

Package ‘CICA’

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Title Clusterwise Independent Component Analysis

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Description

Clustering multi-subject resting state functional Magnetic Resonance Imaging data. This method enables the clustering of subjects based on multi-subject resting state functional Magnetic Resonance Imaging data. Objects are clustered based on similarities and differences in cluster-specific estimated components obtained by Independent Component Analysis.

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CICA

CICA: Clusterwise Independent Component Analysis

Description

Main function to perform Clusterwise Independent Component Analysis

Usage

```

CICA(
  DataList,
  nComp,
  nClus,
  method = "fastICA",
  RanStarts,
  RatStarts = NULL,
  pseudo = NULL,
  pseudoFac,
  userDef = NULL,
  userGrid = NULL,
  scalevalue = 1000,
  center = TRUE,
  maxiter = 100,
  verbose = TRUE,
  ctol = 1e-06,
  checks = TRUE
)

```

Arguments

DataList	a list of matrices
nComp	number or vector of ICA components per cluster
nClus	number or vector of clusters
method	Component method, default is fastICA. EVD for a fast eigen value based estimation
RanStarts	number of random starts
RatStarts	Generate rational starts. Either 'all' or a specific linkage method name (e.g., 'complete'). Use NULL to indicate that Rational starts should not be used.
pseudo	percentage value for perturbing rational starts to obtain pseudo rational starts
pseudoFac	factor to multiply the number of rational starts (7 in total) to obtain pseudorational starts
userDef	a user-defined starting seed stored in a data.frame, if NULL no userDef starting partition is used
userGrid	user supplied data.frame for multiple model CICA. First column are the requested components. Second column are the requested clusters
scalevalue	desired sum of squares of the block scaling procedure
center	mean center matrices
maxiter	maximum number of iterations for each start
verbose	print loss information to console
ctol	tolerance value for convergence criterion
checks	boolean parameter that indicates whether the input checks should be run (TRUE) or not (FALSE).

Value

CICA returns an object of `class` "CICA". It contains the estimated clustering, cluster specific component matrices and subject specific time course matrices

P	partitioning vector of size <code>length(DataList)</code>
Sr	list of size <code>nClus</code> , containing cluster specific independent components
Ais	list of size <code>length(DataList)</code> , containing subject specific time courses
Loss	loss function value of the best start
FinalLossDiff	value of the loss difference between the last two iterations of the algorithm.
IndLoss	a vector with containing the individual loss function values
LossStarts	loss function values of all starts
Iterations	Number of iterations
starts	dataframe with the used starting partitions

Author(s)

Jeffrey Durieux

Examples

```
## Not run:
CICA_data <- Sim_CICA(Nr = 15, Q = 5, R = 4, voxels = 100, timepoints = 10,
E = 0.4, overlap = .25, externalscore = TRUE)

multiple_output = CICA(DataList = CICA_data$X, nComp = 2:6, nClus = 1:5,
method = 'fastICA',userGrid = NULL, RanStarts = 30, RatStarts = NULL,
pseudo = c(0.1, 0.2),pseudoFac = 2, userDef = NULL, scalevalue = 1000,
center = TRUE,maxiter = 100, verbose = TRUE, ctol = .000001)

summary(multiple_output$Q_5_R_4)

plot(multiple_output$Q_5_R_4)

## End(Not run)
```

computeRVmat

Compute modified RV matrix

Description

This function computes a NxN modified RV matrix

Usage

```
computeRVmat(DataList = DataList, dist = TRUE, verbose = TRUE)
```

Arguments

DataList	a list with matrices
dist	boolean if TRUE distance object is returned
verbose	boolean if TRUE progressbar is printed to the console

Value

RVsS	a square similarity matrix of class <code>matrix</code> or distance object of class <code>dist</code> containing the pairwise modified RV values
------	--

