Package 'dartR.base'

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Type Package

Title Analysing 'SNP' and 'Silicodart' Data - Basic Functions

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Description Facilitates the import and analysis of 'SNP' (single nucleotide 'polymorphism') and 'silicodart' (presence/absence) data. The main focus is on data generated by 'DarT' (Diversity Arrays Technology), however, data from other sequencing platforms can be used once 'SNP' or related fragment presence/absence data from any source is imported. Genetic datasets are stored in a derived 'genlight' format (package 'adegenet'), that allows for a very compact storage of data and metadata. Functions are available for importing and exporting of 'SNP' and 'silicodart' data, for reporting on and filtering on various criteria (e.g. 'callrate', 'heterozygosity', 'reproducibility', maximum allele frequency). Additional functions are available for visualization (e.g. Principle Coordinate Analysis) and creating a spatial representation using maps. 'dartR.base' is the 'base' package of the 'dartRverse' suits of packages. To install the other packages, we recommend to install the 'dartRverse' package, that supports the installation of all packages in the 'dartRverse'.
If you want to cite 'dartR', you find the information by typing citation('dartR.base') in the console.

Encoding UTF-8

Depends R (>= 3.5), adegenet (>= 2.0.0), ggplot2, dplyr, dartR.data

Imports ape,crayon,data.table,foreach,gridExtra,methods,patchwork,plyr, reshape2,SNPRelate,StAMPP,stats,stringr,tidyr,utils,MASS,SNPassoc, snpStats, raster

Suggests boot, devtools, directlabels, dismo, doParallel, expm, gdistance, gganimate, ggrepel, grid, gtable, ggthemes, gplots, HardyWeinberg, hierfstat, igraph, iterpc, knitr, label.switching, lattice, leaflet, leaflet.minicharts, markdown, mmod, networkD3, parallel, pegas, pheatmap, plotly, poppr, proxy, purrr, qvalue, RColorBrewer, Rcpp, rgl, rmarkdown, rrBLUP, scales, seqinr, sf, shinyBS, shinyjs, shinythemes, shinyWidgets, SIBER, stringi, terra, tibble, vcfR, zoo, viridis, fields, testthat (>= 3.0.0), ggtern

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Contents

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cbind.dartR adjust cbind for dartR

Description

cbind is a bit lazy and does not take care for the metadata (so data in the other slot is lost). You can get most of the loci metadata back using gl.compliance.check.

Usage

S3 method for class 'dartR'
cbind(...)

Arguments

... list of dartR objects

Value

A genlight object

Examples

t1 <- platypus.gl
class(t1) <- "dartR"
t2 <- cbind(t1[,1:10],t1[,11:20])</pre>

gl.add.indmetrics Adds metadata into a genlight object

Description

This function adds the metadata information to the slot ind.metrics and populates population and coordinates information slots if the they are found in the metadata.

Usage

```
gl.add.indmetrics(x, ind.metafile, verbose = NULL)
```

Arguments

х	Name of the genlight object containing the SNP data, or the genind object con- taining the SilocoDArT data [required].
ind.metafile	path and name of CSV file containing the metadata information for each indi- vidual (see details for explanation) [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

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gl.alf

Details

The ind.metadata file needs to have very specific headings. First a column with a heading named 'id'. Here the ids must match the ids in the genlight object, e.g. indNames(your_genlight). The following column headings are optional:

- 'pop' specifies the population membership of each individual.
- 'lat' latitude coordinates (in decimal degrees WGS1984 format).
- 'lon' longitude coordinates (in decimal degrees WGS1984 format).

Additional columns with individual metadata can be imported (e.g. age, sex, etc).

Value

A genlight object with metadata information for each individual.

Author(s)

Custodian: Luis Mijangos – Post to https://groups.google.com/d/forum/dartr

Examples

```
dartfile <- system.file('extdata','testset_SNPs_2Row.csv', package='dartR.data')
metadata <- system.file('extdata','testset_metadata.csv', package='dartR.data')
gl <- gl.read.dart(dartfile, probar=TRUE)
gl <- gl.add.indmetrics(gl, ind.metafile = metadata)</pre>
```

gl	•	a	lf	
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Calculates allele frequency of the first and second allele for each locus A very simple function to report allele frequencies

Description

Calculates allele frequency of the first and second allele for each locus A very simple function to report allele frequencies

Usage

gl.alf(x)

Arguments

Х

Name of the genlight object [required].

Value

A simple data.frame with ref (reference allele), alt (alternate allele).

