Package 'QCpipeline'

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Description Configuration and plotting code
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bestLegendPos	Best legend position
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Description

Find the section of a plot with the fewest number of points for placing a legend

Usage

```
bestLegendPos(x, y)
```

Arguments

```
x Points on the x axis
y Points on the y axis
```

Value

```
Returns one of the legend placement strings from the list "bottomright", "bottom", "bottomleft", "left", "topleft", "top", "topright", "right" and "center".
```

Author(s)

Stephanie Gogarten

Examples

```
x <- sample(1:100, 50, replace=TRUE)
y <- sample(1:100, 50, replace=TRUE)
plot(x,y)
legend(bestLegendPos(x,y), "points", pch=1)</pre>
```

boxplotMeanSD

Boxplot with mean and SD

Description

Boxplot with mean and SD

Usage

```
boxplotMeanSD(x, y, data=NULL, xlab=NULL, ylab=NULL, nSD=1, ...)
```

Arguments

X	vector or character string denoting column in data
у	vector or character string denoting column in data
data	data.frame
xlab	title for x axis (defaults to x if data is not NULL
ylab	title for y axis (defaults to y if data is not NULL
nSD	number of standard deviations to plot
	additional plotting arguments

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Author(s)

Jess Shen

Examples

```
age <- sample(25:55, 100, replace=TRUE)
sex <- sample(c("M", "F"), 100, replace=TRUE)
boxplotMeanSD(sex, age)

data <- data.frame(age, sex)
boxplotMeanSD("sex", "age", data)</pre>
```

dbgapAnnotation

Write annotation files for dbGaP

Description

dbgapScanAnnotation and dbgapSnpAnnotation create text files appropriate for posting on db-GaP.

Usage

```
dbgapScanAnnotation(scanAnnot, dir=".",
   consentVar="consent", subjVar="subj.plink", dupVar="dup.post", omitVar="no.post",
   annotationCol="annotation", analysisCol="analysis")

dbgapSnpAnnotation(snpAnnot, dir=".",
   annotationCol="annotation", analysisCol="analysis")
```

Arguments

 $\begin{array}{lll} & & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$

dir A character string with the directory for file output.

consentVar The variable in scanAnnot containing consent levels.

subjVar The logical variable in scanAnnot indicating unique subjects to post.

dupVar The logical variable in scanAnnot indicating duplicate scans to post.

omitVar The logical variable in scanAnnot indicating scans to be omitted from posting.

annotationCol The logical column in the metadata indicating which variables should be in-

cluded in the annotation files.

analysisCol The logical column in the metadata indicating which variables should be in-

cluded in the analysis files.

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Details

dbgapScanAnnotation writes the following files to dir:

- Sample_annotation.csv
- Sample_annotation_consent_*.csv
- Sample_annotation_duplicates.csv
- Sample_annotation_duplicates_consent_*.csv
- Sample_annotation_DD.txt
- · Sample_analysis.csv
- · Sample_analysis_duplicates.csv
- Sample_analysis_DD.txt

dbgapSnpAnnotation writes the following files to dir:

- SNP_annotation.csv
- SNP_annotation_DD.txt
- · SNP_analysis.csv
- SNP_analysis_DD.txt

Which variables should be written to the annotation and analysis files are indicated in the metadata columns annotationCol and analysisCol.

The data dictionary files are populated from the metadata. The "type" column is automatically generated from the classes of the variables in scanAnnot and snpAnnot.

Author(s)

Stephanie Gogarten

ideogram

BAF/LRR plots with chromosome ideograms

Description

Plot BAF/LRR with chromosome ideograms at the bottom.

Usage

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Arguments

intenData	IntensityData object
genoData	GenotypeData object
scan.ids	A vector containing the sample indices of the plots.
chrom.ids	A vector containing the chromosome indices of the plots.
anom.stats	data.frame of chromosome anomalies with statistics, usually the output of anomSegStats. Names must include "anom.id", "scanID", "chromosome", "left.index", "right.index", "method", "nmark.all", "nmark.elig", "left.base", "right.base", "nbase", "non.anom.baf.med", "non.anom.lrr.med", "anom.baf.dev.med", "anom.baf.dev.5", "anom.lrr.med", "nmark.baf", "nmark.lrr". Left and right refer to start and end, respectively, of the anomaly, in position order.
snp.ineligible	vector of ineligible snp ids (e.g., intensity-only, failed snps, XTR and HLA regions). See HLA and pseudoautosomal.
win	size of the window (a multiple of anomaly length) surrounding the anomaly to plot
main	Vector of plot titles. If NULL then the title will include scanID, sex, and chromosome.
info	A character vector containing extra information to include in the main title.
	additional arguments to chromIntensityPlot or anomStatsPlot
ideo.zoom	logical for whether to zoom in on the ideogram to match the range of the BAF/LRR plots
ideo.rect	logical for whether to draw a rectangle on the ideogram indicating the range of the BAF/LRR plots
cex.axis	cex value for the axis
cex.lab	cex value for the labels
cex.main	cex value for the title
cex.sub	cex value for the subtitle
cex.leg	cex value for the ideogram legend

