# Package 'MMOC'

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Title Multi-Omic Spectral Clustering using the Flag Manifold

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**Description** Multi-omic (or any multi-view) spectral clustering methods often assume the same number of clusters across all datasets. We supply methods for multi-omic spectral clustering when the number of distinct clusters differs among the omics profiles (views).

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**Encoding** UTF-8

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**Imports** Spectrum (>= 1.1), igraph (>= 1.4.1), MASS (>= 7.3-58.1)

**Suggests** knitr, rmarkdown, testthat (>= 3.0.0), SNFtool (>= 2.3.1), plotly (>= 4.10.0)

Config/testthat/edition 3

VignetteBuilder knitr

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**Repository** CRAN

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```
clustStruct
```

#### Description

Generates multiple data sets from a multivariate normal distribution using the mvrnorm function from the MASS package.

# Usage

clustStruct(n, p, k, noiseDat = "random", randNoise = 2)

#### Arguments

n	An integer, the sample size for all generated data sets
р	An integer, the number of columns (features) in each generated data set
k	An integer or vector, the number of distinct clusters in each generated data set. $n/k$ must be an integer for all values of k
noiseDat	Either the character string 'random', indicating the covariance matrix is a diag- onal matrix with randNoise along the diagonal, or a valid covariance matrix
randNoise	The value along the diagonal when noiseDat='random'

#### Details

The function accepts k as a vector. It splits data into k groups with means  $c(0, 2^{(1:(kk-1))})$ , e.g., when k=3 the data will be split into 3 groups with means 0, 2, and 4, respectively. The covariance matrix is either a diagonal matrix with randNoise (an integer) along the diagonal, or a given matrix.

# Value

A list of  $n \times p$  data frames with the specified number of groups

# Examples

```
## A single view with 30 variables and 3 groups
s1 <- clustStruct(n=120, p=30, k=3, noiseDat='random')[[1]]
## Multiple views with 30 variables
## View 1 has 2 groups and View 2 has 3 groups
s2 <- clustStruct(n=120, p=30, k=c(2,3), noiseDat='random')
## Multiple views with 30 variables
## View 1 has 2 groups, View 2 has 3, and View 3 has 3 groups
s3 <- clustStruct(n=120, p=30, k=c(2,3,3), noiseDat='random')
## Three view study.</pre>
```

#### flagMean

```
# View 1: 2 groups, 30 variables, random noise = 5
# View 2: 3 groups, 60 variables, random noise = 2
# View 3: 4 groups, 45 variables, random noise = 4
s4 <- clustStruct(n=120, k=c(2,3,4), p=c(30,60,45), randNoise=c(5,2,4))</pre>
```

```
flagMean
```

Calculate the Flag mean of multiple subspaces

# Description

Calculate the flag-mean of multiple subspaces. This method allows you to find the extrinsic mean of a finite set of subspaces. You can think of this as a median subspace. This method is also able to handle subspaces with different dimensions. See the references for more details

# Usage

flagMean(LapList, k, laplacian = c("shift", "Ng", "sym", "rw"), plots = TRUE)

#### Arguments

LapList	A list of Laplacian matrices
k	A vector indicating how many eigenvectors to take from each Laplacian, i.e., the number of clusters in each view
laplacian	One of "shift", "Ng", "rw" or "sym". Should be the same type used to calculate your Laplacians
plots	Whether or not to plot the singular values from SVD

# Details

Despite the complex linear algebra to achieve this result, the opperation is very simple. This function concatonates (cbind) the given subspaces and then performs singular value decomposition on the resulting matrix. This gives the 'median' subspace of the given set of subspaces. We would then cluster on the columns of the U matrix just as we do in standard spectral clustering

#### Value

The output from a singular value decomposition. See svd

#### References

https://www.semanticscholar.org/paper/Flag-Manifolds-for-the-Characterization-of-in-Large-Marrinan-Beveridge/7d306512b545df98243f87cb8173df83b4672b18 https://www.sciencedirect.com/science/article/pii/S0024379514

# Examples

