

Package ‘IPCAPS’

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Type Package

Title Iterative Pruning to Capture Population Structure

Version 1.1.8

Description

An unsupervised clustering algorithm based on iterative pruning is for capturing population structure. This version supports ordinal data which can be applied directly to SNP data to identify fine-level population structure and it is built on the iterative pruning Principal Component Analysis ('ipPCA') algorithm as explained in Intarapanich et al. (2009) <[doi:10.1186/1471-2105-10-382](https://doi.org/10.1186/1471-2105-10-382)>. The 'IPCAPS' involves an iterative process using multiple splits based on multivariate Gaussian mixture modeling of principal components and 'Expectation-Maximization' clustering as explained in Lebret et al. (2015) <[doi:10.18637/jss.v067.i06](https://doi.org/10.18637/jss.v067.i06)>. In each iteration, rough clusters and outliers are also identified using the function rubikclust() from the R package 'KRIS'.

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Suggests testthat

BugReports <https://gitlab.com/kris.ccp/ipcaps/-/issues>

URL <https://gitlab.com/kris.ccp/ipcaps>

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'save.html.R' 'save.eigenplots.html.R'
'save.plots.label.html.R' 'save.plots.cluster.html.R'
'save.plots.R' 'postprocess.R' 'preprocess.R' 'ipcaps.R'
'top.discriminator.R'

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cal.eigen.fit	<i>(Internal function) Calculae a vector of EigenFit values, internally used for parallelization</i>
---------------	------------------------------------------------------------------------------------------------------

Description

(Internal function) Calculae a vector of EigenFit values, internally used for parallelization

Usage

```
cal.eigen.fit(eigen.value)
```

Arguments

eigen.value	A vector of Eigenvalues return from <code>svd</code> (\$d), <code>rARPACK::svds</code> (\$d), <code>eigen</code> (\$values) or <code>rARPACK::eigs</code> (\$values).
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Value

A vector of all possible values for EigenFit.

check.stopping	<i>(Internal function) Check whether the IPCAPS process meets the stopping criterion.</i>
----------------	-------------------------------------------------------------------------------------------

Description

(Internal function) Check whether the IPCAPS process meets the stopping criterion.

Usage

```
check.stopping(eigen.value, threshold)
```

Arguments

eigen.value	A vector of Eigenvalues return from <code>svd</code> (\$d), <code>rARPACK::svds</code> (\$d), <code>eigen</code> (\$values) or <code>rARPACK::eigs</code> (\$values).
threshold	A threshold or a cutoff to stop the IPCAPS process. Also see <code>ipcaps</code> (the parameter <code>threshold</code>).

Value

A list containing `status`, `eigen.value`, `eigen.fit`, `threshold`, and `no.significant.PC` as explained below:

- `$status` is either 0 representing that the criterion is not met, or 1 representing that the criterion is met.
- `$eigen.value` is a vector of Eigenvalues as the input parameter.
- `$eigen.fit` is a vector of EigenFit values.
- `$threshold` is a threshold as the input parameter.
- `$no.significant.PC` is an estimated number of significant principal components (PC).

`clustering`

(Internal function) Perform the clustering process of IPCAPS

Description

(Internal function) Perform the clustering process of IPCAPS

Usage

```
clustering(
  dataframe,
  node = 1,
  result.dir,
  threshold,
  min.fst,
  method = "mix",
  min.in.group = 20,
  datatype = "snp",
  nonlinear = FALSE
)
```

Arguments

<code>dataframe</code>	A data frame containing <code>raw.data</code> (matrix or data frame), <code>label</code> (vector), and <code>index</code> (vector). <code>raw.data</code> represents a matrix of subset of input data. <code>label</code> represents a vector of labels for all rows of <code>raw.data</code> . <code>index</code> represents a vector of indexes that are selected from the original input data.
<code>node</code>	An integer representing the current node number which is being analyzed.
<code>result.dir</code>	An output directory
<code>threshold</code>	A threshold or a cutoff to stop the IPCAPS process. Also see ipcaps (the parameter <code>threshold</code>).
<code>min.fst</code>	A number represents a cutoff for minimum Fst value.
<code>method</code>	A clustering method selected from the ipcaps function. See ipcaps for available methods.

<code>min.in.group</code>	A integer represents a minimum number of group members.
<code>datatype</code>	To specify whether the input data are 'snp' or other type. Default = 'snp'.
<code>nonlinear</code>	(Unimplemented) To specify whether linear or non-linear method is used for IPCAPS analysis. If TRUE, non-linear method is used, otherwise linear method is used. Default = FALSE.

Value

A list containing `status`, `node`, and `new.index` as explained below:

- `$status` is either 0 representing that the criterion is not met, or 1 representing that the criterion is met.
- `$node` is an integer representing the current node number which is being analyzed.
- `$new.index` is a list of vectors containing a subset of indexes split from `dataframe$index` according to a clustering result.