Examples

```
## Not run:
CICA_data <- Sim_CICA(Nr = 15, Q = 5, R = 4, voxels = 100, timepoints = 10,
E = 0.4, overlap = .25, externalscore = TRUE)
#Compute single subject ICAs (nClus equals length(ExampleData))
output <- CICA(DataList = CICA_data$X, nStarts = 1,
               nComp = 5, nClus = 9, verbose = FALSE)
```

```
RV <- computeRVmat(DataList = output$Sr, dist = TRUE,
                   verbose = FALSE)

# apply hierarchical clustering on RV output
hcl <- hclust(RV)
plot(hcl)

# low dimensional visualisation using Classical Multidimensional Scaling
mds <- cmdscale(RV)
plot(mds)

## End(Not run)
```

embed_papaya

Embed images with Papaya

Description

Writes temporary images out from nifti objects or passes character filenames of images to papaya JS viewer

Usage

```
embed_papaya(images, outdir = NULL)
```

Arguments

images character filenames or nifti objects to be viewed
outdir output directory for index and all to go

Value

Output html

FindRationalStarts

Plot method for rstarts object

Description

Two step clustering for finding rational start partitions

Usage

```

FindRationalStarts(
  DataList,
  RatStarts = "all",
  nComp,
  nClus,
  scalevalue = NULL,
  center = TRUE,
  verbose = TRUE,
  pseudo = NULL,
  pseudoFac = NULL
)

## S3 method for class 'rstarts'
plot(x, type = 1, mdsdim = 2, nClus = NULL, ...)

```

Arguments

DataList	a list of matrices
RatStarts	type of rational start. 'all' computes all types of hclust methods
nComp	number of ICA components to extract
nClus	Number of clusters for rectangles in dendrogram, default NULL is based on number of clusters present in the object
scalevalue	scale each matrix to have an equal sum of squares
center	mean center matrices
verbose	print output to console
pseudo	percentage value for perturbing rational starts to obtain pseudo rational starts
pseudoFac	how many pseudo starts per rational start
x	an object of class rstarts
type	type of plot, 1 for a dendrogram, 2 for a multidimensional scaling configuration
mdsdim	2 for two dimensional mds configuration, 3 for a three dimensional configuration
...	optional arguments passed to hclust function

Value

dataframe with (pseudo-) rational and dist object based on the pairwise modified RV values

References

Durieux, J., & Wilderjans, T. F. (2019). Partitioning subjects based on high-dimensional fMRI data: comparison of several clustering methods and studying the influence of ICA data reduction in big data. *Behaviormetrika*, 46(2), 271-311.

Examples

```
## Not run:
CICA_data <- Sim_CICA(Nr = 15, Q = 5, R = 4, voxels = 100, timepoints = 10,
E = 0.4, overlap = .25, externalscore = TRUE)
rats <- FindRationalStarts(DataList = CICA_data$X, nComp = 5, nClus = 4, verbose = TRUE, pseudo = .2)
plot(rats, type = 1, method = 'ward.D2')
plot(rats, type = 2, method = 'ward.D2')
plot(rats, type = 2, method = 'ward.D2', mdsdim = 3)

## End(Not run)

## Not run:
CICA_data <- Sim_CICA(Nr = 15, Q = 5, R = 4, voxels = 100, timepoints = 10,
E = 0.4, overlap = .25, externalscore = TRUE)
Out_starts <- FindRationalStarts(DataList = CICA_data$X, nComp = 5, nClus = 4, scalevalue = 1000)
plot(Out_starts)
plot(Out_starts, type = 2)
plot(Out_starts, type = 2, mdsdim = 3, method = 'ward.D2')

## End(Not run)
```

GenRanStarts

Generate random starts

Description

Generate random starts

Usage

```
GenRanStarts(
  RanStarts,
  nClus,
  nBlocks,
  ARIlLim = 0.2,
  itmax = 1000,
  verbose = FALSE
)
```

Arguments

RanStarts	number of randomstarts to generate
nClus	number of clusters
nBlocks	number of objects
ARIlLim	maximal value of adjusted Rand Index

itmax maximum number of iterations used to find suitable random starts
 verbose boolean that indicates whether the output should be printed on the console

Value

a list where the first element is a matrix with random starts, second element all pairwise ARIs #'

GenRatStarts

Title

Description

Title

Usage

```
GenRatStarts(  
  DataList,  
  RatStarts,  
  nComp,  
  nClus,  
  scalevalue,  
  center,  
  verbose,  
  pseudo,  
  pseudoFac  
)
```

