

# Package ‘MixfMRI’

October 12, 2022

**Version** 0.1-2

**Date** 2021-11-06

**Title** Mixture fMRI Clustering Analysis

**Depends** R (>= 4.0.0)

**Imports** MASS, Matrix, RColorBrewer, fftw, MixSim, EMCluster

**Enhances** pbdMPI (>= 0.3-4), AnalyzeFMRI, oro.nifti

**LazyLoad** yes

**LazyData** yes

**Description** Utilizing model-based clustering (unsupervised) for functional magnetic resonance imaging (fMRI) data. The developed methods (Chen and Maitra (2021) <[arXiv:2102.03639](https://arxiv.org/abs/2102.03639)>) include 2D and 3D clustering analyses (for p-values with voxel locations) and segmentation analyses (for p-values alone) for fMRI data where p-values indicate significant level of activation responding to stimulate of interesting. The analyses are mainly identifying active voxel/signal associated with normal brain behaviors. Analysis pipelines (R scripts) utilizing this package (see examples in 'inst/workflow/') is also implemented with high performance techniques.

**License** Mozilla Public License 2.0

**BugReports** <https://github.com/snoweye/MixfMRI/issues>

**URL** <https://github.com/snoweye/MixfMRI>

**NeedsCompilation** yes

**Maintainer** Wei-Chen Chen <[wccsnow@gmail.com](mailto:wccsnow@gmail.com)>

**Author** Wei-Chen Chen [aut, cre],  
Ranjan Maitra [aut],  
Dan Nettleton [ctb]

**Repository** CRAN

**Date/Publication** 2021-11-07 17:20:07 UTC

## R topics documented:

MixfMRI-package	2
algorithm	3
Compute Q values	5
Compute Statistics for Log Odds Ratio of Posterior Probability	6
Covariance Matrices	7
Covariance Matrices of Logit ETA	10
Density function of p-values	12
Example Datasets	13
False Discovery Rates for Spatial Signals	14
Generalized Cluster-Based Analysis (CBA) Method	15
initial	17
Likelihood Mixture Tests with Identity Cov Matrix or Only p-values	18
LRT	20
Main functions	22
MixfMRI Control	25
Plotting	27
Print Objects	29
Simulations	30
Smoothing	32
Summarized Overlap	32
<b>Index</b>	<b>34</b>

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MixfMRI-package	<i>fMRI Clustering Analysis</i>
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### Description

Utilizing model-based clustering (unsupervised) for fMRI data especially in a distributed manner. The methods includes 2D and 3D clustering analyses and segmentation analyses for fMRI signals where p-values are significant levels of active voxels which respond to stimulate of interesting. The analyses are mainly identifying active voxels/signals from normal brain behaviors. Workflows are also implemented utilizing high performance techniques.

### Details

Package:	MixfMRI
Type:	Package
License:	GPL (>= 2)
LazyLoad:	yes

The main function of this package is `fclust()` that implements model-based clustering algorithm for fMRI signal data and provides unsupervised clustering results for the data. Several workflows implemented with high-performance computing techniques are also built in for automatically pro-

ness clustering, hypothesis, cluster merging, and visualizations.

### Author(s)

Wei-Chen Chen and Ranjan Maitra.

### References

Chen, W.-C. and Maitra, R. (2021) “A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies”, *arXiv:2102.03639*.

### See Also

[fclust\(\)](#), [set.global\(\)](#).

### Examples

```
library(MixfMRI, quietly = TRUE)

.rem <- function(){

  demo(fclust3d, 'MixfMRI', ask=FALSE, echo=FALSE)
  demo(fclust2d, 'MixfMRI', ask=FALSE, echo=FALSE)

}
```

---

algorithm

*Main algorithms implemented in fclust*

---

### Description

Main algorithms implemented in fclust.

### Usage

```
ecm.step.gbd(PARAM.org)
```

```
apecma.step.gbd(PARAM.org)
```

```
em.step.gbd(PARAM.org)
```

### Arguments

PARAM.org      an initialized PARAM, usually returned by [set.global\(\)](#), [initial.em.gbd\(\)](#), and [initial.RndEM.gbd\(\)](#).

**Details**

These are main algorithms implemented in `fclust()`.

**Value**

Return an optimized PARAM.

**Author(s)**

Wei-Chen Chen and Ranjan Maitra.

**References**

Chen, W.-C. and Maitra, R. (2021) “A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies”, *arXiv:2102.03639*.

**See Also**

`set.global()`, `fclust()`, `PARAM`, `PARAM.org`.

**Examples**

```
library(MixfMRI, quietly = TRUE)
library(EMcluster, quietly = TRUE)
# .FC.CT$algorithm <- "em"
# .FC.CT$model.X <- "v"
# .FC.CT$ignore.X <- TRUE
.FC.CT$check.X.unit <- FALSE

### Test toy1.
set.seed(1234)
X.gbd <- toy1$X.gbd
PV.gbd <- toy1$PV.gbd
PARAM <- set.global(X.gbd, PV.gbd, K = 2)
PARAM.new <- initial.em.gbd(PARAM)
PARAM.toy1 <- em.step.gbd(PARAM.new)
id.toy1 <- .MixfMRIEnv$CLASS.gbd
print(PARAM.toy1$ETA)
RRand(toy1$CLASS.gbd, id.toy1)

.rem <- function(){

  ### Test toy2.
  set.seed(1234)
  X.gbd <- toy2$X.gbd
  PV.gbd <- toy2$PV.gbd
  PARAM <- set.global(X.gbd, PV.gbd, K = 3)
  PARAM.new <- initial.em.gbd(PARAM)
  PARAM.toy2 <- em.step.gbd(PARAM.new)
```

```
id.toy2 <- .MixfMRIEnv$CLASS.gbd
print(PARAM.toy2$ETA)
RRand(toy2$CLASS.gbd, id.toy2)

}
```

---

Compute Q values      *Q-values using Benjamini and Hochberg (1995)*

---

### Description

Compute q-values Benjamini and Hochberg's (1995) approach for controlling FDR.

### Usage

```
qvalue(p, method = c("BH1995", "BY2001"))
```

### Arguments

p	a p-value vector.
method	using method by either BH1995 or BY2001

### Details

This function compute q-values using Benjamini and Hochberg's (1995) approach for controlling FDR. The function `bh.fdr` is originally written by Dr. Dan Nettleton.

The Benjamini and Yeekutieli's (2001) approach for controlling FDR using the function `by.fdr` is coded by Wei-Chen Chen.

### Value

Return corresponding q-values for the input p-values.

### Author(s)

Dan Nettleton.

Modified by Wei-Chen Chen.

### References

Chen, W.-C. and Maitra, R. (2021) "A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies", *arXiv:2102.03639*.

### See Also

[dpval\(\)](#), [dmixpval\(\)](#).

**Examples**

```
library(MixfMRI, quietly = TRUE)
set.seed(1234)
da <- gendataset(phantom = shepp1fMRI, overlap = 0.01)
p <- da$pval[!is.na(da$pval)][1:100]
qvalue(p)
```

---

Compute Statistics for Log Odds Ratio of Posterior Probability  
*Compute Statistics for Log Odds Ratio of Posterior Probability*

---

**Description**

The function computes statistics for log odds ratio of posterior probability.