`clustering.mode`

(Internal function) Select a clustering method to be used for the IP-CAPS process.

Description

(Internal function) Select a clustering method to be used for the IPCAPS process.

Usage

```
clustering.mode(node, work.dir, method)
```

Arguments

<code>node</code>	An integer representing the current node number which is being analyzed.
<code>work.dir</code>	A working directory.
<code>method</code>	A clustering method selected from the ipcaps function. See ipcaps for available methods.

Value

A vector of cluster assignment, for which cluster each individual belongs.

See Also

[ipcaps](#)

diff.eigen.fit *(Internal function) Calculate a vector of different values from a vector of EigenFit values, internally used for parallelization*

Description

(Internal function) Calculate a vector of different values from a vector of EigenFit values, internally used for parallelization

Usage

```
## S3 method for class 'eigen.fit'
diff(eigen.value)
```

Arguments

eigen.value A vector of Eigenvalues return from svd (\$d), rARPACK::svds (\$d), eigen (\$values) or rARPACK::eigs (\$values).

Value

A vector of different values from a vector of EigenFit values

diff.xy *(Internal function) Check the different value of X and Y, internally used for parallelization*

Description

(Internal function) Check the different value of X and Y, internally used for parallelization

Usage

```
## S3 method for class 'xy'
diff(x, y)
```

Arguments

x	The first number
y	The second number

Value

The different number of x and y.

`do.glm`

(Internal function) Perform regression models, internally used for parallelization

Description

(Internal function) Perform regression models, internally used for parallelization

Usage

```
do.glm(X, PC, method = "linear")
```

Arguments

X	A vector of data
PC	A matrix of principal components
method	Specify a method to be used for regression model, which can be "linear", "poisson", and "negative.binomial". Default = "linear"

Value

A vector of residuals computed from regression model

`export.groups`

Export the IPCAPS result to a text file

Description

Export clustering result of [ipcaps](#) to text file called 'groups.txt'.

Usage

```
export.groups(result.dir)
```

Arguments

result.dir	A result directory as the \$output object returned from the ipcaps function.
------------	----------------------------------------------------------------------------------------------

Details

After running, this function exports the file called 'groups.txt' to the same result directory. If 'groups.txt' already exists in the result directory, the exported file is changed to 'groups1.txt', 'groups2.txt', 'groups3.txt', ..., accordingly.

Value

A data frame of exported data containing 4 columns; `group`, `node`, `label`, `row.number`, as described below for more details:

- `group` represents group membership of IPCAPS result.
- `node` represents node numbers of IPCAPS result.
- `label` represents labels of rows in orginal input data.
- `row.number` represents row numbers of orginal input data.

Examples

```
# Importantly, bed file, bim file, and fam file are required
# Use the example files embedded in the package

BED.file <- system.file("extdata","ipcaps_example.bed",package="IPCAPS")
LABEL.file <- system.file("extdata","ipcaps_example_individuals.txt.gz",package="IPCAPS")

my.cluster <- ipcaps(bed=BED.file,label.file=LABEL.file,lab.col=2,out=tempdir())

#Here, to export the IPCAPS result to a text file
exported.data <- export.groups(my.cluster$output.dir)
print(dim(exported.data))
head(exported.data)
```

`get.node.info`

Get the information for specified node

Description

Obtain the information for specified node from the output list of [ipcaps](#).

Usage

```
get.node.info(cluster.obj, node)
```

Arguments

- | | |
|--------------------------|-------------------------------------------------------------------------------------------------|
| <code>cluster.obj</code> | A list returned from the ipcaps function. |
| <code>node</code> | An integer representing a node number to enquire information as shown in the HTML output files. |

Value

The return value is NULL if node's information does not exist or a list containing PCs, eigen.fit, index, and label as explained below:

- \$PCs is a matrix of principal components of this node.
- \$eigen.fit is a number representing the EigenFit value of this node.
- \$index is a vector of row number (individuals) of raw.data (input data).
- \$label is the vector of labels of all individuals that belongs to this node.

Examples

```
# Importantly, bed file, bim file, and fam file are required
# Use the example files embedded in the package

BED.file <- system.file("extdata","ipcaps_example.bed",package="IPCAPS")
LABEL.file <- system.file("extdata","ipcaps_example_individuals.txt.gz",package="IPCAPS")

my.cluster <- ipcaps(bed=BED.file,label.file=LABEL.file,lab.col=2,out=tempdir())

#Here, to obtain the information of specified node, for example, node 3
node.info <- get.node.info(my.cluster,3)
ls(node.info)
```

ipcaps

Perform unsupervised clustering to capture population structure based on iterative pruning

Description

This version supports ordinal data which can be applied directly to SNP data to identify fine-scale population structure. It was built on the iterative pruning Principal Component Analysis (ipPCA) algorithm (Intarapanich et al., 2009; Limpiti et al., 2011). The IPCAPS involves an iterative process using multiple splits based on multivariate Gaussian mixture modeling of principal components and Clustering EM estimation (Lebret et al., 2015). In each iteration, rough clusters and outliers are also identified using our own method called rubikclust (R package **KRIS**).