Details

 $chrow Intensity PlotI deogram is a \ wrapper for chrow Intensity Plot. \ anom Stats PlotI deogram is a \ wrapper for a nom Stats Plot.$

These functions call paintCytobands to draw ideograms.

Author(s)

Stephanie Gogarten

See Also

chrom Intensity Plot, anom Stats Plot, paint Cytobands

Examples

```
library(GWASdata)
data(illuminaScanADF)
data(illuminaSnpADF)
blfile <- system.file("extdata", "illumina_bl.nc", package="GWASdata")</pre>
blnc <- NcdfIntensityReader(blfile)</pre>
intenData <- IntensityData(blnc, scanAnnot=illuminaScanADF, snpAnnot=illuminaSnpADF)</pre>
genofile <- system.file("extdata", "illumina_geno.nc", package="GWASdata")</pre>
genonc <- NcdfGenotypeReader(genofile)</pre>
genoData <- GenotypeData(genonc, scanAnnot=illuminaScanADF, snpAnnot=illuminaSnpADF)</pre>
\hbox{\tt\#} chromIntensityPlotIdeogram}
scanID <- getScanID(illuminaScanADF, index=1:2)</pre>
#png("tmp_%03d.png", width=720, height=900)
main <- paste("Sample", c("A", "B"), "- Chromosome X")</pre>
chromIntensityPlotIdeogram(intenData=intenData, scan.ids=scanID,
                             chrom.ids=c(23,23), main=main,
                             colorGenotypes=TRUE, genoData=genoData)
#dev.off()
# anomStatsPlotIdeogram
scan.ids <- illuminaScanADF$scanID[1:2]</pre>
chrom.ids <- unique(illuminaSnpADF$chromosome)</pre>
snp.ids <- illuminaSnpADF$snpID[illuminaSnpADF$missing.n1 < 1]</pre>
snp.failed <- illuminaSnpADF$snpID[illuminaSnpADF$missing.n1 == 1]</pre>
# example results from anomDetectBAF
baf.anoms <- data.frame("scanID"=scan.ids[1],"chromosome"=21,</pre>
  "left.index"=100, "right.index"=200, sex="M", method="BAF",
  anom.id=1, stringsAsFactors=FALSE)
# example results from anomDetectLOH
loh.anoms <- data.frame("scanID"=scan.ids[2],"chromosome"=22,</pre>
  "left.index"=400,"right.index"=500, sex="F", method="LOH",
  anom.id=2, stringsAsFactors=FALSE)
anoms <- rbind(baf.anoms, loh.anoms)</pre>
data(centromeres.hg18)
stats <- anomSegStats(intenData, genoData, snp.ids=snp.ids, anom=anoms,</pre>
  centromere=centromeres.hg18)
#png("tmp_%03d.png", width=720, height=900)
main <- paste("Sample", c("A", "B"), "- Chromosome", stats$chromosome)</pre>
anomStatsPlotIdeogram (intenData, \ genoData, \ anom.stats=stats,
  snp.ineligible=snp.failed, centromere=centromeres.hg18, main=main)
#dev.off()
close(genoData)
close(intenData)
```

make.allele.mappings 7

Description

Function to take Illumina build 37 SNP annotation data file and make an Allele Mappings table

Usage

```
make.allele.mappings(snp.dat)
make.allele.annotation(map, alleles=c("top", "design", "fwd", "plus"))
```

Arguments

snp.dat	a data frame made from SNP Illumina annotation file (i.e., "HumanOmni2.5-4v1_D.csv"), with following fields: "IlmnID", "Name", "IlmnStrand", "SNP", "GenomeBuild", "SourceStrand", "SourceSeq", "TopGenomicSeq", "RefStrand"
map	a data frame with allele mappings (output of make.allele.mappings), with following fields: "snp", "alle.AB", "alle.design", "alle.top", "alle.fwd", "alle.plus"
alleles	character string with type of alleles to return in the annotation table ("top", "design", "fwd", or "plus")

Details

In the case of indels, the make.allele.mappings codes A or B as "-" and the other allele with the insertion/deletion sequence. make.allele.annotation codes indels as "D" and "I" (useful for making PLINK files).