**Usage**

```
logor.stat(x, fcobj, post.z, cov.param = NULL, cov.post.z = NULL,
           cov.logit.z = NULL, all.x = FALSE, drop.ETA1 = FALSE)
```

**Arguments**

<code>x</code>	an input list of two elements <code>X.gbd</code> and <code>PV.gbd</code> .
<code>fcobj</code>	a <code>fclust</code> object.
<code>post.z</code>	a matrix of <code>dim = N * K</code> for posterior probabilities, which is also the return value of <code>post.prob()</code> .
<code>cov.param</code>	a covariance matrix of <code>dim = d * d</code> for parameters, which is also a return of <code>cov.param()</code> . <code>d</code> is total number of parameters which is dependent on data and models.
<code>cov.post.z</code>	a covariance list of length equal to number of active voxels, which is also a return of <code>cov.post.z()</code> .
<code>cov.logit.z</code>	a covariance list of length equal to number of active voxels, which is also a return of <code>cov.logit.z()</code> .
<code>all.x</code>	all cov matrices for all observations are returned if <code>TRUE</code> , while for only active observations (those of class ids are greater than 1) if <code>FALSE</code> .
<code>drop.ETA1</code>	if drop the <code>ETA[1]</code> from the cov matrix due to the <code>min.1st.prop</code> constrain.

**Details**

For posterior probability, this function compute log odd ratio, cov matrix of log odd ratio, degrees of freedom, and testing statistics.

**Value**

A list is returned with four elements: `log.or`, `cov.log.or`, `df`, and `test.stat`.

**Author(s)**

Wei-Chen Chen and Ranjan Maitra.

**References**

Chen, W.-C. and Maitra, R. (2021) “A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies”, *arXiv:2102.03639*.

**See Also**

`post.prob()`, `cov.param()`, `cov.post.z()`, `cov.logit.z()`.

**Examples**

```
library(MixfMRI, quietly = TRUE)
.FC.CT$model.X <- "I"
.FC.CT$CONTROL$debug <- 0
K <- 3

### Fit toy1.
set.seed(1234)
X.gbd <- toy1$X.gbd
X.range <- apply(X.gbd, 2, range)
X.gbd <- t((t(X.gbd) - X.range[1,]) / (X.range[2,] - X.range[1,]))
PV.gbd <- toy1$PV.gbd
fcobj <- fclust(X.gbd, PV.gbd, K = K, min.1st.prop = 0.5)

### Test log odds ratio.
x <- list(X.gbd = X.gbd, PV.gbd = PV.gbd)
post.z <- post.prob(x, fcobj)
lor <- logor.stat(x, fcobj, post.z)

### Check if 95% CE covers log odd ratio = 1.
id <- !is.na(lor$df)
id.cover.0 <- which(lor$test.stat[id] < pchisq(0.95, lor$df[id]))

### Get voxels needed for merging.
id.active <- which(fcobj$class != 1)
id.merge <- id.active[id][id.cover.0]

### Check results.
post.z[id.merge,]
cbind(toy1$X.gbd[id.merge,], toy1$PV.gbd[id.merge])
```

**Description**

These functions compute posterior probabilities, Fisher information with covariance matrix of parameters, covariance matrix of posterior probabilities, and covariance matrix of logit posterior probabilities.

**Usage**

```
post.prob(x, fcobj)
cov.param(x, fcobj, post.z, drop.ETA1 = FALSE)
cov.post.z(x, fcobj, post.z, cov.param = NULL, all.x = FALSE,
           drop.ETA1 = FALSE)
cov.logit.z(x, fcobj, post.z, cov.param = NULL, cov.post.z = NULL,
            all.x = FALSE, drop.ETA1 = FALSE)
```

**Arguments**

<code>x</code>	an input list of two elements <code>X.gbd</code> and <code>PV.gbd</code> .
<code>fcobj</code>	a <code>fclust</code> object.
<code>post.z</code>	a matrix of <code>dim = N * K</code> for posterior probabilities, which is also the return value of <code>post.prob()</code> .
<code>cov.param</code>	a covariance matrix of <code>dim = d * d</code> for parameters, which is also a return of <code>cov.param()</code> . <code>d</code> is total number of parameters which is dependent on data and models.
<code>cov.post.z</code>	a covariance list of length equal to number of active voxels, which is also a return of <code>cov.post.z()</code> .
<code>all.x</code>	all cov matrices for all observations are returned if <code>TRUE</code> , while for only active observations (those of class ids are greater than 1) if <code>FALSE</code> .
<code>drop.ETA1</code>	if drop the <code>ETA[1]</code> from the cov matrix due to the <code>min.1st.prop</code> constrain.

**Details**

These functions are required to compute covariance matrices of parameters and posterior probabilities.

Use `post.prob()` to get the posterior probabilities.

Input the returns of `post.prob()` to `cov.param()` to obtain the cov matrix for parameters (inversed Fisher information obtained from inner product of gradient of log observed data likelihood). A list is returned with `I` for Fisher information, and `cov` for the covariance matrix which is inverted by `ginv()`.

Input the returns of `post.prob()` and `cov.param()` to `cov.post.z()` to obtain the cov matrix for posterior probabilities by the multivariate delta method on the cov matrix for parameters.

Input the returns of `post.prob()`, `cov.param()`, and `cov.post.z()` to `cov.logit.z()` to obtain cov matrix for logit posterior probabilities by the multivariate delta method on cov matrix of posterior probabilities.



**Value**

A matrix or a list is returned.

The `cov.param()` will return a list containing two elements `I` for the Fisher information, and `cov` for the covariance matrix by generalized inversed of the Fisher information. The dimension of both elements are  $d * d$  where  $d = K * 7 - 4$  for 2D data and  $d = K * 9 - 4$  for 3D data if `drop.ETA1 = TRUE`, otherwise they are  $d = K * 7 - 3$  and  $d = K * 9 - 4$ , respectively.

The `cov.post.z()` will return a list containing cov matrices of posterior probabilities for each valid/selected voxel.

The `cov.logit.z()` will return a list containing cov matrices of logit posterior probabilities for each valid/selected voxel.

**Author(s)**

Wei-Chen Chen and Ranjan Maitra.

**References**

Chen, W.-C. and Maitra, R. (2021) "A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies", *arXiv:2102.03639*.

**See Also**

`EMCluster::lmt()`, `lmt.I()`.