Author(s)

Bernd Gruber (bugs? Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other utilities: utils.check.datatype(),utils.collapse.matrix(),utils.dart2genlight(),
utils.dist.binary(),utils.flag.start(),utils.hamming(),utils.het.pop(),utils.impute,
utils.is.fixed(),utils.jackknife(),utils.n.var.invariant(),utils.plot.save(),utils.read.fasta(),
utils.read.ped(),utils.recalc.avgpic(),utils.recalc.callrate(),utils.recalc.freqhets(),
utils.recalc.freqhomref(),utils.recalc.freqhomsnp(),utils.recalc.maf(),utils.reset.flags(),
utils.transpose(),utils.vcfr2genlight.polyploid()
```

Examples

```
#for the first 10 loci only
#Deprecated:
gl.alf(possums.gl[,1:10])
barplot(t(as.matrix(gl.alf(possums.gl[,1:10]))))
#Current:
gl.allele.freq(possums.gl[,1:10],simple=TRUE)
barplot(t(as.matrix(gl.allele.freq(possums.gl[,1:10],simple=TRUE))))
```

gl.allele.freq Generates percentage allele frequencies by locus and population

Description

This is a support script, to take SNP data or SilicoDArT presence/absence data grouped into populations in a genlight object {adegenet} and generate a table of allele frequencies for each population and locus

Usage

```
gl.allele.freq(x, percent = FALSE, by = "pop", simple = FALSE, verbose = NULL)
```

Arguments

x	Name of the genlight object containing the SNP or Tag P/A (SilicoDArT) data [required].
percent	If TRUE, percentage allele frequencies are given, if FALSE allele proportions are given [default FALSE]
by	If by='popxloc' then breakdown is given by population and locus; if by='pop' then breakdown is given by population with statistics averaged across loci; if by='loc' then breakdown is given by locus with statistics averaged across indi- viduals [default 'pop']
simple	A legacy option to return a dataframe with the frequency of the reference allele (alf1) and the frequency of the alternate allele (alf2) by locus [default FALSE]

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gl.amova

verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress
	and results summary; 5, full report [default 2 or as specified using gl.set.verbosity]

Value

A matrix with allele (SNP data) or presence/absence frequencies (Tag P/A data) broken down by population and locus

Author(s)

Custodian: Arthur Georges (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other unmatched report: gl.report.basics(), gl.report.diversity(), gl.report.excess.het(),
gl.report.heterozygosity(), gl.report.polyploid_heterozygosity()
```

Examples

```
gl.allele.freq(testset.gl,percent=FALSE,by='pop')
gl.allele.freq(testset.gl,percent=FALSE,by="loc")
gl.allele.freq(testset.gl,percent=FALSE,by="popxloc")
gl.allele.freq(testset.gl,simple=TRUE)
```

. amov	

Performs AMOVA using genlight data

Description

This script performs an AMOVA based on the genetic distance matrix from stamppNeisD() [package StAMPP] using the amova() function from the package PEGAS for exploring within and between population variation. For detailed information use their help pages: ?pegas::amova, ?StAMPP::stamppAmova. Be aware due to a conflict of the amova functions from various packages I had to 'hack' StAMPP::stamppAmova to avoid a namespace conflict.

Usage

```
gl.amova(x, distance = NULL, permutations = 100, verbose = NULL)
```

Arguments

х	Name of the genlight containing the SNP genotypes, with population informa- tion [required].
distance	Distance matrix between individuals (if not provided NeisD from StAMPP::stamppNeisD is calculated) [default NULL].

permutations	Number of permutations to perform for hypothesis testing [default 100]. Please note should be set to 1000 for analysis.
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity]

Value

An object of class 'amova' which is a list with a table of sums of square deviations (SSD), mean square deviations (MSD), and the number of degrees of freedom, and a vector of variance components.

Author(s)

Bernd Gruber (bugs? Post to https://groups.google.com/d/forum/dartr)

Examples

```
#permutations should be higher, here set to 1 because of speed
out <- gl.amova(bandicoot.gl, permutations=1)</pre>
```

gl.check.verbosity Checks the current global verbosity

Description

The verbosity can be set in one of two ways -(a) explicitly by the user by passing a value using the parameter verbose in a function, or (b) by setting the verbosity globally as part of the r environment (gl.set.verbosity).

Usage

```
gl.check.verbosity(x = NULL)
```

Arguments

```
Х
```

User requested level of verbosity [default NULL].