Note that "RefStrand" in build 37 annotations from Illumina is result of BLAST search of DESIGN strand. In previous strand annotations from Illumina for build 36 arrays, "BlastStrand" held +/-result of BLAST search of the SOURCE sequence.

Value

```
make.allele.mappings returns a data frame object with columns ("snp", "alle.AB", "alle.design", "alle.top", "alle.fwd", "alle.plus").

make.allele.annotation returns a data frame with columns "snp", "alleleA.*", "alleleB.*" where
"*"=alleles.
```

Create a data frame with one row per SNP, TOP alleles, and indels coded as D/I

snp.annot <- make.allele.annotation(map.final, alleles="top")</pre>

Author(s)

Sarah Nelson

Examples

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```
## Example allele mappings table:
                 snp alle.AB alle.design alle.top alle.fwd alle.plus
#
 rs1000000
                 Α
                           Т
                                   Α
                                             Τ
#
 rs1000000
                  В
                             С
                                     G
                                              С
                                                        G
# rs1000002
                 Α
                            Α
                                     Α
                                              Α
                                                       Т
# rs1000002
                  В
                             G
                                     G
                                              G
                                                       С
                                              Т
# rs10000023
                             Т
                                     Α
                                                        Т
                  Α
# rs10000023
                  В
                                      С
                                              G
                                                        G
## With the following data dictionary:
# variable
                type
                         description
        text rs id and other comparable identifiers for a snp
# snp
# alle.AB text A or B, per the Illumina genotyping system nomenclature
# alle.design text
                        nucleotide(s) corresponding to A or B allele for design strand
# alle.top text nucleotide(s) corresponding to A or B allele for Illumina TOP strand
# alle.fwd
              text nucleotide(s) corresponding to A or B allele for FORWARD strand, with respect to dbSNP re
# alle.plus
                         \operatorname{nucleotide}(s) corresponding to A or B allele
               text
# for PLUS(+) strand, relative to the forward direction in the human
# reference genome sequence
## End(Not run)
```

Description

plot2DwithHist

plot2DwithHist produces a scatterplot of y vs x, along with histograms of the marginal distributions of x and y.

Usage

```
plot2DwithHist(x, y, xlab=NULL, ylab=NULL, xlim=NULL, ylim=NULL,
    sublab=NULL, mn=NULL, sd=NULL, ...)
```

Scatterplot with density

Arguments

X	vector of x coordinates
у	vector of y coordinates
xlab	x-axis label (defaults to variable name)
ylab	y-axis label (defaults to variable name)
xlim	x-axis limits (defaults to [min,max] of X, plus a bit of space
ylim	y-axis limits (defaults to [min,max] of Y, plus a bit of space
sublab	sub-label (instead of main, since there's no room)
mn	2-element vector with mean of x and y
sd	2-element vector with sd of x and y
	additional arguments to pass to points

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Author(s)

Leila Zelnick

Examples

```
library(MSBVAR)
# generate some multivariate normal example data
n <- 5000
mu <- c(0, 2)
vmat <- matrix(c(1, 0.7, 0.7, 1), nrow=2)

dat <- rmultnorm(n, mu, vmat) # generates n multivariate normal obs.
x <- dat[,1]
y <- dat[,2]

plot2DwithHist(x, y, xlab="This is the X variable", ylab="This is the Y variable.",
    sub="Example Plot!")
# defining axis limits
plot2DwithHist(x, y, xlab="This is the X variable", ylab="This is the Y variable.",
    sub="Example Plot!", xlim=c(0,4), ylim=c(-2,2))</pre>
```

readConfig

Read a configuration file

Description

Read a configuration file

Usage

```
readConfig(file, ...)
```

Arguments

file file where column 1 is parameter name and column 2 is value.
... additional arguments to read.table

Value

Returns a named character vector of parameter values.

Author(s)

Stephanie Gogarten

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