# Package 'ABHgenotypeR'

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<b>Description</b> Easy to use functions to visualize marker data from biparental populations. Useful for both analyzing and presenting genotypes in the ABH format.
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## Description

The ABHgenotypeR packages is meant as a companion packages in between the TASSEL GBS pipeline and R/qtl. It allows easy visualization of ABH-encoded genotypes in a .csv format as outtut by the TASSEL ABHGenosPlugin which is also the format necessary for R/qtl input.

correctStretches

Correct short miscalled stretches based on flanking alleles.

## **Description**

Correct short miscalled stretches based on flanking alleles.

## Usage

```
correctStretches(inputGenos = "genotypes", maxHapLength = 1)
```

## **Arguments**

inputGenos A genotypes list object.

maxHapLength The maximum length of stretches flanked by non-heterzygous sites that are

changed. If set to 1 (default) only AXA or BXB will be corrected. If set to

2, both AXA and AXYA (or BXB and BXYB) will be corrected.

#### Value

A genotype object in which short miscalled stretches are corrected if both flanking alleles match.

#### **Examples**

```
\#\# Not run: corrStretchGenos <- correctStretches(genotypes, maxHapLength = 3)
```

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correctUndercalledHets

Correct undercalled heterozygous sites based on flanking alleles.

## **Description**

Correct undercalled heterozygous sites based on flanking alleles.

## Usage

```
correctUndercalledHets(inputGenos = "genotypes", maxHapLength = 1)
```

#### **Arguments**

inputGenos A genotypes list object.

maxHapLength The maximum length of not heterozygous stretches flanked by heterzygous sites

that are changed to heterozygous. If set to 1 (default) only HAH or HBH will be corrected. If set to 2, both HAH and HAAH (or HBH and HBBH) will be

corrected.

#### Value

A genotype object in which undercalled heterozygous sites are corrected if both flanking alleles match.

## **Examples**

```
## Not run: corrUndHetsGenos <- correctUndercalledHets(genotypes, maxHapLength = 3)</pre>
```

imputeByFlanks

Impute missing genotypes based on flanking alleles

## **Description**

Impute missing genotypes based on flanking alleles

## Usage

```
imputeByFlanks(inputGenos = "genotypes")
```

## Arguments

inputGenos A genotypes list object.

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#### Value

A genotype object in which missing data is imputed based on flanking alleles. Any number of N is replaced by either A, B or N if the alleles which flank the N match

#### **Examples**

```
## Not run: imputedGenos <- imputeByFlanks(genotypes)</pre>
```

plotAlleleFreq

Plot the parental allele frequencies along the chromosomes.

#### **Description**

Plot the parental allele frequencies along the chromosomes.

## Usage

```
plotAlleleFreq(genos = "genotypes")
```

#### **Arguments**

genos

The output of readABHgenotypes

#### Value

A plot of parental allele frequencies along the chromosomes. If the output is assigned a name a ggplot2 object is returned for further manipulation.

## **Examples**

```
## Not run: plotAlleleFreq(genotypes)
## Not run: p <- plotAlleleFreq(genotypes)</pre>
```

plotCompareGenos

Compare to genotype matrices

## Description

Compare to genotype matrices

## Usage

```
plotCompareGenos(genos_1 = "genotypes_1", genos_2 = "genotypes_2",
   markerToPlot = "all", individualsToPlot = "all", chromToPlot = "all",
   CompColors = c("#000000", "#E69F00"), textSize = 12,
   showMarkerNames = FALSE, showIndividualNames = FALSE)
```

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## **Arguments**

genos_1	Output of readABHgenotypes								
genos_2	Output of readABHgenotypes. Note that both genos object need to have identical numbers of marker x individuals.								
markerToPlot	A character vector of marker names which appear in the plot. Defaults to all.								
individualsToPlot									
	A character vector of individual names which appear in the plot. Defaults to all.								
chromToPlot	A character vector of chromosome names which appear in the plot. Defaults to all.								
CompColors	A character vector of length 2 giving the color names or values to use for differnt and identical markers. Defaults to black and orange.								
textSize	The size of all text elements in the plot. Useful for making a nice plot. Defaults to 12.								
showMarkerNames									
	Show the marker names along the x axis. This and showIndividualnames are useful when you display only a few markers and want them labeled. Defaults to FALSE.								
showIndividualNames									
	Show individual names along the y axis.								

## Value

A graphical comparison of genotypes.