**Examples**

```
library(MixfMRI, quietly = TRUE)
library(EMCluster, quietly = TRUE)
.FC.CT$model.X <- "I"
.FC.CT$CONTROL$debug <- 0
K <- 3

.rem <- function(){

  ### Fit toy1.
  set.seed(1234)
  X.gbd <- toy1$X.gbd
  X.range <- apply(X.gbd, 2, range)
  X.gbd <- t((t(X.gbd) - X.range[1,]) / (X.range[2,] - X.range[1,]))
  PV.gbd <- toy1$PV.gbd
  fcobj <- fclust(X.gbd, PV.gbd, K = K, min.1st.prop = 0.5)

  ### Test cov matrix of posterior z and logit posterior z.
  x <- list(X.gbd = X.gbd, PV.gbd = PV.gbd)
  post.z <- post.prob(x, fcobj)
  cov.param <- cov.param(x, fcobj, post.z = post.z)
  cov.post.z <- cov.post.z(x, fcobj, post.z = post.z,
                           cov.param = cov.param$cov)
```

```

cov.logit.z <- cov.logit.z(x, fcobj, post.z = post.z,
                          cov.param = cov.param$cov,
                          cov.post.z = cov.post.z)

### Compute cov matrix of log odds ratio for all k > 1.
A <- cbind(rep(-1, K - 1), diag(1, K - 1))
logit.p <- log(post.z[fcobj$class != 1,] / (1 - post.z[fcobj$class != 1,]))
log.or <- logit.p %>% t(A)
cov.log.or <- lapply(cov.logit.z, function(x) A %>% x %>% t(A))

### Check if 0 vector covered by 95% confidence ellipsoid.
id <- 1
plot(log.or[id,],
      xlim = log.or[id, 1] + c(-5, 5),
      ylim = log.or[id, 2] + c(-5, 5),
      main = "1st observation", xlab = "x", ylab = "y")
plotBN(log.or[id,], cov.log.or[[id]])
points(0, 0, col = 2)
}

```

---

Covariance Matrices of Logit ETA

*Covariance Matrices of Logit ETA*

---

## Description

These functions computes covariance matrix of logit ETA.

## Usage

```
cov.logit.ETA(x, fcobj, cov.param = NULL)
```

## Arguments

<code>x</code>	an input list of two elements <code>X.gbd</code> and <code>PV.gbd</code> .
<code>fcobj</code>	a <code>fclust</code> object.
<code>cov.param</code>	a covariance matrix of $\text{dim} = d * d$ for parameters, which is also a return of <code>cov.param()</code> . <code>d</code> is total number of parameters which is dependent on data and models.

## Details

These functions are required to compute covariance matrices of logit ETA.

Input the returns of `cov.param()` to `cov.logit.ETA()` to obtain the cov matrix for logit ETA by the multivariate delta method on the cov matrix for parameters.

**Value**

A matrix.

**Author(s)**

Wei-Chen Chen and Ranjan Maitra.

**References**

Chen, W.-C. and Maitra, R. (2021) “A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies”, *arXiv:2102.03639*.

**See Also**

EMCluster::lmt(), lmt.I().

**Examples**

```
library(MixfMRI, quietly = TRUE)
.FC.CT$model.X <- "I"
.FC.CT$CONTROL$debug <- 0
K <- 3

.rem <- function(){

  ### Fit toy1.
  set.seed(1234)
  X.gbd <- toy1$X.gbd
  X.range <- apply(X.gbd, 2, range)
  X.gbd <- t((t(X.gbd) - X.range[1,]) / (X.range[2,] - X.range[1,]))
  PV.gbd <- toy1$PV.gbd
  fcobj <- fclust(X.gbd, PV.gbd, K = K, min.1st.prop = 0.5)

  ### Test cov matrix of posterior z.
  x <- list(X.gbd = X.gbd, PV.gbd = PV.gbd)
  post.z <- post.prob(x, fcobj)
  cov.param <- cov.param(x, fcobj, post.z)
  cov.logit.ETA <- cov.logit.ETA(x, fcobj, cov.param = cov.param$cov)

  ### Compute cov matrix of eta_k - eta_1 for all k > 1.
  A <- cbind(rep(-1, K - 1), diag(1, K - 1))
  ETA <- fcobj$param$ETA
  log.or <- log(ETA / (1 - ETA)) %*% t(A)
  cov.log.or <- A %*% cov.logit.ETA %*% t(A)

}
```

---

Density function of p-values

*Density function of p-values*

---

### Description

These functions based on normal assumption and transformation to derive a (mixture) density function of p-values.

### Usage

```
dpval(x, mu = 0, log = FALSE)
dmixpval(x, eta, mu)
```

### Arguments

x	support of p-values which should be between 0 and 1.
mu	hypothetical mean of testing statistics (in normal distribution) for producing p-values.
log	if return log of density.
eta	mixing proportion of K components if a mixture is assumed.

### Details

Note that eta and mu in `dmixpval()` are of length K for K component mixtures.

### Value

Corresponding density values (to the input x) are returned.

### Author(s)

Wei-Chen Chen and Ranjan Maitra.

### References

Chen, W.-C. and Maitra, R. (2021) "A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies", *arXiv:2102.03639*.

### See Also

[gendataset\(\)](#), [qvalue\(\)](#).

**Examples**

```
library(MixfMRI, quietly = TRUE)
set.seed(1234)
da <- gendataset(phantom = shepp1fMRI, overlap = 0.01)
x <- da$pval[!is.na(da$pval)][1:100]
dpval(x)
dmixpval(x, mu = da$mu, eta = da$eta)
```

---

Example Datasets

*Example datasets in MixfMRI*

---

**Description**

These are datasets used to demo examples and workflows in this package.

**Format**

Objects may contain several information and data.

**Details**

pstats is a 3D example.

pval.2d.complex and pval.2d.mag are 2D examples.

shepp0fMRI, shepp1fMRI, shepp2fMRI and sheppAnat are phantoms generated by Dr. Maitra for simulation studies with different overlap levels for p-values.

toy\* are 2D toy examples.

**Author(s)**

Wei-Chen Chen and Ranjan Maitra.

**References**

Chen, W.-C. and Maitra, R. (2021) “A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies”, *arXiv:2102.03639*.

**Examples**

```
library(MixfMRI, quietly = TRUE)

### Plotting.
demo(shepp, 'MixfMRI', ask=FALSE, echo=FALSE)
```

---

**False Discovery Rates for Spatial Signals**

*False Discovery Rates for Spatial Signals using Benjamini and Heller (2007)*

---

**Description**

Compute q-values Benjamini and Heller's (2007) approach for controlling FDR for spatial signals.

**Usage**

```
fdr.bh.p1(p, w = rep(1, length(p)), q = 0.05)
fdr.bh.p2(p, w = rep(1, length(p)), q = 0.05)
```

**Arguments**

p	a p-value vector. No NA is allowed and all values are in [0, 1].
w	a weight vector for p-values.
q	a desired cutoff for adjusting p-values.

**Details**

These functions implement first two procedures in Benjamini and Heller (2007) for controlling FDR for spatial signals.

**Value**

Return the number of rejected hypotheses and all corresponding q-values for the input p-values.

**Author(s)**

Wei-Chen Chen.

**References**

Chen, W.-C. and Maitra, R. (2021) "A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies", *arXiv:2102.03639*.

**See Also**

[qvalue\(\)](#).

**Examples**

```
library(MixfMRI, quietly = TRUE)
set.seed(1234)
da <- gendataset(phantom = shepp1fMRI, overlap = 0.01)
p <- da$pval[!is.na(da$pval)][1:100]
fdr.bh.p1(p)
fdr.bh.p2(p)
```

---

Generalized Cluster-Based Analysis (CBA) Method

*Generalized Cluster-Based Analysis (CBA) Method*

---

**Description**

Find clusters in 2D or 3D based on a generalized CBA method. The CBA method is originally proposed by Heller, et.al. (2006) using the correlation of two time series as the similarity of two spatial locations.