Usage

```
ipcaps(
  bed = NA,
  rdata = NA,
  files = NA,
  label.file = NA,
  lab.col = 1,
  out,
  plot.as.pdf = FALSE,
```

```

method = "mix",
missing = NA,
covariate = NA,
cov.col.first = NA,
cov.col.last = NA,
threshold = 0.18,
min.fst = 8e-04,
min.in.group = 20,
no.plot = FALSE
)

```

Arguments

<code>bed</code>	A PLINK binary format consists of 3 files; bed, bim, and fam. To generate these files from PLINK, use option <code>-make-bed</code> . See more details at: http://zzz.bwh.harvard.edu/plink/data.shtml .
<code>rdata</code>	In case of re-analysis, it is convenient to run IPCAPS using the file <code>rawdata.RData</code> generated by IPCAPS. This file contains a matrix of SNPs (<code>raw.data</code>) and a vector of labels (<code>label</code>).
<code>files</code>	IPCAPS supports SNPs encoded as 0, 1 and 2 (dosage encoding). Rows represent SNPs and columns represent subjects. Each column needs to be separated by a space or a tab. A big text file should be divided into smaller files to load faster. For instance, to input 3 files, use as: <code>files=c('input1.txt', 'input2.txt', 'input3.txt')</code> .
<code>label.file</code>	An additional useful information (called "labels" in IPCAPS) related subject, for example, geographic location or disease phenotype. These labels (one at a time) are used in displaying the clustering outcome of IPCAPS. A label file must contain at least one column. However, it may contain more than one column in which case each column need to be separated by a space or a tab.
<code>lab.col</code>	The label in the label file to be used in the tree-like display of IPCAPS clustering results.
<code>out</code>	To set an absolute path for IPCAPS output. If the specified output directory already exists, result files are saved in sub-directories <code>cluster_out</code> , <code>cluster_out1</code> , <code>cluster_out2</code> , etc.
<code>plot.as.pdf</code>	To export plots as PDF. When omitted, plots are saved as PNG.
<code>method</code>	The internal clustering method. It can be set to "mix" (rubikclust & mixmod), "mixmod" (Lebret et al., 2015), "clara" (R: Clustering Large Applications), "pam" (R: Partitioning Around Medoids (PAM) Object), "meanshift" (Wang, 2016), "apcluster" (Bodenhofer et al., 2016), and "hclust" (R: Hierarchical Clustering). Default = "mix".
<code>missing</code>	Symbol used for missing genotypes. Default = NA.
<code>covariate</code>	A file of covariates; one covariate per column. SNPs can be adjusted for these covariates via regression modeling and residual computation.
<code>cov.col.first</code>	Refer to a covariate file, the first covariate to be considered as confounding variable.

<code>cov.col.last</code>	Refer to a covariate file, the last covariate to be considered as confounding variable. All the variables in between the <code>cov.col.first</code> and <code>cov.col.last</code> will be considered in the adjustment process.
<code>threshold</code>	Cutoff value for EigenFit. Possible values range from 0.03 to 0.18. The smaller, the potentially finer the substructure can be. Default = 0.18.
<code>min.fst</code>	Minimum Fst between a pair of subgroups. Default = 0.0008.
<code>min.in.group</code>	Minimum number of individuals to constitute a cluster or subgroup. Default = 20.
<code>no.plot</code>	No plot is generated if this option is TRUE. This option is useful when the system does not support X Windows in the unix based system. Default = FALSE.

Details

The computational time depends on the number of individuals. Consequentially, if large data sets are analyzed, it may be necessary first to split data into smaller files, and then load into computer memory (with parameter 'files'). Internally, the split files are merged to construct a comprehensive matrix.

Value

Returns the list object containing 2 internal objects; `output.dir` as class character and `cluster` as class `data.frame`. The object `output.dir` stores a result directory. The object `cluster` contains 4 columns, `group`, `node`, `label`, and `row.number`. The column `group` contains the assigned groups from IPCAPS. The column `node` contains node numbers in a tree as illustrated in the HTML result files. The column `label` contains the given labels that is set to the parameter `label`. The column `row.number` contains indices to an input data, which is matched to row numbers of input matrix. All clustering result files are saved in one directory (as specified by `out`) containing assigned groups, hierarchical trees of group membership, PC plots, and binary data for further analysis.

- `$snp` is a SNP matrix from BED file.
- `$snp.info` is a `data.frame` of SNP information from BIM file.
- `$ind.info` is a `data.frame` of individual information from FAM file.