## **Examples**

```
## Not run: plotCompareGenos(preImpGenotypes,postImpGenotypes)
## Not run: #for more examples see plotGenos()
```

plotGenos	Plot graphical genotypes.

## Description

Plot graphical genotypes.

## Usage

```
plotGenos(genos = "genotypes", markerToPlot = "all",
  individualsToPlot = "all", chromToPlot = "all",
  alleleColors = c("#56B4E9", "#E69F00", "#009E73", "#000000"),
  textSize = 12, showMarkerNames = FALSE, showIndividualNames = FALSE)
```

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#### **Arguments**

genos The output of readABHgenotypes

markerToPlot A character vector of marker names which appear in the plot. Defaults to all.

individualsToPlot

A character vector of individual names which appear in the plot. Defaults to all.

chromToPlot A character vector of chromosome names which appear in the plot. Defaults to

all.

alleleColors A character vector of length 4 giving the color names or values to use for the

A,B,H and n.d genotypes. Defaults to orange, blue, green and black.

textSize The size of all text elements in the plot. Useful for making a nice plot. Defaults

to 12.

showMarkerNames

Show the marker names along the x axis. This and showIndividualNames are useful when you display only a few markers and want them labeled. Defaults to

FALSE.

showIndividualNames

Show individual names along the y axis.

#### Value

Graphical genotypes.

#### **Examples**

```
## Not run: plotGenos(genotypes)
markerNames <- c("marker1", "marker2", "marker3")
individualNames <- c("F2_100", "F2_101", "F2_102", "F2_103")
someColors <- c("black", "red", "gold", "white")
## Not run: plotgenos(genotypes, markerNames, individualNames, 1:3, someColors)
## Not run: p <- plotGenos(genotypes)</pre>
```

plotMarkerDensity

Plot the marker density along the chromosomes.

## Description

Plot the marker density along the chromosomes.

#### Usage

```
plotMarkerDensity(genos = "genotypes")
```

#### **Arguments**

genos

The output of readABHgenotypes

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#### Value

A plot of marker densities along the chromosomes. If the output is assigned a name a ggplot2 object is returned for further manipulation.

## **Examples**

```
## Not run: plotMarkerDensity(genotypes)
## Not run: p <- plotMarkerDensity(genotypes)</pre>
```

readABHgenotypes

Read in the output of the genosToABH plugin.

#### **Description**

Read in the output of the genosToABH plugin.

#### Usage

```
readABHgenotypes(pathToABH, nameA = "A", nameB = "B", readPos = TRUE)
```

#### Arguments

pathToABH The path and filename of the input file.

nameA Name of the parent represented by "A" in the input file.

Name of the parent represented by "B" in the input file.

readPos Should the function attempt read the physical position of markers from the input

?

#### **Details**

The input files should be a .csv file holding genotypes as specified by the qtl package and its "csvs" format. All characters in the genotype matrix which are not either A,B or H will be set to N. If readPos = TRUE (default) marker names must conform to S1\_123456 meaning 123456 bp on chromosome 1. If FALSE, pos is set to NULL and needs to be manually constructed as shown in the examples. Note that this might throw off some plotting function.

## Value

A genotype list object which holds the information from the input file. This list is the fundamental datastructure used by the other functions in this package. See the vignette for what each item in the list is.

#### **Examples**

```
## Not run: genotypes <- readABHgenotypes("./genotypes.csv", "NB", "OL")

## Not run: otherGenotypes <- readABHgenotypes("./otherGenotypes.csv", readPos = FALSE)
#arbitrary position to keep marker order intact

## Not run: therGenotypes$pos <- 1:length(otherGenotypes$marker_names)</pre>
```

writeABHgenotypes

reportGenos

Report the total and relative number of each allele in a genotype ob-

ject.

## **Description**

Report the total and relative number of each allele in a genotype object.

#### Usage

```
reportGenos(genos1)
```

#### **Arguments**

genos1

A genotypes list object.

#### Value

Console output of the total an relative number of each allele.

## **Examples**

```
## Not run: reportGenos(preImputation)
```

writeABHgenotypes

Export a genotype list to a .csv file.

#### **Description**

Export a genotype list to a .csv file.

#### Usage

```
writeABHgenotypes(genos = "genotypes", outfile = "./outfile.csv")
```

#### **Arguments**

genos The output of readABHgenotypes or one of the impuation/error correction func-

tions.

outfile The path and filename of the output file.

#### Value

A file which can be used in R/qtl or elsewhere.

#### **Examples**

```
## Not run: writeABHgenotypes(genotypes, outfile = "./outfile_name.csv")
```

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