**Usage**

```
cba.cor(da.ts, da.m = NULL, adj.dist = TRUE, fun.sim = stats::cor)
cba.cor.2d(da.ts, da.m = NULL, adj.dist = TRUE, fun.sim = stats::cor)
cba.cor.3d(da.ts, da.m = NULL, adj.dist = TRUE, fun.sim = stats::cor)
```

**Arguments**

<code>da.ts</code>	a time series array of dimensions $x * y * z * t$ .
<code>da.m</code>	a mask determining inside of brain or not.
<code>adj.dist</code>	if adjust correlations by distance.
<code>fun.sim</code>	a function computing similarity of two locations.

**Details**

These functions implement the 2D and 3D versions of CBA proposed by Heller, et.al. (2006).

`da.ts` should have dimensions  $x * y * z * t$  for 3D data and  $x * y * t$  for 2D data. Similarly, `da.m` would have  $x * y * z$  and  $x * y$  correspondingly.

`da.m` has values 0 or 1 indicating outside or inside a brain, respectively.

`fun.sim(a, B)` is a function return similarity between a location `a` and `N` neighboring locations `B` where `a` is of dimension  $t * 1$  and `B` is of dimension  $t * N$ . Ideally, `fun.sim()` should return values of similarity which take values between 0 and 1 where 0 means totally different and 1 means completely identical of two spatial locations. By default, `stats::cor` is used. See the example section next for user defined functions for `fun.sim()`.

**Value**

Return the cluster ids for each voxel. NA for outside of brain if `da.m` is provided.

**Author(s)**

Wei-Chen Chen.

**References**

- Heller, et.al. (2006) “Cluster-based analysis of FMRI data”, *NeuroImage*, 33(2), 599-608.
- Chen, W.-C. and Maitra, R. (2021) “A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies”, *arXiv:2102.03639*.

**See Also**

[fdr.bh.p1\(\)](#), [fdr.bh.p2\(\)](#).

**Examples**

```
### Simulated data
library(MixfMRI, quietly = TRUE)
dim <- c(4, 5, 4, 10)
set.seed(123)
da.ts <- array(rnorm(prod(dim)), dim = dim)
id.class <- suppressWarnings(cba.cor(da.ts))
table(id.class)

fun.tanh <- function(a, B){
  d <- 1 / apply(B, 2, function(b){ dist(rbind(as.vector(a), b)) })
  tanh(d)
}
id.class.tanh <- suppressWarnings(cba.cor(da.ts, fun.sim = fun.tanh))
table(id.class.tanh)

fun.logit <- function(a, B){
  d <- dist(t(cbind(a, B)))[1:ncol(B)]
  (1 / (1 + exp(-d))) * 2 - 1
}
id.class.logit <- suppressWarnings(cba.cor(da.ts, fun.sim = fun.logit))
table(id.class.logit)

.rem <- function(){

### Real data
library(AnalyzeFMRI, quietly = TRUE)
library(oro.nifti, quietly = TRUE)

fn <- "pb02_volreg_tlrc.nii"
da <- readNIFTI(fn)
da.ts <- da@.Data

fn <- "mask_anat.nii"
da <- readNIFTI(fn)
```



```
da.m <- da@.Data

id.class <- suppressWarnings(cba.cor(da.ts, da.m))
dim(id.class) <- dim(da.m)
length(table(id.class))

}
```

---

**initial***Main initialization functions*

---

**Description**

Main initialization functions.

**Usage**

```
initial.em.gbd(PARAM)
```

```
initial.RndEM.gbd(PARAM)
```

**Arguments**

PARAM            a list of uninitialized parameters, as usual, the returned values of `set.global()`, to be initialized according to data (inside PARAM).

**Details**

`initial.em.gbd()` takes in a template of PARAM (uninitialized), and usually is available by calling `set.global()`, then return an initialized PARAM which is ready for EM runs.

Internally, there are six different initializations implemented for the function `initial.em.gbd()` including `prob.extend`, `prob.simple`, `qnorm.extend`, `qnorm.simple`, `extend`, and `simple`. These methods are mainly based on transformation of original space of data (p-values and voxel locations) into more linear space such that the Euclidean distance more makes sense (fairly) to classify data in groups.

`initial.RndEM.gbd()` implement RndEM initialization algorithm based on repeated calling `initial.em.gbd()`.

Note that all configurations are included in PARAM set by `set.global()`.

**Value**

These functions return an initialized PARAM for EM runs based on pre-stored configuration within the input uninitialized PARAM.

**Author(s)**

Wei-Chen Chen and Ranjan Maitra.

## References

Chen, W.-C. and Maitra, R. (2021) “A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies”, *arXiv:2102.03639*.

## See Also

[set.global\(\)](#), [fclust\(\)](#), [PARAM](#).

## Examples

```
library(MixfMRI, quietly = TRUE)
library(EMcluster, quietly = TRUE)
# .FC.CT$algorithm <- "em"
# .FC.CT$model.X <- "v"
# .FC.CT$ignore.X <- TRUE
.FC.CT$check.X.unit <- FALSE

### Test toy1.
set.seed(1234)
X.gbd <- toy1$X.gbd
PV.gbd <- toy1$PV.gbd
PARAM <- set.global(X.gbd, PV.gbd, K = 2)
PARAM.new <- initial.em.gbd(PARAM)
PARAM.toy1 <- em.step.gbd(PARAM.new)
id.toy1 <- .MixfMRIEnv$CLASS.gbd
print(PARAM.toy1$ETA)
RRand(toy1$CLASS.gbd, id.toy1)

.rem <- function(){

  ### Test toy2.
  set.seed(1234)
  X.gbd <- toy2$X.gbd
  PV.gbd <- toy2$PV.gbd
  PARAM <- set.global(X.gbd, PV.gbd, K = 3)
  PARAM.new <- initial.em.gbd(PARAM)
  PARAM.toy2 <- em.step.gbd(PARAM.new)
  id.toy2 <- .MixfMRIEnv$CLASS.gbd
  print(PARAM.toy2$ETA)
  RRand(toy2$CLASS.gbd, id.toy2)

}
```

**Description**

These functions test two mixture Gaussian fMRI models with identity covariance matrices and different numbers of clusters. These functions are similar to the `EMCluster::lmt()`, but is coded for fMRI models in **MixfMRI**.

**Usage**

```
lmt.I(fcobj.0, fcobj.a, X.gbd, PV.gbd, tau = 0.5, n.mc.E.delta = 1000,
      n.mc.E.chi2 = 1000, verbose = FALSE)
lmt.pv(fcobj.0, fcobj.a, X.gbd, PV.gbd, tau = 0.5, n.mc.E.delta = 1000,
       n.mc.E.chi2 = 1000, verbose = FALSE)
```

**Arguments**

<code>fcobj.0</code>	a <code>fclust</code> object for the null hypothesis.
<code>fcobj.a</code>	a <code>fclust</code> object for the alternative hypothesis.
<code>X.gbd</code>	a data matrix of $N$ voxel locations. $\dim(X.gbd) = N \times 3$ for 3D data and $N \times 2$ for 2D data.
<code>PV.gbd</code>	a p-value vector of signals associated with voxels. $\text{length}(PV.gbd) = N$ .
<code>tau</code>	proportion of null and alternative hypotheses.
<code>n.mc.E.delta</code>	number of Monte Carlo simulations for expectation of delta (difference of logL).
<code>n.mc.E.chi2</code>	number of Monte Carlo simulations for expectation of chisquare statistics.
<code>verbose</code>	if verbose.

