# Package 'CAnD'

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Type Package			
Title Perform Chromosomal Ancestry Differences (CAnD) Analyses			
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Imports methods, ggplot2, reshape			
Suggests RUnit, BiocGenerics, BiocStyle			
Description Functions to perform the  CAnD test on a set of ancestry proportions. For a particular ancestral subpopulation, a user will supply the estimated ancestry proportion for each sample, and each chromosome or chromosomal segment of interest. A p-value for each chromosome as well as an overall CAnD p-value will be returned for each test. Plotting functions are also available.			
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ancestries barPlotAncest BonfCorr-methods calc_combP CAnD CAnDResult-class getDiffMatrices overallpValue-methods overallStatistic-methods plotPvals			

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ancestries		Sample Chromosomal Proportions of Ancestral Subpopulations to Use for CAnD Testing	

#### **Description**

This dataset provides simulated ancestral proportions by chromosomes for 50 samples. The samples are assumed to have three different ancestral proportions, denoted by Euro\_, Afr\_ and Asian\_. Furthermore, these proportions are included as averages across each chromosome 1-22 and the X chromosome.

## Usage

ancestries

#### **Format**

A data.frame containing 50 rows and 70 columns:

IID unique sample id

Euro\_AA simulated proportion European ancestry on each chromosome, where AA ranges from 1-22, X

Afr\_AA simulated proportion African ancestry on each chromosome, where AA ranges from 1-22, X

Asian\_AA simulated proportion Asian ancestry on each chromosome, where AA ranges from 1-22, X

#### Value

A character value of the name of the dataset.

barPlotAncest	Create a Barplot of Ancestry Proportion Estimates for Every Sample
	and a Given Chromosome or Chromosomal Region

#### **Description**

Plots ancestry proportion estimates for each sample

## Usage

```
barPlotAncest(set, order = TRUE, title = "", xlab = "Sample",
  ylab = "Ancestry Proportion", ...)
```

#### **Arguments**

set

A data. frame with columns of the proportion ancestry for a given chromosome or chromosomal region, and one row per sample (bar).

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order	A logical argument determining whether the samples should be ordered in increasing proportion of the first ancestry. Default is TRUE.
title	A character string containing the title of the plot. Default is "", a blank title.
xlab	A character vector with the label for the x-axis on the plot. Default is Sample.
ylab	A character vector holding the label for the y-axis on the plot. Default is Ancestry Proportion.
	Further arguments to be passed to the plotting methods, such as graphical parameters.

#### **Details**

Creates a barplot of ancestry proportions for each sample for a given chromosome or chromosomal region.

## Value

Creates a plot.

## Author(s)

Caitlin McHugh <mchughc@uw.edu>

## **Examples**

```
data(ancestries)
chr1 <- ancestries[,c("Euro_1","Afr_1","Asian_1")]
#barPlotAncest(chr1,title="Chr 1 Ancestry Proportions")</pre>
```

BonfCorr-methods ~~ Methods for Function BonfCorr ~~

## Description

This function returns whether the Bonferroni multiple testing correction was applied in calculating the final p-values by chromosome/chromosomal segment.

## Usage

```
BonfCorr(object)
```

## Arguments

object An object of type CAnDResult

## Value

A logical indicator of whether the Bonferroni multiple testing correction was used.

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#### Methods

Defined methods include:

Returns the whether the Bonferroni multiple testing correction was applied in calculating the final p-values for a CAnD object

#### Author(s)

```
signature(object = "CAnD") Caitlin McHugh
```

#### See Also

```
CAnDResult-class
```

#### **Examples**

```
data(ancestries)
BonfCorr( CAnD(ancestries[,c(2:5)]) )
```

calc\_combP

Calculate the Combined CAnD Test Statistic P-value

#### **Description**

Calculate the combined CAnD test statistic p-value on a set of ancestry proportions estimated for a particular ancestral subpopulation of interest

#### Usage

```
calc_combP(chrAncest)
```

## **Arguments**

chrAncest

A data frame holding the ancestral proportions; each row corresponds to a sample and each column corresponds to a chromosomal/chromosomal segment ancestry proportion. Note: only include the proportions for one ancestral population at a time.