Value

The verbosity, in variable verbose

Author(s)

Bernd Gruber (Post to https://groups.google.com/d/forum/dartr)

See Also

Other environment: gl.check.wd(), gl.print.history(), gl.set.wd(), theme_dartR()

gl.check.wd

Examples

gl.check.verbosity()

gl.check.wd

Checks the global working directory

Description

The working directory can be set in one of two ways -(a) explicitly by the user by passing a value using the parameter plot.dir in a function, or (b) by setting the working directory globally as part of the r environment (gl.setwd). The default is in accordance to CRAN set to tempdir().

Usage

gl.check.wd(wd = NULL, verbose = NULL)

Arguments

wd	path to the working directory [default: tempdir()].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress
	and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Value

the working directory

Author(s)

Custodian: Bernd Gruber (Post to https://groups.google.com/d/forum/dartr)

See Also

Other environment: gl.check.verbosity(), gl.print.history(), gl.set.wd(), theme_dartR()

Examples

gl.check.wd()

gl.colors

Description

Creates a vector of colors in hex (e.g. "#3B9AB2" "#78B7C5") based on user selected category (parameter type).

- "2" [two colors]
- "2c" [two colors contrast]
- "3" [three colors]
- 4" [four colors]
- "pal" [need to be specify the palette type and the number of colors]

A palette of colors can be specified via "div" [divergent], "dis" [discrete], "con" [convergent], "vir" [viridis]. Be aware a palette needs the number of colors specified as well. It returns a function and therefore the number of colors needs to be a part of the function call. Check the examples to see how this works.

Usage

gl.colors(type = 2, verbose = NULL)

Arguments

type	Type of color (2, 3 or 4 colors, or palette, see description) [default 2].
verbose	- verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

returns colors as a vector

Author(s)

Custodian: Bernd Gruber - Post to https://groups.google.com/d/forum/dartr

See Also

```
Other graphics: gl.map.interactive(), gl.plot.heatmap(), gl.report.ld.map(), gl.select.colors(),
gl.select.shapes(), gl.smearplot(), gl.tree.nj()
```

gl.compliance.check

Examples

```
gl.colors(2)
gl.colors("2")
gl.colors("2c")
#five discrete colors
gl.colors(type="dis")(5)
#seven divergent colors
gl.colors("div")(7)
```

gl.compliance.check Checks a genlight object to see if it complies with dartR expectations and amends it to comply if necessary @family environment

Description

This function will check to see that the genlight object conforms to expectation in regard to dartR requirements (see details), and if it does not, will rectify it.

Usage

gl.compliance.check(x, verbose = NULL)

Arguments

Х	Name of the input genlight object [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress
	and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Details

A genlight object used by dartR has a number of requirements that allow functions within the package to operate correctly. The genlight object comprises:

- 1. The SNP genotypes or Tag Presence/Absence data (SilicoDArT);
- An associated dataframe (gl@other\$loc.metrics) containing the locus metrics (e.g. Call Rate, Repeatability, etc);
- 3. An associated dataframe (gl@other\$ind.metrics) containing the individual/sample metrics (e.g. sex, latitude (=lat), longitude(=lon), etc);
- 4. A specimen identity field (indNames(gl)) with the unique labels applied to each individual/sample;
- 5. A population assignment (popNames) for each individual/specimen;
- 6. Flags that indicate whether or not calculable locus metrics have been updated.

Value

A genlight object that conforms to the expectations of dartR

Author(s)

Custodian: Luis Mijangos – Post to https://groups.google.com/d/forum/dartr

Examples

x <- gl.compliance.check(testset.gl)
x <- gl.compliance.check(testset.gs)</pre>

gl.define.pop Defines a new population in a genlight object for specified individuals

Description

The script reassigns existing individuals to a new population and removes their existing population assignment. The script returns a genlight object with the new population assignment.

Usage

gl.define.pop(x, ind.list, new, verbose = NULL)

Arguments

х	Name of the genlight object containing SNP genotypes [required].
ind.list	A list of individuals to be assigned to the new population [required].
new	Name of the new population [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

A genlight object with the redefined population structure.