**Details**

This function calls several subroutines to compute information, likelihood ratio statistics, degrees of freedom, non-centrality of chi-squared distributions ...etc. Based on Monte Carlo methods to estimate parameters of likelihood mixture tests, this function return a p-value for testing  $H_0: fcobj.0$  v.s.  $H_a: fcobj.a$ .

`lmt.pv()` only uses `PV.gbd`.

**Value**

A list of class `lmt.I` are returned.

**Author(s)**

Wei-Chen Chen and Ranjan Maitra.

**References**

Chen, W.-C. and Maitra, R. (2021) "A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies", *arXiv:2102.03639*.

**See Also**

EMCluster::lmt().

**Examples**

```
library(MixfMRI, quietly = TRUE)
library(EMCluster, quietly = TRUE)
.FC.CT$model.X <- "I"
.FC.CT$check.X.unit <- FALSE
.FC.CT$CONTROL$debug <- 0

.rem <- function(){

  ### Fit toy1.
  set.seed(1234)
  X.gbd <- toy1$X.gbd
  PV.gbd <- toy1$PV.gbd
  ret.2 <- fclust(X.gbd, PV.gbd, K = 2)
  ret.3 <- fclust(X.gbd, PV.gbd, K = 3)
  ret.4 <- fclust(X.gbd, PV.gbd, K = 4)
  ret.5 <- fclust(X.gbd, PV.gbd, K = 5)

  ### ARI
  RRand(toy1$CLASS.gbd, ret.2$class)
  RRand(toy1$CLASS.gbd, ret.3$class)
  RRand(toy1$CLASS.gbd, ret.4$class)
  RRand(toy1$CLASS.gbd, ret.5$class)

  ### Test toy1.
  (lmt.23 <- lmt.I(ret.2, ret.3, X.gbd, PV.gbd))
  (lmt.24 <- lmt.I(ret.2, ret.4, X.gbd, PV.gbd))
  (lmt.25 <- lmt.I(ret.2, ret.5, X.gbd, PV.gbd))
  (lmt.34 <- lmt.I(ret.3, ret.4, X.gbd, PV.gbd))
  (lmt.35 <- lmt.I(ret.3, ret.5, X.gbd, PV.gbd))
  (lmt.45 <- lmt.I(ret.4, ret.5, X.gbd, PV.gbd))

  ### Test toy1 using p-values only.
  (lmt.pv.23 <- lmt.pv(ret.2, ret.3, X.gbd, PV.gbd))
  (lmt.pv.24 <- lmt.pv(ret.2, ret.4, X.gbd, PV.gbd))
  (lmt.pv.25 <- lmt.pv(ret.2, ret.5, X.gbd, PV.gbd))
  (lmt.pv.34 <- lmt.pv(ret.3, ret.4, X.gbd, PV.gbd))
  (lmt.pv.35 <- lmt.pv(ret.3, ret.5, X.gbd, PV.gbd))
  (lmt.pv.45 <- lmt.pv(ret.4, ret.5, X.gbd, PV.gbd))

}
```

**Description**

Likelihood ratio tests for merging clusters.

**Usage**

```
lrt(PV.gbd, CLASS.gbd, K, H0.alpha = .FC.CT$LRT$H0.alpha,
    H0.beta = .FC.CT$LRT$H0.beta)

lrt2(PV.gbd, CLASS.gbd, K, H0.mean = .FC.CT$LRT$H0.mean,
     upper.beta = .FC.CT$INIT$BETA.beta.max, proc = c("1", "2", "weight"))

lrt.betamean(PV.gbd, CLASS.gbd, K, proc = c("1", "2"))

lrt.betaab(PV.gbd, CLASS.gbd, K, proc = c("1", "2"))
```

**Arguments**

PV.gbd	a p-value vector of signals associated with voxels. $\text{length}(\text{PV.gbd}) = N$ .
CLASS.gbd	a classification vector of signals associated with voxels. $\text{length}(\text{CLASS.gbd}) = N$ .
K	number of clusters.
H0.alpha	null hypothesis for the alpha parameter of Beta distribution.
H0.beta	null hypothesis for the beta parameter of Beta distribution.
H0.mean	null hypothesis for the mean of Beta distribution.
upper.beta	BETA.beta.max, maximum value of beta parameter of Beta distribution.
proc	q-value procedure for adjusting p-values.

**Details**

These functions perform likelihood ratio tests for merging clusters. Only p-values coordinates (Beta density) are tested, while voxel location coordinates (multivariate Normal density) are not involved in testing.

`lrt.betamean` tests if means of any two pairs of mixture (p-value) component were the same. The chi-square distribution with 1 degree of freedom is used.

`lrt.betaab` tests if alpha and beta of any two pairs of mixture (p-value) components were the same. The chi-square distribution with 2 degrees of freedom is used.

**Value**

A matrix contains MLEs of parameters of Beta distribution under the null hypothesis and the union of null and alternative hypotheses. The matrix also contains testing statistics and p-values.

**Author(s)**

Wei-Chen Chen and Ranjan Maitra.

## References

Chen, W.-C. and Maitra, R. (2021) “A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies”, *arXiv:2102.03639*.

## See Also

[PARAM](#).

## Examples

```
library(MixfMRI, quietly = TRUE)
set.seed(1234)

### Test 2d data.
da <- pval.2d.mag
id <- !is.na(da)
PV.gbd <- da[id]
id.loc <- which(id, arr.ind = TRUE)
X.gbd <- t(t(id.loc) / dim(da))
ret <- fclust(X.gbd, PV.gbd, K = 2, min.1st.prop = 0.95)
# print(ret)

### p-values of rest clusters.
ret.lrt <- lrt(PV.gbd, ret$class, K = 2)
print(ret.lrt)

.rem <- function(){

  ret.lrt2 <- lrt2(PV.gbd, ret$class, K = 3)
  print(ret.lrt2)

}
```

---

Main functions

*Main MixfMRI function*

---

## Description

Main MixfMRI functions.

