# Package 'CONCUR'

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<b>Description</b> Implements a kernel-based association test for copy number variation (CNV) aggregate analysis in a certain genomic region (e.g., gene set, chromosome, or genome) that is robust to the within-locus and across-locus etiological heterogeneity, and bypass the need to define a ``locus'' unit for CNVs.  Brucker, A., et al. (2020) <doi:10.1101 666875="">.</doi:10.1101>
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cnvData

Pseudo Copy Number Variants Data

# Description

This data set includes simulated CNV data in PLINK CNV data format. The data are also available from the authors through the url provided below. These data were generated following the simulation study used to illustrate the method in the original manuscript also referenced below; it has been reduced to include only 600 individuals. These data are not meaningful and are intended for demonstration purposes only.

# Usage

data(cnvData)

#### **Format**

cnvData is a data.frame containing 522 observations with 5 columns:

ID character patient identifier.

CHR CNV chromosome.

**BP1** starting location in base pairs.

**BP2** ending location in base pairs.

**TYPE** copy number (0,1,3,or 4).

# References

Brucker, A., Lu, W., Marceau West, R., Yu, Q-Y., Hsiao, C. K., Hsiao, T-H., Lin, C-H., Magnusson, P. K. E., Holloway, S. T., Sullivan, P. F., Szatkiewicz, J. P., Lu, T-P., and Tzeng, J-Y. Association testing using Copy Number Profile Curves (CONCUR) enhances power in copy number variant analysis. <doi:10.1101/666875>.

https://www4.stat.ncsu.edu/~jytzeng/Software/CONCUR/

concur

Copy Number Profile Curve-Based Association Test

# **Description**

Implements a kernel-based association test for CNV aggregate analysis in a certain genomic region (e.g., gene set, chromosome, or genome) that is robust to the within-locus and across-locus etiologoical heterogeneity, and bypass the need to define a "locus" unit for CNVs.

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# Usage

```
concur(
   cnv,
   X,
   pheno,
   phenoY,
   phenoType,
   ...,
   nCore = 1L,
   outFileKernel = NULL,
   verbose = TRUE
)
```

# **Arguments**

cnv

A character or data.frame object. If character, the name of the data file containing the CNV data (with a header). If data.frame, the CNV data. The data must contain the following columns: "ID", "CHR", "BP1", "BP2", "TYPE", where "ID" is a unique patient id, "CHR" is the CNV chromosome, "BP1" is the start location in base pairs or kilo-base pairs, "BP2" is the end location in base pairs or kilo-base pairs, and "TYPE" is the CNV copy number.

Χ

A character or data frame object. If character, the name of the data file containing the covariate data (with a header). If data frame, the covariate data. The data must contain a column titled "ID" containing a unique patient id. This column must contain the patient identifiers of the CNV data specified in input cnv; however, it can contain patient identifiers not contained in cnv. Further, inputs X and pheno must contain the same patient identifiers. Categorical variables must be translated into design matrix format.

pheno

A character or data.frame object. If character, the name of the data file containing the phenotype data (with a header). If data.frame, the phenotype data. The data must contain a column titled "ID" containing a unique patient id. This column must contain the patient identifiers of the CNV data specified in input cnv; however, it can contain patient identifiers not contained in cnv. Further, inputs X and pheno must contain the same patient identifiers.

phenoY

A character object. The column name in input pheno containing the phenotype of interest.

phenoType

A character object. Must be one of of {"bin", "cont"} indicating if input phenoY (i.e., the phenotype of interest) is binary or continuous.

Ignored. Included to require named inputs.

nCore

An integer object. If nCore > 1, package parallel is used to calculate the kernel. Though the methods of package CompQuadForm dominate the time profile, setting nCore > 1L can improve computation times.

outFileKernel

A character object or NULL. If a character, the file in which the kernel is to be saved. If NULL, the kernel is returned by the function.

verbose

A logical object. If TRUE, progress information is printed to the screen.