If function return NULL, it means the input files are not in proper format.

References

- Bodenhofer, U., Palme, J., Melkonian, C., and Kothmeier, A. (2016). `apcluster` : Affinity Propagation Clustering. Available at: <https://CRAN.R-project.org/package=apcluster> (Accessed March 7, 2017).
- Intarapanich, A., Shaw, P. J., Assawamakin, A., Wangkumhang, P., Ngamphiw, C. , Chaichoompu, K., et al. (2009). Iterative pruning PCA improves resolution of highly structured populations. *BMC Bioinformatics* 10, 382. doi:10.1186/1471-2105-10-382.
- Lebret, R., Iovleff, S., Langrognet, F., Biernacki, C., Celeux, G., and Govaert, G. (2015). `Rmixmod`: The R Package of the Model-Based Unsupervised, Supervised, and Semi-Supervised Classification-`MixmodLibrary`. *J. Stat. Softw.* 67. doi:10.18637/jss.v067.i06.

Limpiti, T., Intarapanich, A., Assawamakin, A., Shaw, P. J., Wangkumhang, P., Piriyapongsa, J., et al. (2011). Study of large and highly stratified population datasets by combining iterative pruning principal component analysis and structure. BMC Bioinformatics 12, 255. doi:10.1186/1471-2105-12- 255.

Maechler, M., Rousseeuw, P., Struyf, A., Hubert, M., and Hornik, K. (2017). cluster: Cluster Analysis Basics and Extensions.

R: Clustering Large Applications Available at: <https://stat.ethz.ch/R-manual/R-devel/library/cluster/html/clara.html> (Accessed March 7, 2017).

R Core Team (2017). R: A Language and Environment for Statistical Computing. Vienna, Austria: R Foundation for Statistical Computing Available at: <https://www.R-project.org/>.

R: Hierarchical Clustering Available at: <https://stat.ethz.ch/R-manual/R-devel/library/stats/html/hclust.html> (Accessed March 7, 2017).

R: Partitioning Around Medoids (PAM) Object Available at: <https://stat.ethz.ch/R-manual/R-devel/library/cluster/html/pam.object.html> (Accessed March 7, 2017).

Wang, M. C. and D. (2016). MeanShift: Clustering via the Mean Shift Algorithm. Available at: <https://CRAN.R-project.org/package=MeanShift> (Accessed March 7, 2017).

Examples

```
#Use the BED format as input
#Importantly, bed file, bim file, and fam file are required
#Use the example files embedded in the package

BED.file <- system.file("extdata", "ipcaps_example.bed", package = "IPCAPS")
LABEL.file <- system.file("extdata", "ipcaps_example_individuals.txt.gz",
                         package = "IPCAPS")
my.cluster1 <- ipcaps(bed = BED.file, label.file = LABEL.file, lab.col = 2,
out = tempdir())

table(my.cluster1$cluster$label, my.cluster1$cluster$group)

# Alternatively, use a text file as input
# Use the example files embedded in the package

#text.file <- system.file("extdata", "ipcaps_example_rowVar_colInd.txt.gz",
#                         package="IPCAPS")
#LABEL.file <- system.file("extdata", "ipcaps_example_individuals.txt.gz",
#                         package="IPCAPS")

#my.cluster2 <- ipcaps(files = c(text.file), label.file = LABEL.file, lab.col = 2,
#                       #                     out=tempdir())
#table(my.cluster2$cluster$label, my.cluster2$cluster$group)

# The other alternative way, use an R Data file as input
# Use the example file embedded in the package

#rdata.file <- system.file("extdata", "ipcaps_example.rda", package = "IPCAPS")
```

```
#my.cluster3 <- ipcaps(rdata = rdata.file, out = tempdir())
#table(my.cluster3$cluster$label, my.cluster3$cluster$group)
```

label	<i>Synthetic dataset containing population labels for the dataset raw.data</i>
--------------	--------------------------------------------------------------------------------

Description

A dataset contains a character vector of 1,004 elements containing labels or populations of 1,004 individuals which they belong. Five populations and outliers were labeled as "pop1", "pop2", "pop3", "pop4", "pop5", and "outlier".

Usage

```
data(ipcaps_example)
```

Format

A vector with 1,004 elements.

See Also

[raw.data](#) and [PC](#)

output.template	<i>(Internal object) The HTML output template for IPCAPS</i>
------------------------	--------------------------------------------------------------

Description

`output.template` contains `$lno_title`, `$lno_data`, `$lno_leafnode`, `$lno_body`, and `$template` as explained below:

- lno_title** A index number of `$template` that a title of HTML file is replaced.
- lno_data** A index number of `$template` that a data section in the HTML file is replaced.
- lno_leafnode** A index number of `$template` that a leaf-node section in the HTML file is replaced.
- lno_body** A index number of `$template` that a body section in the HTML file is replaced.
- template** A vector of characters for HTML file.

