Control message

labels Labels for external evaluation

out.dir Output directory (default: tempdir())

seed Arguments passed to set.seed (default: 1234)

tsne.dims Output dimensionality (default: 2)

tsne.perplexity

Perplexity paramete (default: 15)

tsne.verbose logical; Whether progress updates should be printed (default: TRUE)

tsne.max_iter The number of iterations (default: 1000)

umap.n_neighbors

The size of local neighborhood (default: 15)

umap.n_components

The dimension of the space to embed into (default: 2)

silhouette.summary

logical; If true a summary of cluster silhouettes are printed.

distance matrices.

Description

Generates WormTensor object A WormTensor object is generated from distance matrices.

Usage

```
as_worm_tensor(Ds)
```

Arguments

Ds A list containing distance matrices

Value

An object containing distance matrices and metadata

Examples

```
worm_download("mSBD", qc = "PASS")$Ds |> as_worm_tensor() -> object
```

WormTensor-class

S4 class used by as_worm_tensor.R, worm_membership.R, worm_clustering.R, worm_evaluate.R, worm_visualize.R

Description

S4 class used by as_worm_tensor.R, worm_membership.R, worm_clustering.R, worm_evaluate.R, worm_visualize.R

Slots

```
dist_matrices list
n_animals numeric
union_cellnames character
n_union_cells numeric
membership_tensor Tensor
k numeric
clustering_algorithm character
clustering numeric
weight numeric
factor matrix
consensus matrix
eval list
dimension_reduction_algorithm character
```

worm_clustering,WormTensor-method

Generates clustering result A clustering result is generated from a membership tensor.

Description

Generates clustering result A clustering result is generated from a membership tensor.

Usage

```
## S4 method for signature 'WormTensor'
worm_clustering(
  object,
  algorithm = c("MCMI", "OINDSCAL", "CSPA"),
  num.iter = 30,
  thr = 1e-10,
  verbose = FALSE
)
```

worm_distance 5

Arguments

object WormTensor object with a membership tensor

algorithm Clustering methods

num.iter The upper limit of iterations (Default value is 30)

thr The lower limit of relative change in estimates (Default value is 1E-10)

verbose Control message

Value

WormTensor object with a clustering result added

Examples

```
# Pipe Operation
worm_download("Euclid", qc = "WARN")$Ds |>
    as_worm_tensor() |>
        worm_membership(k = 6) -> object
worm_clustering(object, verbose = TRUE) -> ob_mcmi
worm_clustering(object, algorithm = "OINDSCAL", verbose = TRUE) -> ob_oind
worm_clustering(object, algorithm = "CSPA", verbose = TRUE) -> ob_cspa
```

worm_distance Generates distance matrices Distance matrices are generated between

the cells of multiple animals (Caenorhabditis elegans) from time-

series matrices.

Description

Generates distance matrices Distance matrices are generated between the cells of multiple animals (Caenorhabditis elegans) from time-series matrices.

Usage

```
worm_distance(data, distance = c("mSBD", "SBD", "Euclid"))
```

Arguments

data Time-series matrices

distance "mSBD" or "SBD" or "Euclid" can be specified. mSBD means modified Shape-

based distance.

Value

A list containing distance matrices

6 worm_download

Examples

```
# Toy data
n_cell_x <- 13
n_cell_y <- 24
n_cell_z <- 29
n_cells <- 30
n_time_frames <- 100
# 13 cells, 100 time frames
animal_x <- matrix(runif(n_cell_x * n_time_frames),</pre>
    nrow = n_cell_x, ncol = n_time_frames
rownames(animal_x) <- sample(seq(n_cells), n_cell_x)</pre>
colnames(animal_x) <- seq(n_time_frames)</pre>
# 24 cells, 100 time frames
animal_y <- matrix(runif(n_cell_y * n_time_frames),</pre>
    nrow = n_cell_y, ncol = n_time_frames
)
rownames(animal_y) <- sample(seq(n_cells), n_cell_y)</pre>
colnames(animal_y) <- seq(n_time_frames)</pre>
# 29 cells, 100 time frames
animal_z <- matrix(runif(n_cell_z * n_time_frames),</pre>
    nrow = n_cell_z, ncol = n_time_frames
)
rownames(animal_z) <- sample(seq(n_cells), n_cell_z)</pre>
colnames(animal_z) <- seq(n_time_frames)</pre>
# Positive Control of Difference between SBD and mSBD
animal_z[2, ] \leftarrow -animal_x[1, ]
X <- list(</pre>
    animal_x = animal_x,
    animal_y = animal_y,
    animal_z = animal_z
Ds_mSBD <- worm_distance(X, "mSBD")</pre>
```

worm_download

Downloads distance matrices 28 animals' data including 24 normal and 4 noisy are retrieved from figshare.

Description

Downloads distance matrices 28 animals' data including 24 normal and 4 noisy are retrieved from figshare.

Usage

```
worm_download(distance = c("mSBD", "Euclid"), qc = c("PASS", "WARN", "FAIL"))
```

Arguments

distance "mSBD" or "Euclid" can be specified

qc "PASS" or "WARN" or "FAIL" can be specified. "PASS" downloads 24 data

except 4 noisy data. "WARN" downloads 27 data except 1 noisy data. "FAIL"

downloads all 28 data.