#### Value

A vector of length two where 'statistic' is the combined CAnD statistic and 'pvalue' is it's corresponding p-value, where the combined statistic is combined over all chromosomes/chromosomal segments included in chrAncest.

#### Author(s)

Caitlin McHugh <mchughc@uw.edu>

#### References

McHugh, C., Brown, L., Thornton, T. Detecting heterogeneity in population structure across the genome in admixed populations. Genetics, 2016.

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#### **Examples**

```
data(ancestries)
euroCols <- grep("Euro",colnames(ancestries))
euro <- ancestries[,euroCols]
res <- calc_combP(euro)
res</pre>
```

CAnD

Peform the CAnD Test

#### **Description**

Perform the CAnD test on a set of ancestry proportions estimated for a particular ancestral subpopulation of interest

## Usage

```
CAnD(chrAncest, bonfCorr = TRUE)
```

## **Arguments**

chrAncest A data.frame holding the ancestral proportions; each row corresponds to a sam-

ple and each column corresponds to a chromosomal/chromosomal segment ancestry proportion. Note: only include the proportions for one ancestral popula-

tion at a time.

bonfCorr A logical argument indicating whether the p-value should be corrected for mul-

tiple testing using Bonferroni correction. The default is TRUE.

#### Value

A CAnDResult object holding the p-value for each chromosome/chromosomal segment, the overall CAnD p-value, the CAnD statistic and whether the Bonferroni multiple testing correction was used.

#### Author(s)

Caitlin McHugh <mchughc@uw.edu>

#### References

McHugh, C., Brown, L., Thornton, T. Detecting heterogeneity in population structure across the genome in admixed populations. Genetics, 2016.

```
data(ancestries)
euroCols <- grep("Euro",colnames(ancestries))
euro <- ancestries[,euroCols]
res <- CAnD(euro)
res</pre>
```

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CAnDResult-class

Class "CAnDResult"

#### **Description**

Objects of this class store results from running CAnD methods.

#### **Objects from the Class**

Objects can be created by calls of nonParam\_CAnD or CAnD.

#### **Slots**

```
{\tt test:} \ \ Object\ of\ class\ "{\tt character"}, \ the\ type\ of\ CAnD\ test\ performed
```

pValues: Object of class "numeric", the p-values for each chromosome/chromosomal segment tested

overallStatistic: Object of class "numeric", the overall CAnD test statistic for the set

overallpValue: Object of class "numeric", the overall p-value for the set

BonfCorr: Object of class "logical", whether Bonferroni multiple testing correction was applied to the p-values for each chromosome/chromosomal segment tested

#### Methods

No methods defined with class "CAnDResult" in the signature.

## Author(s)

Caitlin McHugh

#### **Examples**

```
showClass("CAnDResult")
```

 ${\tt getDiffMatrices}$ 

Calculate the Mean Ancestry Proportion Excluding Each Chromosome/Chromosomal Segment in Turn

#### **Description**

A helper function to calculate the mean ancestry proportion for a given subpopulation, excluding each chromosome/chromosomal segment in turn.

## Usage

```
getDiffMatrices(chrAncest, diff = TRUE)
```

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#### **Arguments**

chrAncest A data.frame holding the ancestral proportions; each row corresponds to a sam-

ple and each column corresponds to a chromosomal/chromosomal segment ancestry proportion. Note: only include the proportions for one ancestral popula-

tion at a time.

diff A logical argment indicating whether the difference between the pooled mean

and the chromosomal mean should be returned, or whether simply the pooled

mean should be returned.

#### **Details**

This function calculates the mean ancestry proportion of a given subpopulation excluding each chromosome in turn.