Usage

```
output.template
```

Format

A list with with 5 objects

`pasre.categorical.data`

(Internal function) Manipulate categorical input files

Description

(Internal function) Manipulate categorical input files

Usage

```
pasre.categorical.data(files)
```

Arguments

<code>files</code>	IPCAPS supports categorical data, which rows represent features and columns represent subjects or individuals. Each column needs to be separated by a space or a tab. A big text file should be divided into smaller files to load faster. For instance, to input 3 files, use as: <code>files=c('input1.txt', 'input2.txt', 'input3.txt')</code> .
--------------------	-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------

Value

A data frame of input data

`PC`

Synthetic dataset containing the top 10 principal components (PC) from the dataset `raw.data`

Description

A dataset contains a numeric matrix of 1,004 rows and 10 columns of top 10 PCs calculated from the dataset `raw.data`. The PCs were calculated using linear principal component analysis (PCA), see more details at `KRIS::cal.pc.linear`

Usage

```
data(ipcaps_example)
```

Format

A matrix with 10 columns and 1,004 rows

See Also

[raw.data](#) and [label](#)

postprocess*(Internal function) Perform the post-processing step of IPCAPS*

Description

(Internal function) Perform the post-processing step of IPCAPS

Usage

```
postprocess(result.dir, reanalysis = FALSE)
```

Arguments

- | | |
|-------------------------|--------------------------------------------------------------------------------------------------------------------------------|
| <code>result.dir</code> | A result directory as the \$output object returned from the ipcaps function. |
| <code>reanalysis</code> | (Unimplemented) To specify whether it is re-analysis or not. If TRUE, it is re-analysis, otherwise it is not. Default = FALSE. |

Value

A data frame of clustering result containing 4 columns; group, node, label, row.number, as described below for more details:

- `group` represents group membership of IPCAPS result.
- `node` represents node numbers of IPCAPS result.
- `label` represents labels of rows in orginal input data.
- `row.number` represents row numbers of orginal input data.

preprocess*(Internal function) Perform the pre-processing step of IPCAPS*

Description

(Internal function) Perform the pre-processing step of IPCAPS

Usage

```
preprocess(  
  files,  
  label.file,  
  lab.col,  
  rdata.infile,  
  bed.infile,  
  cate.list,  
  result.dir,  
  threshold,
```

```

min.fst,
max.thread = NA,
reanalysis = FALSE,
method = "mix",
min.in.group = 20,
datatype = "snp",
nonlinear = FALSE,
missing.char = NA,
regression.file = NA,
regression.col.first = NA,
regression.col.last = NA,
reg.method = "linear",
plot.as.pdf = NA,
no.plot = NA
)

```

Arguments

<code>files</code>	IPCAPS supports SNPs encoded as 0, 1 and 2 (dosage encoding). Rows represent SNPs and columns represent subjects. Each column needs to be separated by a space or a tab. A big text file should be divided into smaller files to load faster. For instance, to input 3 files, use as: <code>files=c('input1.txt', 'input2.txt', 'input3.txt')</code> .
<code>label.file</code>	An additional useful information (called "labels" in IPCAPS) related subject, for example, geographic location or disease phenotype. These labels (one at a time) are used in displaying the clustering outcome of IPCAPS. A label file must contain at least one column. However, it may contain more than one column in which case each column need to be separated by a space or a tab.
<code>lab.col</code>	The label in the label file to be used in the tree-like display of IPCAPS clustering results.
<code>rdata.infile</code>	In case of re-analysis, it is convenient to run IPCAPS using the file <code>rawdata.RData</code> generated by IPCAPS. This file contains a matrix of SNPs (<code>raw.data</code>) and a vector of labels (<code>label</code>).
<code>bed.infile</code>	A PLINK binary format consists of 3 files; bed, bim, and fam. To generate these files from PLINK, use option <code>-make-bed</code> . See more details at: http://zzz.bwh.harvard.edu/plink/data.shtml .
<code>cate.list</code>	(Unimplemented) A list of categorical input file (text). For instance, to input 3 files, use as: <code>files=c('input1.txt', 'input2.txt', 'input3.txt')</code> .
<code>result.dir</code>	To set an absolute path for IPCAPS output. If the specified output directory already exists, result files are saved in sub-directories <code>cluster_out</code> , <code>cluster_out1</code> , <code>cluster_out2</code> , etc.
<code>threshold</code>	Cutoff value for EigenFit.
<code>min.fst</code>	Minimum Fst between a pair of subgroups.
<code>max.thread</code>	(Require the parallelization patch) Maximum number of concurrent threads.
<code>reanalysis</code>	(Unimplemented) To specify whether it is re-analysis or not. If TRUE, it is re-analysis, otherwise it is not. Default = FALSE.