```
## Generating data with 2 and 3 distinct clusters
## Note that 'clustStruct' returns a list
n=120; k <- c(2,3)
dd <- clustStruct(n=n, p=30, k=k, noiseDat='random')</pre>
## Laplacians
L_list <- lapply(dd, kernelLaplacian, kernel="Spectrum", plots=FALSE, verbose=FALSE)
## Calculating the flag mean
fm <- flagMean(L_list, k=k, laplacian='shift')</pre>
## Knowing the true structure makes it much easier to know how
## many right singular vectors to grab. There are 4 distinct
## groups in these data from 'clustStruct'
trueGroups(n=n, k=k)
kmeans(fm$u[, 1:4], centers=4)
```

kernelLaplacian Calculate the graph Laplacian of a given data set

# Description

Calculate the graph laplacian from a given data set with subjects as rows and features as columns.

# Usage

```
kernelLaplacian(
  dat.
  kernel = c("Gaussian", "ZM", "Spectrum", "Linear"),
 laplacian = c("shift", "Ng", "sym", "rw"),
 grf.type = c("full", "knn", "e-graph"),
 k = 5,
  p = 5,
  rho = NULL,
  epsilon = 0,
 mutual = FALSE,
 binary.grf = FALSE,
 plots = TRUE,
  verbose = TRUE
```

# )

#### Arguments

dat

A matrix like object with subjects as rows and features as columns.

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kernel	The type of kernel used to calculate the graph's adjacency matrix: "Gaussian" for the standard Gaussian kernel, "ZM" for the Zelnik-Manor kernel, "Spectrum" for the spectrum kernel, "Linear" for the linear kernel (dot product), and "Cor" for a kernel of pairwise correlations. See references for more details.
laplacian	One of "shift", "Ng", "rw" or "sym". See details for description
grf.type	Type of graph to calculate: "full" for adjacency matrix equal to the kernel, "knn" for a k-nearest neighbors graph, "e-graph" for an "epsilon graph"
k	An integer value for k in the k-nearest neighbors graph. Only the k largest edges (most similar neighbors) will be kept
р	An integer value for the p-nearest neighbor in the ZM kernel
rho	A value for the dispersion parameter in the Gaussian kernel. It is in the de- nominator of the exponent, so higher values correspond to lower similarity. By default it is the median pairwise Gaussian distance
epsilon	The cutoff value for the e-graph. Edges lower than this value will be removed
mutual	Make a "mutual" knn graph. Only keeps edges when two nodes are both in each others k-nearest set
binary.grf	Set all edges >0 to 1
plots	Whether or not to plot the final graph, a heatmap of calculated kernel, and the eigen values of the Laplacian
verbose	Whether or not to give some summary statistics of the pairwise distances

# Details

The four Lapalacians are defined as  $L_{shift} = I + D^{-1/2}AD^{-1/2}$ ,  $L_{Ng} = D^{-1/2}AD^{-1/2}$ ,  $L_{sym} = I - D^{-1/2}AD^{-1/2}$ , and  $L_{rw} = I - D^{-1}A$ . The shifted Laplacian,  $L_{shift} = I + D^{-1/2}AD^{-1/2}$ , is recommended for multi-view spectral clustering.

# Value

An  $n \times n$  matrix where n is the number of rows in dat.

#### References

https://academic.oup.com/bioinformatics/article/36/4/1159/5566508#199177546

# Examples

```
## Generating data with 3 distinct clusters
## Note that 'clustStruct' returns a list
dd <- clustStruct(n=120, p=30, k=3, noiseDat='random')[[1]]</pre>
```

kernelLaplacian(dd, kernel="Spectrum")

Laplacian

# Description

Calculate the graph laplacian from a given kernel matrix that represents the full graph weighted adjacency matrix

# Usage