#### Value

A matrix of chromosomal ancestry differences.

## Author(s)

Caitlin McHugh <mchughc@uw.edu>

overallpValue-methods ~~ Methods for Function overallpValue ~~

## Description

This function returns the CAnD test p-value from performing a CAnD test on a set of ancestral proportions.

#### Usage

overallpValue(object)

## **Arguments**

object An object of type CAnDResult

#### Value

The CAnD p-value.

#### Methods

Defined methods include:

Returns the CAnD test p-value from applying the CAnD test to a set of ancestral proportions for a CAnD object

#### Author(s)

```
signature(object = "CAnD") Caitlin McHugh
```

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#### See Also

```
CAnDResult-class
```

### **Examples**

```
data(ancestries)
overallpValue( CAnD(ancestries[,c(2:5)]) )
```

overallStatistic-methods

~~ Methods for Function overallStatistic ~~

## Description

This function returns the calculated CAnD test statistic from applying the CAnD test to a set of ancestral proportions.

## Usage

```
overallStatistic(object)
```

#### **Arguments**

object

An object of type CAnDResult

#### Value

The CAnD test statistic stored in the object.

#### Methods

Defined methods include:

Returns the statistic calculated from performing the CAnD test on a set of ancestral proportions for a CAnD object

## Author(s)

```
signature(object = "CAnD") Caitlin McHugh
```

## See Also

```
CAnDResult-class
```

```
data(ancestries)
overallStatistic( CAnD(ancestries[,c(2:5)]) )
```

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plotPvals	Create a Plot of P-Values for Each Chromosome or Chromosomal Region

## Description

Plots CAnD p-values for each chromosome/chromosomal region

## Usage

```
plotPvals(set, title = "", xlab = "Chromosome", ylab = "-log10(PValue)",
...)
```

## Arguments

set	An object of class CAnDResult.
title	A character string containing the title of the plot. Default is "", a blank title.
xlab	A character vector with the label for the x-axis on the plot. Default is Chromosome.
ylab	A character vector holding the label for the y-axis on the plot. Default is -log10(Bonferroni PValue) or -log10(PValue), depending on whether Bonferroni correction was used.
•••	Further arguments to be passed to the plotting methods, such as graphical parameters.

## **Details**

Creates a plot of all p-values for each chromosome or chromosomal region.

#### Value

Creates a plot.

## Author(s)

Caitlin McHugh <mchughc@uw.edu>

```
data(ancestries)
euroEsts <- ancestries[,c(seq(from=2,to=24))]
res <- CAnD(euroEsts)
#plotPvals(res,main="CAnD P-Values")</pre>
```

pValues-methods

pValues-methods

~~ Methods for Function pValues ~~

## Description

This function returns each p-value calculated from the CAnD test on all chromosomes/chromosomal segments.

## Usage

```
pValues(object)
```

## **Arguments**

object

An object of type CAnDResult

#### Value

A vector of p-values from the CAnD test.

#### Methods

Defined methods include:

Returns the p-values for each chromosome/chromosomal segment calculated using the CAnD test for a CAnD object

#### Author(s)

```
signature(object = "CAnD") Caitlin McHugh
```

## See Also

```
CAnDResult-class
```

```
data(ancestries)
pValues( CAnD(ancestries[,c(5:12)]) )
```

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test-methods

~~ Methods for Function test ~~

## **Description**

This function returns the type of CAnD test applied to a set of ancestral proportions.

## Usage

```
test(object)
```

#### **Arguments**

object

An object of type CAnDResult

#### Value

'Parametric.' Note the non-parametric CAnD test is defunct now.

#### Methods

Defined methods include:

Returns the type of CAnD test applied to calculate the p-values for a CAnD object

## Author(s)

```
signature(object = "CAnD") Caitlin McHugh
```

#### See Also

```
CAnDResult-class
```

```
data(ancestries)
test( CAnD(ancestries[,c(2:5)]) )
```

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