method	The internal clustering method. It can be set to "mix" (rubikclust & mixmod), "mixmod" (Lebret et al., 2015), "clara" (R: Clustering Large Applications), "pam" (R: Partitioning Around Medoids (PAM) Object), "meanshift" (Wang, 2016), "apcluster" (Bodenhofer et al., 2016), and "hclust" (R: Hierarchical Clustering). Default = "mix".
min.in.group	Minimum number of individuals to constitute a cluster or subgroup.
datatype	To specify whether the input data are 'snp' or other type. Default = 'snp'.
nonlinear	(Unimplemented) To specify whether linear or non-linear method is used for IPCAPS analysis. If TRUE, non-linear method is used, otherwise linear method is used. Default = FALSE.
missing.char	Symbol used for missing genotypes. Default = NA.
regression.file	A file of covariates; one covariate per column. SNPs can be adjusted for these covariates via regression modeling and residual computation.
regression.col.first	Refer to a covariate file, the first covariate to be considered as confounding variable.
regression.col.last	Refer to a covariate file, the last covariate to be considered as confounding variable. All the variables in between the cov.col.first and cov.col.last will be considered in the adjustment process.
reg.method	(Fixed) Specify a method for regression analysis. Default = 'linear'.
plot.as.pdf	To export plots as PDF. When omitted, plots are saved as PNG.
no.plot	No plot is generated if this option is TRUE. This option is useful when the system does not support X Windows in the unix based system. Default = FALSE.

Value

A data frame of input data.

process.each.node

(Internal function) Perform the iterative process for each node

Description

(Internal function) Perform the iterative process for each node

Usage

```
process.each.node(node, work.dir)
```

Arguments

node	An integer representing the current node number which is being analyzed.
work.dir	A working directory.

Value

NULL

raw.data

*Synthetic dataset containing single nucleotide polymorphisms (SNP)***Description**

The raw.data is the simulated dataset which consists of 3,000 independent SNPs and 1,004 individuals belonging to one of 5 populations (200 individuals each) and 4 outlying individuals. The matrix raw.data contains the number 0, 1, and 2 representing SNP in additive coding. The pairwise genetic distance between populations are listed below (see Balding, 1995):

	pop1	pop2	pop3	pop4	pop5
pop1		0.0040	0.0059	0.0085	0.0101
pop2	0.0040		0.0055	0.0082	0.0099
pop3	0.0059	0.0055		0.0104	0.0119
pop4	0.0085	0.0082	0.0104		0.0139
pop5	0.0101	0.0099	0.0119	0.0139	

Usage

```
data(ipcaps_example)
```

Format

A matrix with 3,000 columns and 1,004 rows

References

Balding, D.J., and Nichols, R.A. (1995). A method for quantifying differentiation between populations at multi-allelic loci and its implications for investigating identity and paternity. *Genetica* 96, 3-12.

See Also

[label](#) and [PC](#)

replace.missing

*(Internal function) Replace missing values by specified values, internally used for parallelization***Description**

(Internal function) Replace missing values by specified values, internally used for parallelization

Usage

```
replace.missing(X, missing = NA, rep)
```

Arguments

X	A vector of data
missing	The old characters representing a missing value, for example, 'NA' or '-'
rep	The new characters to replace the old characters, for example, 'NULL' or '1'

Value

A vector of data with replaced character for miss values

save.eigenplots.html *Generate HTML file for EigenFit plots*

Description

Generate HTML file called 'tree_scree.html' from the result of [ipcaps](#). This function is a part of workflow in [save.plots](#). The clustering result is shown as a tree rendering by the online Google Organizational Chart library. Note that the Internet is required to view the HTML file.

Usage

```
save.eigenplots.html(output.dir)
```

Arguments

output.dir	A result directory as the \$output object returned from the ipcaps function.
------------	----------------------------------------------------------------------------------------------

Details

After running, this function generates the file called 'tree_scree.html' in the same result directory. All plots are generated and saved as image files in the sub-directory 'images'.

Value

NULL

See Also

[save.html](#), [save.plots](#), [save.plots.cluster.html](#), and [save.plots.label.html](#)

Examples

```
# Importantly, bed file, bim file, and fam file are required
# Use the example files embedded in the package

BED.file <- system.file("extdata","ipcaps_example.bed",package="IPCAPS")
LABEL.file <- system.file("extdata","ipcaps_example_individuals.txt.gz",package="IPCAPS")

my.cluster <- ipcaps(bed=BED.file,label.file=LABEL.file,lab.col=2,out=tempdir())

#Here, to generate HTML file
save.eigenplots.html(my.cluster$output.dir)
```

save.html

Generate HTML file for clustering result in text mode

Description

Generate HTML file called 'tree_text.html' from the result of [ipcaps](#). The clustering result is shown as a tree rendering by the online Google Organizational Chart library. Note that the Internet is required to view the HTML file.