## Usage

```
fclust(X.gbd, PV.gbd, K = 2,
       PARAM.init = NULL,
       min.1st.prop = .FC.CT$INIT$min.1st.prop,
       max.PV = .FC.CT$INIT$max.PV,
       class.method = .FC.CT$INIT$class.method[1],
```

```

RndEM.iter = .FC.CT$CONTROL$RndEM.iter,
algorithm = .FC.CT$algorithm[1],
model.X = .FC.CT$model.X[1],
ignore.X = .FC.CT$ignore.X,
stop.unstable = TRUE,
MPI.gbd = .FC.CT$MPI.gbd, common.gbd = .FC.CT$common.gbd)

set.global(X.gbd, PV.gbd, K = 2,
min.1st.prop = .FC.CT$INIT$min.1st.prop,
max.PV = .FC.CT$INIT$max.PV,
class.method = .FC.CT$INIT$class.method[1],
RndEM.iter = .FC.CT$CONTROL$RndEM.iter,
algorithm = .FC.CT$algorithm[1],
model.X = .FC.CT$model.X[1],
ignore.X = .FC.CT$ignore.X,
check.X.unit = .FC.CT$check.X.unit,
MPI.gbd = .FC.CT$MPI.gbd, common.gbd = .FC.CT$common.gbd)

```

### Arguments

<code>X.gbd</code>	a data matrix of N voxel locations. $\dim(X.gbd) = N \times 3$ for 3D data and $N \times 2$ for 2D data.
<code>PV.gbd</code>	a p-value vector of signals associated with voxels. $\text{length}(PV.gbd) = N$ .
<code>K</code>	number of clusters to be estimated.
<code>PARAM.init</code>	initial parameters.
<code>min.1st.prop</code>	lower bound of mixing proportion (ETA) of the 1st cluster (uniform).
<code>max.PV</code>	upper bound of p-values where initializations pick from.
<code>class.method</code>	classification method for initializations.
<code>RndEM.iter</code>	number of RndEM iterations.
<code>algorithm</code>	either “ecm” (ECM), “apecma” (APECMA) or “em” (EM) algorithm.
<code>model.X</code>	either “I” or “V” for covariance matrix.
<code>ignore.X</code>	if <code>X.gbd</code> used in model, TRUE for <code>PV.gbd</code> only.
<code>check.X.unit</code>	if <code>X.gbd</code> are all in $[0, 1]$ .
<code>stop.unstable</code>	if <code>fclust</code> stops if unstable results occur.
<code>MPI.gbd</code>	if MPI (“EGM” algorithm) is used.
<code>common.gbd</code>	if <code>X.gbd</code> and <code>PV.gbd</code> are in common across all ranks when <code>MPI.gbd = TRUE</code> .

### Details

The `fclust()` contains initialization and EM algorithms for clustering fMRI signal data which have two parts: `X.gbd` for voxel information either 2D or 3D, `PV.gbd` for p-value of signals associated with voxels. Each signal is assumed as a mixture distribution with `K` components with mixing proportion `ETA`, and each component has two independent coordinates with density functions: Beta and multivariate Normal distributions.

Beta density: The 1st component is restricted by `min.1st.prop` and Beta(1, 1) distribution. The other  $K - 1$  components have Beta(alpha, beta) distribution with  $\alpha < 1 < \beta$ .

Multivariate Normal density: `model.X = "I"` is for identity cov matrix of multivariate Normal distribution, and "V" for unstructured cov matrix. `ignore.X = TRUE` is to ignore `X.gbd` and normal density, i.e. only Beta density is used.

Currently, APECMa and EM algorithms are implemented with EGM algorithm to speed up convergence if MPI is available. RndEM initialization is also implemented for better chance of good initial values for convergence.

The `set.global()` has purposes: create a template/storage of parameters, save configurations, and called by `fclust()` to initial the parameters, such as `initial.em.gbd()` or `initial.RndEM.gbd()`.

### Value

A list with class `fclust` by `fclust()` is returned which can be summarized by `print.fclust()`.

A list `PARAM` or `PARAM.org` is returned by `set.global()`:

<code>N.gbd</code>	number of observations (within the rank), and should be equal to <code>N.all</code> if <code>MPI.gbd = FALSE</code> .
<code>N.all</code>	numbers of observations (of all ranks if <code>MPI.gbd = TRUE</code> ).
<code>N</code>	total number of observations ( <code>sum(N.all)</code> ).
<code>p</code>	dimension of an observation (3 for 2D signals, 4 for 3D signals), equivalent to total number of coordinates.
<code>p.X</code>	dimension of <code>X.gbd</code> (2 for 2D signals, 3 for 3D signals, 0 when <code>ignore.X = TRUE</code> , number of voxel coordinates.
<code>K</code>	number of clusters.
<code>ETA</code>	mixing proportion, length <code>K</code> .
<code>log.ETA</code>	<code>log(ETA)</code> .
<code>BETA</code>	a list of length <code>K</code> containing parameters (alpha, beta) of Beta density.
<code>MU</code>	a matrix of dimension <code>p.X</code> by <code>K</code> .
<code>SIGMA</code>	a list of length <code>K</code> , and each is of dimension <code>K x K</code> .
<code>logL</code>	log likelihood value.
<code>min.1st.prop</code>	carried from input.
<code>max.PV</code>	carried from input.
<code>class.method</code>	classification method of initializations.
<code>min.N.CLASS</code>	<code>p + 1</code> .
<code>model.X</code>	carried from input.

### Author(s)

Wei-Chen Chen and Ranjan Maitra.



## References

Chen, W.-C. and Maitra, R. (2021) “A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies”, *arXiv:2102.03639*.

## See Also

[print.fclust\(\)](#).

## Examples

```
library(MixfMRI, quietly = TRUE)
library(EMCluster, quietly = TRUE)
# .FC.CT$algorithm <- "em"
# .FC.CT$model.X <- "V"
# .FC.CT$ignore.X <- TRUE
.FC.CT$check.X.unit <- FALSE
set.seed(1234)

### Test toy1.
X.gbd <- toy1$X.gbd[, -3]
PV.gbd <- toy1$PV.gbd
PARAM <- fclust(X.gbd, PV.gbd, K = 2)
print(PARAM)
id.toy1 <- .MixfMRIEnv$CLASS.gbd
print(RRand(toy1$CLASS.gbd, id.toy1))

.rem <- function(){

  ### Test toy2.
  X.gbd <- toy2$X.gbd[, -3]
  PV.gbd <- toy2$PV.gbd
  PARAM <- fclust(X.gbd, PV.gbd, K = 3)
  print(PARAM)
  id.toy2 <- .MixfMRIEnv$CLASS.gbd
  print(RRand(toy2$CLASS.gbd, id.toy2))

}
```

## Description

These sets of controls are used to provide default values in this package.

## Format

Objects contain several parameters for methods.

## Details

The elements of `.FC.CT` are default values for main controls of **MixfMRI** including

Elements	Default	Usage
<code>algorithm</code>	"apecma"	implemented algorithm
<code>optim.method</code>	"BFGS"	optimization method
<code>model.X</code>	"I"	cov matrix structure
<code>ignore.X</code>	FALSE	if using voxel information
<code>check.X.unit</code>	TRUE	if checking X in $[0, 1]$
<code>CONTROL</code>	a list	see CONTROL next for details
<code>INIT</code>	a list	see INIT next for details
<code>LRT</code>	a list	see LRT next for details
<code>MPI.gbd</code>	FALSE	if MPI speedup available
<code>common.gbd</code>	TRUE	if X in common gbd format

The elements of `CONTROL` are default values for optimization controls of implemented EM algorithm including

Elements	Default	Usage
<code>max.iter</code>	1000	maximum number of EM iterations
<code>abs.err</code>	1e-4	absolute error of convergence
<code>rel.err</code>	1e-6	relative error of convergence
<code>debug</code>	1	debugging level
<code>RndEM.iter</code>	10	RndEM iterations
<code>exp.min</code>	<code>log(.Machine\$double.xmin)</code>	minimum exponential power
<code>exp.max</code>	<code>log(.Machine\$double.xmax)</code>	maximum exponential power
<code>sigma.ill</code>	1e-6	ill condition limit
<code>DS.max</code>	1e+4	maximum <code>chol()</code> cov matrix
<code>DS.min</code>	1e-6	minimum <code>chol()</code> cov matrix