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

```
Other data manipulation: gl.drop.ind(),gl.drop.loc(),gl.drop.pop(),gl.edit.recode.pop(),
gl.impute(),gl.join(),gl.keep.ind(),gl.keep.loc(),gl.keep.pop(),gl.make.recode.ind(),
gl.merge.pop(),gl.reassign.pop(),gl.recode.ind(),gl.recode.pop(),gl.rename.pop(),
gl.sample(),gl.sim.genotypes(),gl.sort(),gl.subsample.ind(),gl.subsample.loc()
```

gl.diagnostics.hwe

Examples

```
popNames(testset.gl)
gl <- gl.define.pop(testset.gl, ind.list=c('AA019073','AA004859'),
new='newguys')
popNames(gl)
indNames(gl)[pop(gl)=='newguys']</pre>
```

gl.diagnostics.hwe Provides descriptive stats and plots to diagnose potential problems with Hardy-Weinberg proportions @family matched report

Description

Different causes may be responsible for lack of Hardy-Weinberg proportions. This function helps diagnose potential problems.

Usage

```
gl.diagnostics.hwe(
    x,
    alpha_val = 0.05,
    bins = 20,
    stdErr = TRUE,
    colors.hist = gl.colors(2),
    colors.barplot = gl.colors("2c"),
    plot.theme = theme_dartR(),
    n.cores = "auto",
    plot.file = NULL,
    plot.dir = NULL,
    verbose = NULL
)
```

Arguments

x	Name of the genlight object containing the SNP data [required].
alpha_val	Level of significance for testing [default 0.05].
bins	Number of bins to display in histograms [default 20].
stdErr	Whether standard errors for Fis and Fst should be computed (default: TRUE)
colors.hist	List of two color names for the borders and fill of the histogram [default gl.colors(2)].
colors.barplot	Vector with two color names for the observed and expected number of significant HWE tests [default gl.colors("2c")].
plot.theme	User specified theme [default theme_dartR()].
n.cores	The number of cores to use. If "auto", it will use all but one available cores [default "auto"].

plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL]
plot.dir	Directory in which to save files [default = working directory]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity]

Details

This function initially runs gl.report.hwe and reports the ternary plots. The remaining outputs follow the recommendations from Waples (2015) paper and De Meeûs 2018. These include:

- 1. A histogram with the distribution of p-values of the HWE tests. The distribution should be roughly uniform across equal-sized bins.
- 2. A bar plot with observed and expected (null expectation) number of significant HWE tests for the same locus in multiple populations (that is, the x-axis shows whether a locus results significant in 1, 2, ..., n populations. The y axis is the count of these occurrences. The zero value on x-axis shows the number of non-significant tests). If HWE tests are significant by chance alone, observed and expected number of HWE tests should have roughly a similar distribution.
- 3. A scatter plot with a linear regression between Fst and Fis, averaged across subpopulations. De Meeûs 2018 suggests that in the case of Null alleles, a strong positive relationship is expected (together with the Fis standard error much larger than the Fst standard error, see below). **Note**, this is not the scatter plot that Waples 2015 presents in his paper. In the lower right corner of the plot, the Pearson correlation coefficient is reported.
- 4. The Fis and Fst (averaged over loci and subpopulations) standard errors are also printed on screen and reported in the returned list (if stdErr=TRUE). These are computed with the Jack-knife method over loci (See De Meeûs 2007 for details on how this is computed) and it may take some time for these computations to complete. De Meeûs 2018 suggests that under a global significant heterozygosity deficit: if the correlation between Fis and Fst is strongly positive, and StdErrFis » StdErrFst, Null alleles are likely to be the cause. if the correlation between Fis and Fst is ~0 or mildly positive, and StdErrFis > StdErrFst, Wahlund may be the cause. if the correlation between Fis and Fst is or sib mating could to be the cause. It is important to realise that these statistics only suggest a pattern (pointers). Their absence is not conclusive evidence of the absence of the problem, as their presence does not confirm the cause of the problem.
- 5. A table where the number of observed and expected significant HWE tests are reported by each population, indicating whether these are due to heterozygosity excess or deficiency. These can be used to have a clue of potential problems (e.g. deficiency might be due to a Wahlund effect, presence of null alleles or non-random sampling; excess might be due to sex linkage or different selection between sexes, demographic changes or small Ne. See Table 1 in Wapples 2015). The last two columns of the table generated by this function report chisquare values and their associated p-values. Chisquare is computed following Fisher's procedure for a global test (Fisher 1970). This basically tests whether there is at least one test that is truly significant in the series of tests conducted (De Meeûs et al 2009).

gl.dist.ind

Value

A list with the table with the summary of the HWE tests and (if stdErr=TRUE) a named vector with the StdErrFis and StdErrFst.