Usage

```
save.html(output.dir)
```

Arguments

output.dir A result directory as the \$output object returned from the [ipcaps](#) function.

Details

After running, this function generates the file called 'tree_text.html' in the same result directory.

Value

NULL

See Also

[save.plots](#), [save.plots.cluster.html](#), [save.eigenplots.html](#), and [save.plots.label.html](#)

Examples

```
# Importantly, bed file, bim file, and fam file are required
# Use the example files embedded in the package

BED.file <- system.file("extdata","ipcaps_example.bed",package="IPCAPS")
LABEL.file <- system.file("extdata","ipcaps_example_individuals.txt.gz",package="IPCAPS")

my.cluster <- ipcaps(bed=BED.file,label.file=LABEL.file,lab.col=2,out=tempdir())

#Here, to generate HTML file
save.html(my.cluster$output.dir)
```

save.plots

Workflow to generate HTML files for all kinds of plots

Description

Generate HTML files and all image files (plots) from the result of [ipcaps](#). The clustering result is shown as a tree rendering by the online Google Organizational Chart library. Note that the Internet is required to view the HTML files.

Usage

```
save.plots(output.dir)
```

Arguments

`output.dir` A result directory as the `$output` object returned from the [ipcaps](#) function.

Details

After running, this function generates all plots and saves as image files in the sub-directory 'images'. It calls [save.plots.cluster.html](#), [save.eigenplots.html](#), and [save.plots.label.html](#) to generate all HTML files.

Value

NULL

See Also

[save.html](#), [save.plots.cluster.html](#), [save.eigenplots.html](#), and [save.plots.label.html](#)

Examples

```
# Importantly, bed file, bim file, and fam file are required
# Use the example files embedded in the package

BED.file <- system.file("extdata","ipcaps_example.bed",package="IPCAPS")
LABEL.file <- system.file("extdata","ipcaps_example_individuals.txt.gz",package="IPCAPS")

my.cluster <- ipcaps(bed=BED.file,label.file=LABEL.file,lab.col=2,out=tempdir())

#Here, to generate all plots and HTML files
save.plots.label.html(my.cluster$output.dir)
```

save.plots.cluster.html

Generate HTML file for scatter plots which all data points are highlighted by IPCAPS clusters

Description

Generate HTML file called 'tree_scatter_cluster.html' from the result of [ipcaps](#). This function is a part of workflow in [save.plots](#). The clustering result is shown as a tree rendering by the online Google Organizational Chart library. Note that the Internet is required to view the HTML file.

Usage

```
save.plots.cluster.html(output.dir)
```

Arguments

output.dir	A result directory as the \$output object returned from the ipcaps function.
------------	----------------------------------------------------------------------------------------------

Details

After running, this function generates the file called 'tree_scatter_cluster.html' in the same result directory. All plots are generated and saved as image files in the sub-directory 'images'.

Value

NULL

See Also

[save.html](#), [save.plots](#), [save.eigenplots.html](#), and [save.plots.label.html](#)

Examples

```
# Importantly, bed file, bim file, and fam file are required
# Use the example files embedded in the package

BED.file <- system.file("extdata","ipcaps_example.bed",package="IPCAPS")
LABEL.file <- system.file("extdata","ipcaps_example_individuals.txt.gz",package="IPCAPS")

my.cluster <- ipcaps(bed=BED.file,label.file=LABEL.file,lab.col=2,out=tempdir())

#Here, to generate HTML file
save.plots.cluster.html(my.cluster$output.dir)
```

save.plots.label.html *Generate HTML file for scatter plots which data points are highlighted by given labels*

Description

Generate HTML file called 'tree_scatter_label.html' from the result of [ipcaps](#). This function is a part of workflow in [save.plots](#). The clustering result is shown as a tree rendering by the online Google Organizational Chart library. Note that the Internet is required to view the HTML file.

Usage

```
save.plots.label.html(output.dir)
```

Arguments

output.dir A result directory as the \$output object returned from the [ipcaps](#) function.

Details

After running, this function generates the file called 'tree_scatter_label.html' in the same result directory. All plots are generated and saved as image files in the sub-directory 'images'.

Value

NULL

See Also

[save.html](#), [save.plots](#), [save.plots.cluster.html](#), and [save.eigenplots.html](#)

Examples

```
# Importantly, bed file, bim file, and fam file are required
# Use the example files embedded in the package

BED.file <- system.file("extdata","ipcaps_example.bed",package="IPCAPS")
LABEL.file <- system.file("extdata","ipcaps_example_individuals.txt.gz",package="IPCAPS")

my.cluster <- ipcaps(bed=BED.file,label.file=LABEL.file,lab.col=2,out=tempdir())

#Here, to generate HTML file
save.plots.label.html(my.cluster$output.dir)
```

top.discriminator *Detecting top discriminators between two groups*

Description

Detects top discriminators that contribute to group separation based on the fixation index (Fst).