The elements of `INIT` are default values or limitations for initial parameters implemented for EM algorithm including

Elements	Default	Usage
<code>min.1st.prop</code>	0.8	minimum proportion of 1st cluster
<code>max.PV</code>	0.1	maximum p-value for initialization
<code>BETA.alpha.min</code>	$0 + 1e-6$	minimum value of alpha parameter of Beta distribution
<code>BETA.alpha.max</code>	$1 - 1e-6$	maximum value of alpha parameter of Beta distribution
<code>BETA.beta.min</code>	$1 + 1e-6$	minimum value of beta parameter of Beta distribution
<code>BETA.beta.max</code>	1e+6	maximum value of beta parameter of Beta distribution
<code>max.try.iter</code>	10	maximum retry iterations if result is unstable
<code>class.method</code>	"prob.extned"	classification method at initializations

The elements of LRT are default values or limitations for likelihood ratio tests including

Elements	Default	Usage
<code>H0.alpha</code>	1	null hypothesis alpha parameter of Beta distribution
<code>H0.beta</code>	1	null hypothesis beta parameter of Beta distribution
<code>H0.mean</code>	0.05	null hypothesis mean of Beta distribution

### Author(s)

Wei-Chen Chen and Ranjan Maitra.

### References

Chen, W.-C. and Maitra, R. (2021) “A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies”, *arXiv:2102.03639*.

### See Also

[set.global\(\)](#), [fclust\(\)](#).

---

Plotting

*Main plotting function*

---

### Description

Main plotting function in **MixfMRI**.

### Usage

```
plotfclust(da, posterior, main = NULL, xlim = NULL, ylim = NULL)
plotfclustpv(da, posterior, main = NULL, xlim = NULL, ylim = NULL)

plotpv(da, posterior, PARAM, zlim = c(0, 0.01), plot.mean = TRUE,
        xlab = "", ylab = "", main = NULL, xlim = NULL, ylim = NULL,
        col = my.YlOrRd(), ignore.bg = FALSE)
plotpvlegend(zlim = c(0, 0.01), n.level = 20, main = NULL,
             col = my.YlOrRd())
```

**Arguments**

<code>da</code>	a data set to be plotted.
<code>posterior</code>	a posterior data set to be plotted.
<code>PARAM</code>	a returning parameter object from <code>fclust()</code> .
<code>main</code>	title of the plot.
<code>xlim</code>	limits of x-axis.
<code>ylim</code>	limits of y-axis.
<code>zlim</code>	limits of z-axis.
<code>xlab</code>	labels of x-axis.
<code>ylab</code>	labels of y-axis.
<code>plot.mean</code>	if plotting mean values of each cluster.
<code>col</code>	colors to be drawn.
<code>ignore.bg</code>	if ignoring the background.
<code>n.level</code>	number of levels to be plotted.

**Details**

These are example functions to plot results, simulations, and datasets.

**Value**

Return plots.

**Author(s)**

Wei-Chen Chen and Ranjan Maitra.

**References**

Chen, W.-C. and Maitra, R. (2021) “A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies”, *arXiv:2102.03639*.

**See Also**

`set.global()`.

**Examples**

```
library(MixfMRI, quietly = TRUE)
set.seed(1234)
```

```
.rem <- function(){
  ### Check 2d data.
```

```

da <- pval.2d.complex
id <- !is.na(da)
PV.gbd <- da[id]
hist(PV.gbd, nclass = 100, main = "p-value")

### Test 2d data.
id.loc <- which(id, arr.ind = TRUE)
X.gbd <- t(t(id.loc) / dim(da))
ret <- fclust(X.gbd, PV.gbd, K = 3)
print(ret)

### p-values of rest clusters.
ret.lrt <- lrt(PV.gbd, ret$class, K = 3)
print(ret.lrt)
ret.lrt2 <- lrt2(PV.gbd, ret$class, K = 3)
print(ret.lrt2)

### Plotting.
par(mfrow = c(2, 2), mar = c(0, 0, 2, 0))
plotpv(da, ret$posterior, ret$param,
       zlim = c(0.005, 0.008), main = "Mean of Beta Distribution")
plotpv(da, ret$posterior, ret$param,
       plot.mean = FALSE, main = "p-value")
par(mar = c(5.1, 4.1, 4.1, 2.1))
plotpvlegend(zlim = c(0.005, 0.008), main = "Mean of Beta Distribution")
plotpvlegend(zlim = c(0, 0.01), main = "p-value")

}

```

---

Print Objects

*Print fclust related outputs*


---

## Description

Print fclust related outputs.

## Usage

```
## S3 method for class 'fclust'
print(x, ...)
```

## Arguments

x                    an object with the class attributes.  
...                   other arguments to the print function.

## Details

x is the return result from `fclust()`.

**Value**

A summary of `fclust` object is printed.

**Author(s)**

Wei-Chen Chen and Ranjan Maitra.

**References**

Chen, W.-C. and Maitra, R. (2021) “A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies”, *arXiv:2102.03639*.

**See Also**

`set.global()`, `fclust()`.

**Examples**

```
library(MixfMRI, quietly = TRUE)
set.seed(1234)

### Check 2d data.
da <- pval.2d.complex
id <- !is.na(da)
PV.gbd <- da[id]
# hist(PV.gbd, nclass = 100, main = "p-value")

### Test 2d data.
id.loc <- which(id, arr.ind = TRUE)
X.gbd <- t(t(id.loc) / dim(da))
ret <- fclust(X.gbd, PV.gbd, K = 2)
print(ret)
```

---

Simulations

*Generate datasets for MixfMRI simulations*

---

**Description**

Generate datasets for MixfMRI simulations

**Usage**

```
gendataset(phantom, overlap, smooth = FALSE)
```

**Arguments**

<code>phantom</code>	a phantom dataset.
<code>overlap</code>	a desired overlap level.
<code>smooth</code>	if <code>gcv.smooth2d()</code> be applied to the data.

## Details

This is a function to generate simulated fMRI data based on the input phantom and the desired overlap level for the fMRI p-value.

## Value

Return a list contains `eta` for mixing proportion, `overlap` for the desired level, `mu` for center of p-values, `class.id` for the true classifications where p-values belong to, `tval` for the testing statistics, and `pval` for the p-values of interesting in simulations.

## Author(s)

Wei-Chen Chen and Ranjan Maitra.

## References

Chen, W.-C. and Maitra, R. (2021) “A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies”, *arXiv:2102.03639*.

## See Also

[set.global\(\)](#).