```
Laplacian(
    A,
    laplacian = c("shift", "Ng", "sym", "rw"),
    grf.type = c("full", "knn", "e-graph"),
    k = 5,
    rho = NULL,
    epsilon = 0,
    mutual = FALSE,
    binary.grf = FALSE,
    plots = TRUE
)
```

# Arguments

A	An n by n kernel matrix, where n is the sample size, that represents your initial adjacency matrix. Kernel matrices are symmetric, positive semi-definite distance matrices
laplacian	One of "shift", "Ng", "rw" or "sym". See details for description
grf.type	Type of graph to calculate: "full" for adjacency matrix equal to the kernel, "knn" for a k-nearest neighbors graph, "e-graph" for an "epsilon graph"
k	An integer value for k in the k-nearest neighbors graph. Only the k largest edges (most similar neighbors) will be kept
rho	A value for the dispersion parameter in the Gaussian kernel. It is in the de- nominator of the exponent, so higher values correspond to lower similarity. By default it is the median pairwise Gaussian distance
epsilon	The cutoff value for the e-graph. Edges lower than this value will be removed
mutual	Make a "mutual" knn graph. Only keeps edges when two nodes are both in each others k-nearest set
binary.grf	Set all edges >0 to 1
plots	Whether or not to plot the final graph, a heatmap of calculated kernel, and the eigen values of the Laplacian

# Lapprox

#### Details

The four Lapalacians are defined as  $L_{shift} = I + D^{-1/2}AD^{-1/2}$ ,  $L_{Ng} = D^{-1/2}AD^{-1/2}$ ,  $L_{sym} = I - D^{-1/2}AD^{-1/2}$ , and  $L_{rw} = I - D^{-1}A$ . The shifted Laplacian,  $L_{shift} = I + D^{-1/2}AD^{-1/2}$ , is recommended for multi-view spectral clustering.

# Value

An  $n \times n$  matrix where n is the number of rows in dat.

#### References

https://academic.oup.com/bioinformatics/article/36/4/1159/5566508#199177546

#### Examples

```
## Generating data with 3 distinct clusters
## Note that 'clustStruct' returns a list
dd <- clustStruct(n=120, p=30, k=3, noiseDat='random')[[1]]
## Gaussian kernel
rho <- median(dist(dd))
A <- exp(-(1/rho)*as.matrix(dist(dd, method = "euclidean", upper = TRUE)^2))
Laplacian(A, laplacian='shift', grf.type = 'knn')</pre>
```

Lapprox

Compute a rank k approximation of a graph Laplacian

# Description

This function calculates the rank-k approximation of a graph Laplacian (or any symmetric matrix). This function performs eigen decomposition on the given matrix L and reconstructs it using only the LAST k eigenvectors and eigenvalues.

#### Usage

Lapprox(LapList, k, laplacian = c("shift", "Ng", "sym", "rw"), plots = TRUE)

#### Arguments

LapList	A list of Laplacian matrices
k	A vector indicating how many eigenvectors to take from each Laplacian, i.e., the number of clusters in each view
laplacian	One of "shift", "Ng", "rw" or "sym". Should be the same type used to calculate your Laplacians
plots	Whether or not to plot the eigenvalues from the rank approximated Laplacians

# Value

An  $n \times n$  matrix

#### Examples

```
## Generating data with 2 and 3 distinct clusters
## Note that 'clustStruct' returns a list
n=120; k <- c(2,3)
set.seed(23)
dd <- clustStruct(n=n, p=30, k=k, noiseDat='random')
## Laplacians
L_list <- lapply(dd, kernelLaplacian, kernel="Spectrum",
laplacian='shift', plots=FALSE, verbose=FALSE)
trueGroups(n,k)
La <- Lapprox(L_list, k=k, plots=FALSE)
kmeans(La$vectors[,1:4], centers=4)</pre>
```

trueGroups

Get the groups created by the clustStruct function

# Description

Get the unique groups generated by the clustStruct function for a given k. The number of rows of the resulting matrix gives the number of unique groups.

# Usage

trueGroups(n, k)

# Arguments

n	An integer, the sample size for all generated data sets
k	An integer or vector, the number of distinct clusters in each generated data set.
	n/k must be an integer for all values of k

#### Value

A matrix with the unique groups/clusters from the multi-view data generated from clustStruct. The final column Grps enumerates these groups.

# Examples

trueGroups(n=120, k=c(2,3,4))

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