Arguments

DataList	DataList
RatStarts	Type of rational start
nComp	number of components
nClus	number of clusters
scalevalue	value for blockscaling procedure
center	center
verbose	verbose
pseudo	percentage used for perturbation rational starts (between 0)
pseudoFac	multiplication factor for pseudo rational starts

Value

out

get_papaya_version	<i>Get Papaya Version</i>
--------------------	---------------------------

Description

Reads the papaya.js file installed and determines version and build

Usage

```
get_papaya_version()
```

Value

List of build and version, both characters

loadNIFTIs	<i>Load Nifti files from directory</i>
------------	--

Description

Load Nifti files from directory

Usage

```
loadNIFTIs(dir, toMatrix = TRUE)
```

Arguments

dir	Input directory containing nifti files
toMatrix	logical if TRUE nifti's are converted to matrices

Value

list object containing Voxel by Time course matrices

Examples

```
## Not run:  
nifs <- loadNIFTIs('<FolderPath>', toMatrix = T)  
outnif <- CICA(DataList = nifs, RanStarts = 2, nComp = 10, nClus = 2)  
  
## End(Not run)
```

matcher	<i>Match components between cluster specific spatial maps</i>
---------	---

Description

Match components between cluster specific spatial maps

Usage

```
matcher(x, reference, RV = FALSE, ...)
```

Arguments

x	object of class CICA
reference	integer cluster index that serves as the reference. If nifti path is supplied, clusters will be matched to this template
RV	compute modified-RV between cluster components
...	other arguments

Value

out

Examples

```
## Not run:
CICA_data <- Sim_CICA(Nr = 15, Q = 5, R = 4, voxels = 100, timepoints = 10,
E = 0.4, overlap = .25, externalscore = TRUE)

multiple_output = CICA(DataList = CICA_data$X, nComp = 2:6, nClus = 1:5,
userGrid = NULL, RanStarts = 30, RatStarts = NULL, pseudo = c(0.1, 0.2),
pseudoFac = 2, userDef = NULL, scalevalue = 1000, center = TRUE,
maxiter = 100, verbose = TRUE, ctol = .000001)

matcher(multiple_output$Q_5_R_4, reference = 1, RV = TRUE)

## End(Not run)
```

matcher.CICA	<i>Match components between cluster specific spatial maps</i>
--------------	---

Description

Match components between cluster specific spatial maps

Usage

```
## S3 method for class 'CICA'  
matcher(x, reference = 1, RV = FALSE, ...)
```

Arguments

x	object of class CICA
reference	integer cluster index that serves as the reference. If nifti path is supplied, clusters will be matched to this template
RV	compute modified-RV between cluster components
...	other arguments

Value

out

Examples

```
## Not run:  
CICA_data <- Sim_CICA(Nr = 15, Q = 5, R = 4, voxels = 100, timepoints = 10,  
E = 0.4, overlap = .25, externalscore = TRUE)  
  
multiple_output = CICA(DataList = CICA_data$X, nComp = 2:6, nClus = 1:5,  
userGrid = NULL, RanStarts = 30, RatStarts = NULL, pseudo = c(0.1, 0.2),  
pseudoFac = 2, userDef = NULL, scalevalue = 1000, center = TRUE,  
maxiter = 100, verbose = TRUE, ctol = .000001)  
  
matcher(multiple_output$Q_5_R_4, reference = 1, RV = TRUE)  
  
## End(Not run)
```

`mpinv` *Moore Penrose inverse*

Description

Moore Penrose inverse

Usage

`mpinv(X)`

Arguments

`X` input matrix

Value

mp Moore Penrose inverse of matrix X

`papaya` *View images with Papaya*

Description

Writes temporary images out from nifti objects or passes character filenames of images to papaya JS viewer

Usage

`papaya(images, outdir = NULL, ...)`

Arguments

`images` character filenames or nifti objects to be viewed
`outdir` output directory for index and all to go
`...` Options to be passed to [pass_papaya](#)