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# **Details**

The CNV data must adhere to the following conditions:

- CNVs must be at least 1 unit long.
- CNVs cannot end at the exact location another begins

Violations of these conditions typically occur when data are rounded to a desired resolution. For example

```
ID CHR BP1 BP2 TYPE 1 13 10112087 10112414 3
```

becomes upon rounding to kilo

```
ID CHR BP1 BP2 TYPE
1 13 10112 10112 3
```

These cases should either be discarded or modified to be of length 1, e.g.,

```
ID CHR BP1 BP2 TYPE
1 13 10112 10113 3
```

As an example of condition 2

```
ID CHR BP1 BP2 TYPE
1 13 100768 101100 3
1 13 101100 101299 1
```

should be modified to one of

```
ID CHR BP1 BP2 TYPE
1 13 100768 101100 3
1 13 101101 101299 1
```

or

```
ID CHR BP1 BP2 TYPE
1 13 100768 101099 3
1 13 101100 101299 1
```

Additionally,

```
ID CHR BP1 BP2 TYPE
1 13 100768 101100 3
1 13 101100 101299 3
```

should be combined as

```
ID CHR BP1 BP2 TYPE
1 13 100768 101299 3
```

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#### Value

A list containing the kernel (or its file name) and the p-value.

#### References

Brucker, A., Lu, W., Marceau West, R., Yu, Q-Y., Hsiao, C. K., Hsiao, T-H., Lin, C-H., Magnusson, P. K. E., Holloway, S. T., Sullivan, P. F., Szatkiewicz, J. P., Lu, T-P., and Tzeng, J-Y. Association testing using Copy Number Profile Curves (CONCUR) enhances power in copy number variant analysis. <doi:10.1101/666875>.

#### **Examples**

```
data(cnvData)
# limit data for examples
exCNV <- cnvData$ID %in% paste0("P", 1:150)
exCOV <- covData$ID %in% paste0("P", 1:150)
exPHE <- phenoData$ID %in% paste0("P", 1:150)</pre>
# binary phenoType
results <- concur(cnv = cnvData[exCNV,],
                  X = covData[exCOV,],
                  pheno = phenoData[exPHE,],
                  phenoY = 'PHEB',
                  phenoType = 'bin',
                  nCore = 1L,
                  outFileKernel = NULL,
                  verbose = TRUE)
# continuous phenoType
results <- concur(cnv = cnvData[exCNV,],
                  X = covData[exCOV,],
                  pheno = phenoData[exPHE,],
                  phenoY = 'PHEC',
                  phenoType = 'cont',
                  nCore = 1L,
                  outFileKernel = NULL,
                  verbose = TRUE)
```

covData

Pseudo Covariate Data

# **Description**

This data set includes simulated covariate data. These data were generated as draws from a Binom(1,0.5) distribution for the 800 individuals in the example data provided with the package. These data are not meaningful and are intended for demonstration purposes only.

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#### Usage

data(cnvData)

# **Format**

covData is a data.frame containing 400 observations with 2 columns

**ID** character patient identifier.

**SEX** binary indicator of M/F.

#### References

Brucker, A., Lu, W., Marceau West, R., Yu, Q-Y., Hsiao, C. K., Hsiao, T-H., Lin, C-H., Magnusson, P. K. E., Holloway, S. T., Sullivan, P. F., Szatkiewicz, J. P., Lu, T-P., and Tzeng, J-Y. Association testing using Copy Number Profile Curves (CONCUR) enhances power in copy number variant analysis. <doi:10.1101/666875>.

phenoData

Pseudo Phenotype Data

# Description

This data set includes simulated phenotype data. These data include a binary phenotype and a normally distributed continuous phenotype that are randomly generated independent of the CNV data. These data are not meaningful and are intended for demonstration purposes only.

# Usage

data(cnvData)

# **Format**

phenoData is a data.frame containing 400 observations with 3 columns

**ID** character patient identifier.

PHEB binary phenotype.

PHEC continuous phenotype.

#### References

Brucker, A., Lu, W., Marceau West, R., Yu, Q-Y., Hsiao, C. K., Hsiao, T-H., Lin, C-H., Magnusson, P. K. E., Holloway, S. T., Sullivan, P. F., Szatkiewicz, J. P., Lu, T-P., and Tzeng, J-Y. Association testing using Copy Number Profile Curves (CONCUR) enhances power in copy number variant analysis. <doi:10.1101/666875>.

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