Usage

```
top.discriminator(
  cluster.obj = NULL,
  group1,
  group2,
  bim.file,
  use.node.number = FALSE,
  num.top = 100,
  percentile = 0.9,
  use.percentile = FALSE,
  use.path = FALSE,
  result.path = NULL
)
```

Arguments

<code>cluster.obj</code>	The object which is returned from <code>ipcaps</code> . This parameter is used when <code>use.path</code> is FALSE.
<code>group1</code>	To specify the first group number to be compared. (also see <code>use.node.number</code>)
<code>group2</code>	To specify the second group number to be compared. (also see <code>use.node.number</code>)
<code>bim.file</code>	Option: In case that SNP information is not provided to <code>ipcaps</code> , an absolute path of SNP information file is required. It needs to be in PLINK format (bim). See more details at: http://zzz.bwh.harvard.edu/plink/data.shtml .
<code>use.node.number</code>	To specify whether a group number or a node number is be used. If TRUE, a node number is used instead. Default = FALSE.

num.top	A number of top Fst SNPs to be returned. This parameter is used when percentile is FALSE. Default = 100.
percentile	A percentile for top Fst SNPs to be returned. This parameter is used when percentile is TRUE. Default = 0.999.
use.percentile	A logical value to indicate whether percentile is used instead of num.top. This parameter is used when percentile is TRUE. Default = FALSE.
use.path	A logical value to indicate whether result.path is used instead of cluster.obj. Importantly, result.path needs to be set. This parameter only work with the IPCAPS's result from version 1.1.7 onward. Default = FALSE.
result.path	A path to an result directory of IPCAPS. This parameter is used when use.path is TRUE. This parameter only work with the IPCAPS's result from version 1.1.7 onward.

Value

The returned value is a data.frame of SNP information sorting by Fst in descending order, which contains 7 columns, chr, SNP, centimorgans, position, allele1, allele2, and Fst. The column 1-6 are SNP information from the bim file. The column Fst contains estimated Fst between group1 and group2.

Examples

```
# Importantly, bed file, bim file, and fam file are required
# Use the example files embedded in the package
BED.file <- system.file("extdata","ipcaps_example.bed",package="IPCAPS")
LABEL.file <- system.file("extdata","ipcaps_example_individuals.txt.gz",package="IPCAPS")
my.cluster <- ipcaps(bed=BED.file,label.file=LABEL.file,lab.col=2,out=tempdir())
table(my.cluster$cluster$label,my.cluster$cluster$group)
# 1 2 3 4 5 6
# outlier4 5 4 1 0 0 0
# pop1 0 0 0 0 250 0
# pop2 0 0 0 0 0 250
# pop3 0 0 0 250 0 0

#Identify top discriminators between groups, for example, group 4 and group 5
top.snp <- top.discriminator(my.cluster,4,5)
#or, specify the bim file
#top.snp <- top.discriminator(my.cluster,4,5,bim.file="ipcaps_example.bim")
head(top.snp)
# chr SNP centimorgans position allele1 allele2 Fst
#V5452 1 marker5452 0 54520000 A T 0.11337260
#V2348 1 marker2348 0 23480000 A T 0.11194490
#V8244 1 marker8244 0 82440000 A T 0.09556580
#V5972 1 marker5972 0 59720000 A T 0.08747794
#V3561 1 marker3561 0 35610000 A T 0.08725860
#V8419 1 marker8419 0 84190000 A T 0.08293494

#Alternatively, specify the previous result directory of IPCAPS and identify
#top discriminators between groups, for example, group 4 and group 5
```

```
previous.res.path <- my.cluster$output.dir
top.snp <-top.discriminator(result.path = previous.res.path, use.path = TRUE,
group1 = 4, group2 = 5)
head(top.snp)

#Identify top discriminators between groups, for example, group 4 and group 5
top.snp <-top.discriminator(my.cluster,4,5)
#or, specify the bim file
#top.snp <-top.discriminator(my.cluster,4,5,bim.file="ipcaps_example.bim")
dim(top.snp)
head(top.snp)

#Here, it is possible to select the top Fst SNPs based on a percentile.
top.snp <-top.discriminator(my.cluster,4,5, percentile = 0.9,
use.percentile = TRUE)
dim(top.snp)
head(top.snp)

#Identify top discriminators between groups, for example, node 7 and node 8
top.snp2 <-top.discriminator(my.cluster,7,8,use.node.number=TRUE)
head(top.snp2)
# chr SNP centimorgans position allele1 allele2 Fst
#V5452 1 marker5452 0 54520000 A T 0.11337260
#V2348 1 marker2348 0 23480000 A T 0.11194490
```

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