Value

Output directory where index.html, js, and copied nii.gz files

Examples

```
## Not run:
library(neurobase)
x = nifti(img = array(rnorm(100^3), dim= rep(100, 3)), dim=rep(100, 3), datatype=16)
thresh = datatyper(x > 1)
index.file = papaya(list(x, thresh))

## End(Not run)
```

papaya_div

Papaya Div element output

Description

Get the necessary div output for embedding a papaya image

Usage

```
papaya_div()
```

Value

Character string

Examples

```
papaya_div()
```

pass_papaya

View images with Papaya

Description

Writes temporary images out from nifti objects or passes character filenames of images to papaya JS viewer

Usage

```
pass_papaya(
  L = NULL,
  outdir = NULL,
  daemon = FALSE,
  close_on_exit = TRUE,
  sleeper = 3,
  version = "0.8",
  build = "982"
)
```

Arguments

L	list of arguments passed to papaya using params
outdir	output directory for index and all to go
daemon	Argument passed to server_config
close_on_exit	Should the server close once the function finishes?
sleeper	Time in seconds to sleep if close_on_exit = TRUE. This allows the server to start up.
version	Version of papaya.js and papaya.css to use
build	Build of papaya.js and papaya.css to use

plot.CICA

Plot method for CICA

Description

Plot method for CICA. This function shows the cluster specific independent components in an interactive viewer using the papayar package

Usage

```
## S3 method for class 'CICA'
plot(x, brain = "auto", cluster = 1, ...)
```

Arguments

x	Object of class CICA
brain	auto
cluster	Components of cluster to plot. Only used when non fMRI related data is used
...	other arguments

Examples

```
## Not run:
CICA_data <- Sim_CICA(Nr = 15, Q = 5, R = 4, voxels = 100, timepoints = 10,
E = 0.4, overlap = .25, externalscore = TRUE)

multiple_output = CICA(DataList = CICA_data$X, nComp = 2:6, nClus = 1:5,
userGrid = NULL, RanStarts = 30, RatStarts = NULL, pseudo = c(0.1, 0.2),
pseudoFac = 2, userDef = NULL, scalevalue = 1000, center = TRUE,
maxiter = 100, verbose = TRUE, ctol = .000001)

plot(multiple_output$Q_5_R_4, cluster = 2)

## End(Not run)
```

`plot.ModSel`*Plot method for sequential model selection*

Description

Plot method for the sequential model selection option for CICA

Usage

```
## S3 method for class 'ModSel'  
plot(x, ...)
```

Arguments

<code>x</code>	Object of class <code>ModSel</code>
<code>...</code>	other arguments

Examples

```
## Not run:  
CICA_data <- Sim_CICA(Nr = 15, Q = 5, R = 4, voxels = 100, timepoints = 10,  
E = 0.4, overlap = .25, externalscore = TRUE)  
  
multiple_output = CICA(DataList = CICA_data$X, nComp = 2:6, nClus = 1:5,  
userGrid = NULL, RanStarts = 30, RatStarts = NULL, pseudo = c(0.1, 0.2),  
pseudoFac = 2, userDef = NULL, scalevalue = 1000, center = TRUE,  
maxiter = 100, verbose = TRUE, ctol = .000001)  
  
ModSelOutput <- SequentialScree(multiple_output)  
  
plot(ModSelOutput)  
  
## End(Not run)
```

`SequentialScree`*Sequential Model Selection for Multiple CICA model*

Description

Sequential Model Selection for Multiple CICA model

Usage

```
SequentialScree(x)
```

Arguments

x an object of class MultipleCICA

Value

a list object

Examples

```
## Not run:
CICA_data <- Sim_CICA(Nr = 15, Q = 5, R = 4, voxels = 100, timepoints = 10,
E = 0.4, overlap = .25, externalscore = TRUE)

multiple_output = CICA(DataList = CICA_data$X, nComp = 2:6, nClus = 1:5,
userGrid = NULL, RanStarts = 30, RatStarts = NULL, pseudo = c(0.1, 0.2),
pseudoFac = 2, userDef = NULL, scalevalue = 1000, center = TRUE,
maxiter = 100, verbose = TRUE, ctol = .000001)

ModSelOutput <- SequentialScree(multiple_output)

plot(ModSelOutput)

## End(Not run)
```

Sim_CICA

Simulate CICA data

Description

Simulate CICA data

Usage

```
Sim_CICA(
  Nr,
  Q,
  R,
  voxels,
  timepoints,
  E,
  overlap = NULL,
  externalscore = FALSE
)
```