## Examples

```
.rem <- function(){  
  
  library(MixfMRI, quietly = TRUE)  
  set.seed(1234)  
  da <- gendataset(phantom = shepp1fMRI, overlap = 0.01)$pval  
  da2 <- gendataset(phantom = shepp2fMRI, overlap = 0.01)$pval  
  
  par(mfrow = c(2, 2), mar = rep(0.05, 4))  
  image(shepp1fMRI[50:210, 50:210], axes = FALSE)  
  image(shepp2fMRI[50:210, 50:210], axes = FALSE)  
  image(da[50:210, 50:210], axes = FALSE)  
  image(da2[50:210, 50:210], axes = FALSE)  
  
}
```

---

Smoothing

*Generate datasets with smoothing for MixfMRI simulations*

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### **Description**

Generate datasets with smoothing for MixfMRI simulations

### **Usage**

```
gcv.smooth2d(y, interval)
```

### **Arguments**

y                    a set of p-values in 2d phantom  
interval            an interval for optimize function.

### **Details**

The function is used to smooth for Dr. Maitra's 2d phantom simulation. The smoothing method is based on Garcia (2010), CSDA.

### **Value**

Return a list containing two elements `im.smooth` and `par.val`.

### **Author(s)**

Ranjan Maitra.

### **References**

Chen, W.-C. and Maitra, R. (2021) "A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies", *arXiv:2102.03639*.

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Summarized Overlap

*Summarized Overlap*

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### **Description**

Compute summarized overlap on a given overlap (symmetric) matrix.

### **Usage**

```
summarized.overlap(overlap.mat)
```



**Arguments**

overlap.mat      an overlap (symmetric) matrix.

**Details**

overlap.mat is a  $p \times p$  matrix containing pair wise overlaps of  $p$  experiments. overlap.mat is assumed a symmetric matrix. This function returns a summarized overlap based on the input overlap.mat that characterizes the overall overlap behavior of the  $p$  experiments.

**Value**

A single value is returned.

**Author(s)**

Ranjan Maitra.

**References**

Chen, W.-C. and Maitra, R. (2021) “A Practical Model-based Segmentation Approach for Accurate Activation Detection in Single-Subject functional Magnetic Resonance Imaging Studies”, *arXiv:2102.03639*.

**Examples**

```
library(MixfMRI, quietly = TRUE)
set.seed(1234)
p <- 10 # 10 experiments.
overlap.mat <- diag(1, p)
overlap.mat[lower.tri(overlap.mat)] <- runif(p * (p - 1) / 2)
overlap.mat[upper.tri(overlap.mat)] <- t(overlap.mat)[upper.tri(overlap.mat)]
summarized.overlap(overlap.mat)
```

# Index

- \* **datasets**
  - Example Datasets, [13](#)
- \* **global variables**
  - MixfMRI Control, [25](#)
- \* **package**
  - MixfMRI-package, [2](#)
- \* **programming**
  - algorithm, [3](#)
  - Compute Q values, [5](#)
  - Compute Statistics for Log Odds Ratio of Posterior Probability, [6](#)
  - Covariance Matrices, [7](#)
  - Covariance Matrices of Logit ETA, [10](#)
  - Density function of p-values, [12](#)
  - False Discovery Rates for Spatial Signals, [14](#)
  - Generalized Cluster-Based Analysis (CBA) Method, [15](#)
  - initial, [17](#)
  - Likelihood Mixture Tests with Identity Cov Matrix or Only p-values, [18](#)
  - LRT, [20](#)
  - Main functions, [22](#)
  - Plotting, [27](#)
  - Print Objects, [29](#)
  - Simulations, [30](#)
  - Smoothing, [32](#)
  - Summarized Overlap, [32](#)
- .FC.CT (MixfMRI Control), [25](#)
  
- algorithm, [3](#)
- apexma.step.gbd (algorithm), [3](#)
  
- cba.cor (Generalized Cluster-Based Analysis (CBA) Method), [15](#)
- Compute Q values, [5](#)
  
- Compute Statistics for Log Odds Ratio of Posterior Probability, [6](#)
- cov.logit.ETA (Covariance Matrices of Logit ETA), [10](#)
- cov.logit.z (Covariance Matrices), [7](#)
- cov.param (Covariance Matrices), [7](#)
- cov.post.z (Covariance Matrices), [7](#)
- Covariance Matrices, [7](#)
- Covariance Matrices of Logit ETA, [10](#)
  
- Density function of p-values, [12](#)
- dmixpval, [5](#)
- dmixpval (Density function of p-values), [12](#)
- dpval, [5](#)
- dpval (Density function of p-values), [12](#)
  
- ecm.step.gbd (algorithm), [3](#)
- em.step.gbd (algorithm), [3](#)
- Example Datasets, [13](#)
  
- False Discovery Rates for Spatial Signals, [14](#)
- fclust, [2-4](#), [18](#), [27-30](#)
- fclust (Main functions), [22](#)
- fdr.bh.p1, [16](#)
- fdr.bh.p1 (False Discovery Rates for Spatial Signals), [14](#)
- fdr.bh.p2, [16](#)
- fdr.bh.p2 (False Discovery Rates for Spatial Signals), [14](#)
  
- gcv.smooth2d (Smoothing), [32](#)
- gendataset, [12](#)
- gendataset (Simulations), [30](#)
- Generalized Cluster-Based Analysis (CBA) Method, [15](#)
  
- initial, [17](#)
- initial.em.gbd, [3](#), [24](#)
- initial.RndEM.gbd, [3](#), [24](#)

- Likelihood Mixture Tests with Identity Cov Matrix or Only p-values, [18](#)
- lmt.I (Likelihood Mixture Tests with Identity Cov Matrix or Only p-values), [18](#)
- lmt.pv (Likelihood Mixture Tests with Identity Cov Matrix or Only p-values), [18](#)
- logor.stat (Compute Statistics for Log Odds Ratio of Posterior Probability), [6](#)
- LRT, [20](#)
- lrt (LRT), [20](#)
- lrt2 (LRT), [20](#)
  
- Main functions, [22](#)
- MixfMRI (MixfMRI-package), [2](#)
- MixfMRI Control, [25](#)
- MixfMRI-package, [2](#)
  
- PARAM, [4](#), [18](#), [22](#)
- PARAM (Main functions), [22](#)
- PARAM.org, [4](#)
- plotfclust (Plotting), [27](#)
- plotfclustpv (Plotting), [27](#)
- plotpv (Plotting), [27](#)
- plotpvlegend (Plotting), [27](#)
- Plotting, [27](#)
- post.prob (Covariance Matrices), [7](#)
- Print Objects, [29](#)
- print.fclust, [24](#), [25](#)
- print.fclust (Print Objects), [29](#)
- pstats (Example Datasets), [13](#)
- pval.2d.complex (Example Datasets), [13](#)
- pval.2d.mag (Example Datasets), [13](#)
  
- qvalue, [12](#), [14](#)
- qvalue (Compute Q values), [5](#)
  
- set.global, [3](#), [4](#), [17](#), [18](#), [27](#), [28](#), [30](#), [31](#)
- set.global (Main functions), [22](#)
- shepp0fMRI (Example Datasets), [13](#)
- shepp1fMRI (Example Datasets), [13](#)
- shepp2fMRI (Example Datasets), [13](#)
- sheppAnat (Example Datasets), [13](#)
- Simulations, [30](#)
- Smoothing, [32](#)
- Summarized Overlap, [32](#)
- summarized.overlap (Summarized Overlap), [32](#)
  
- toy1 (Example Datasets), [13](#)
- toy2 (Example Datasets), [13](#)