Value

A List of containing distance matrices. The list also includes metadata for each animals.

Examples

```
Ds_Euclid <- worm_download("Euclid", qc = "WARN")
Ds_mSBD <- worm_download("mSBD", qc = "PASS")</pre>
```

worm_evaluate, WormTensor-method

Evaluates clustering result An evaluation result is generated from a WormTensor object.

Description

Evaluates clustering result An evaluation result is generated from a WormTensor object.

Usage

```
## S4 method for signature 'WormTensor'
worm_evaluate(object, labels = NULL)
```

Arguments

object WormTensor object with a result of worm_clustering

labels Labels for external evaluation

Value

WormTensor object with an evaluation result added

Examples

```
# Pipe Operation
worm_download("mSBD", qc = "PASS")$Ds |>
    as_worm_tensor() |>
        worm_membership(k = 6) |>
        worm_clustering() -> object
# Internal evaluation
worm_evaluate(object) -> object_internal
```

```
# External evaluation by sample labels
labels <- list(</pre>
   label1 = sample(3, length(object@clustering), replace = TRUE),
   label2 = sample(4, length(object@clustering), replace = TRUE),
   label3 = sample(5, length(object@clustering), replace = TRUE)
worm_evaluate(object, labels) -> object_external
# External evaluation by worm_download labels
Ds_mSBD <- worm_download("mSBD", qc = "PASS")</pre>
labels <- list(</pre>
   label1 = replace(
        Ds_mSBD$labels$Class,
        which(is.na(Ds_mSBD$labels$Class)),
        "NA"
   ),
   label2 = sample(4, length(object@clustering), replace = TRUE),
   label3 = sample(5, length(object@clustering), replace = TRUE)
worm_evaluate(object, labels) -> object_external_Class
```

worm_membership, WormTensor-method

Generates membership tensor A membership tensor is generated from distance matrices.

Description

Generates membership tensor A membership tensor is generated from distance matrices.

Usage

```
## S4 method for signature 'WormTensor'
worm_membership(object, k)
```

Arguments

object WormTensor object with distance matrices
k Assumed number of clusters

Value

WormTensor object with membership tensor added

Examples

```
# Pipe Operation
worm_download("mSBD", qc = "PASS")$Ds |>
    as_worm_tensor() -> object
# k=6
worm_membership(object, k = 6) -> object_k6
```

worm_visualize, WormTensor-method

Plots evaluation result A visualization result is generated from a WormTensor object.

Description

Plots evaluation result A visualization result is generated from a WormTensor object.

Usage

```
## S4 method for signature 'WormTensor'
worm_visualize(
   object,
   out.dir = tempdir(),
   algorithm = c("tSNE", "UMAP"),
   seed = 1234,
   tsne.dims = 2,
   tsne.perplexity = 15,
   tsne.verbose = FALSE,
   tsne.max_iter = 1000,
   umap.n_neighbors = 15,
   umap.n_components = 2,
   silhouette.summary = FALSE
)
```

Arguments

```
object
                  WormTensor object with a result of worm_evaluate
out.dir
                  Output directory (default: tempdir())
algorithm
                  Dimensional reduction methods
seed
                  Arguments passed to set.seed (default: 1234)
tsne.dims
                  Output dimensionality (default: 2)
tsne.perplexity
                  Perplexity parameter (default: 15)
                  logical; Whether progress updates should be printed (default: TRUE)
tsne.verbose
                  Number of iterations (default: 1000)
tsne.max_iter
```

```
umap.n_neighbors

The size of the local neighborhood (default: 15)

umap.n_components

The dimension of the space to embed into (default: 2)

silhouette.summary

logical; If true a summary of cluster silhouettes are printed.
```

Value

Silhouette plots. ARI with a merge result and each animal(with MCMI). Dimensional reduction plots colored by cluster, no. of identified cells, consistency(with labels), Class_label(with labels).

References

The .dist_nn function is quoted from dist_nn (not exported function) in package uwot(https://github.com/jlmelville/uwot/tree/f467185c8cbcd158feb60dde608c9da153ed10d7).

Examples

```
# Temporary directory to save figures
out.dir <- tempdir()</pre>
# Labels
worm_download("mSBD", qc = "PASS")$Ds |>
    as_worm_tensor() |>
        worm\_membership(k = 6) \mid >
            worm_clustering() -> object
Ds_mSBD <- worm_download("mSBD", qc = "PASS")</pre>
labels <- list(</pre>
    label1 = replace(
        Ds_mSBD$labels$Class,
        which(is.na(Ds_mSBD$labels$Class)),
        "NA"
    label2 = sample(4, length(object@clustering), replace = TRUE),
    label3 = sample(5, length(object@clustering), replace = TRUE)
)
# Pipe Operation (without Labels)
worm_download("mSBD", qc = "PASS")$Ds |>
    as_worm_tensor() |>
        worm_membership(k = 6) >
            worm_clustering() |>
                worm_evaluate() |>
                     worm_visualize(out.dir) -> object_no_labels
# Pipe Operation (with Labels)
worm_download("mSBD", qc = "PASS")$Ds |>
    as_worm_tensor() |>
        worm\_membership(k = 6) \mid >
            worm_clustering() |>
                worm_evaluate(labels) |>
```

worm_visualize(out.dir) -> object_labels

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