Arguments

Nr	number of subjects per cluster
Q	number of components
R	number of clusters
voxels	number of voxels
timepoints	number of time points
E	proportion of independent gaussian noise
overlap	amount of overlap between S across clusters. Smaller value means more overlap
externalscore	add simulated external score (default is FALSE)

Value

a list with simulated CICA data

Examples

```
## Not run:

#Use set.seed(1) to obtain the dataset used in the article "Clusterwise
#Independent Component Analysis (CICA): an R package for clustering subjects
#based on ICA patterns underlying three-way (brain) data"

Xe <- Sim_CICA(Nr = 15, Q = 5, R = 4, voxels = 100, timepoints = 10,
E = 0.4, overlap = .25, externalscore = TRUE)

## End(Not run)
```

Sr_to_nifti

Convert Cluster specific independent components to NIFTI format

Description

Convert Cluster specific independent components to NIFTI format

Usage

```
Sr_to_nifti(x, write = FALSE, ...)
```

Arguments

x	an object of class CICA
write	if TRUE, NIFTI files are written to current working directory
...	other arguments passed to RNifti::writeNifti

Value

a list with niftiImage files

Examples

```
## Not run:
nifs <- loadNIFTIs('<FolderPath>', toMatrix = T)
outnif <- CICA(DataList = nifs, RanStarts = 2, nComp = 10, nClus = 2)
test <- Sr_to_nifti(outnif, write = T, datatype = 'int16', version = 2)

## End(Not run)
```

summary.CICA

Summary method for class CICA

Description

Summarize a CICA analysis

Usage

```
## S3 method for class 'CICA'
summary(object, ...)
```

Arguments

object Object of the type produced by [CICA](#)
... Additional arguments

Value

summary.CICA returns an overview of the estimated clustering of a [CICA](#) analysis

PM Partitioning matrix
tab tabulation of the clustering
Loss Loss function value of the solution

Examples

```
## Not run:
CICA_data <- Sim_CICA(Nr = 15, Q = 5, R = 4, voxels = 100, timepoints = 10,
E = 0.4, overlap = .25, externalscore = TRUE)

multiple_output = CICA(DataList = CICA_data$X, nComp = 2:6, nClus = 1:5,
userGrid = NULL, RanStarts = 30, RatStarts = NULL, pseudo = c(0.1, 0.2),
pseudoFac = 2, userDef = NULL, scalevalue = 1000, center = TRUE,
maxiter = 100, verbose = TRUE, ctol = .000001)
```

```
summary(multiple_output$Q_5_R_4)

## End(Not run)
```

summary.MultipleCICA *Summary method for class MultipleCICA*

Description

Summarize a CICA analysis

Usage

```
## S3 method for class 'MultipleCICA'
summary(object, ...)
```

Arguments

object	Object of the type produced by CICA
...	Additional arguments

Value

summary.MultipleCICA returns an overview of the estimated clustering of a [CICA](#) analysis

PM	Partitioning matrix
tab	tabulation of the clustering
Loss	Loss function value of the solution

Examples

```
## Not run:
CICA_data <- Sim_CICA(Nr = 15, Q = 5, R = 4, voxels = 100, timepoints = 10,
E = 0.4, overlap = .25, externalscore = TRUE)

multiple_output = CICA(DataList = CICA_data$X, nComp = 2:6, nClus = 1:5,
userGrid = NULL, RanStarts = 30, RatStarts = NULL, pseudo = c(0.1, 0.2),
pseudoFac = 2, userDef = NULL, scalevalue = 1000, center = TRUE,
maxiter = 100, verbose = TRUE, ctol = .000001)

summary(multiple_output$Q_5_R_4)

## End(Not run)
```

update_papaya_build *Update Papaya build version from GitHub*

Description

Updates the papaya version in the papayar package to the most current on GitHub

Usage

```
update_papaya_build(  
  type = c("standard", "minimal", "nodicom", "nojquery", "standard-with-atlas-local",  
          "standard-with-atlas"),  
  verbose = TRUE  
)
```

Arguments

type	Type of release. Standard is default
verbose	Should download progress be shown?

Value

Result of [get_papaya_version](#) after downloading

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