Author(s)

Custodian: Carlo Pacioni – Post to https://groups.google.com/d/forum/dartr

References

- de Meeûs, T., McCoy, K.D., Prugnolle, F., Chevillon, C., Durand, P., Hurtrez-Boussès, S., Renaud, F., 2007. Population genetics and molecular epidemiology or how to "débusquer la bête". Infection, Genetics and Evolution 7, 308-332.
- De Meeûs, T., Guégan, J.-F., Teriokhin, A.T., 2009. MultiTest V.1.2, a program to binomially combine independent tests and performance comparison with other related methods on proportional data. BMC Bioinformatics 10, 443-443.
- De Meeûs, T., 2018. Revisiting FIS, FST, Wahlund Effects, and Null Alleles. Journal of Heredity 109, 446-456.
- Fisher, R., 1970. Statistical methods for research workers Edinburgh: Oliver and Boyd.
- Waples, R. S. (2015). Testing for Hardy–Weinberg proportions: have we lost the plot?. Journal of heredity, 106(1), 1-19.

See Also

gl.report.hwe

Examples

```
require("dartR.data")
res <- gl.diagnostics.hwe(x = gl.filter.allna(platypus.gl[,1:50]),
stdErr=FALSE, n.cores=1)</pre>
```

gl.dist.ind	Calculates a distance matrix for individuals defined in a genlight ob-
	ject

Description

Calculates various distances between individuals based on allele frequencies or presence-absence data

Usage

```
gl.dist.ind(
    x,
    method = NULL,
    scale = FALSE,
    swap = FALSE,
    type = "dist",
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
    plot.file = NULL,
    verbose = NULL
)
```

Arguments

х	Name of the genlight [required].
method	Specify distance measure [SNP: Euclidean; P/A: Simple].
scale	If TRUE, the distances are scaled to fall in the range [0,1] [default TRUE]
swap	If TRUE and working with presence-absence data, then presence (no disrupt- ing mutation) is scored as 0 and absence (presence of a disrupting mutation) is scored as 1 [default FALSE].
type	Specify the type of output, dist or matrix [default dist]
plot.display	If TRUE, resultant plots are displayed in the plot window [default TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.colors	List of two color names for the borders and fill of the plots [default c("#2171B5", "#6BAED6")].
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL]
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log ; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Details

The distance measure for SNP genotypes can be one of:

- Euclidean Distance [method = "Euclidean"]
- Scaled Euclidean Distance [method='Euclidean", scale=TRUE]
- Simple Mismatch Distance [method="Simple"]
- Absolute Mismatch Distance [method="Absolute"]
- Czekanowski (Manhattan) Distance [method="Manhattan"]

The distance measure for Sequence Tag Presence/Absence data (binary) can be one of:

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gl.dist.phylo

- Euclidean Distance [method = "Euclidean"]
- Scaled Euclidean Distance [method='Euclidean", scale=TRUE]
- Simple Matching Distance [method="Simple"]
- Jaccard Distance [method="Jaccard"]
- Bray-Curtis Distance [method="Bray-Curtis"]

Refer to the documentation of functions in https://doi.org/10.1101/2023.03.22.533737 for algorithms and definitions.

Value

An object of class 'matrix' or dist' giving distances between individuals

Author(s)

Author(s): Custodian: Arthur Georges – Post to #'https://groups.google.com/d/forum/dartr

See Also

Other distance: gl.dist.pop(), gl.fdsim(), utils.dist.ind.snp()

Examples

```
D <- gl.dist.ind(testset.gl[1:20,], method='manhattan')
D <- gl.dist.ind(testset.gs[1:20,], method='Jaccard',swap=TRUE)
D <- gl.dist.ind(testset.gl[1:20,], method='euclidean',scale=TRUE)</pre>
```

gl.dist.phylo	Generates a distance matrix from a SNP genlight object taking into
	account a substitution model

Description

Generates a distance matrix for individuals or populations in a genlight object using one of a selection of substitution models.

Usage

```
gl.dist.phylo(
    xx,
    subst.model = "F81",
    min.tag.len = NULL,
    pairwise.missing = TRUE,
    by.pop = TRUE,
    verbose = NULL
)
```

Arguments

xx	Name of the genlight object containing the SNP data [required].	
subst.model	The evolutionary model of nucleotide substitutions to employ in calculating ge- netic distance between individuals [default "F81"]	
min.tag.len	Minimum tag length of sequence tags to be used in the analysis [default NULL]	
pairwise.missing		
	How to handle missing sequences [default TRUE]	
by.pop	If TRUE, the distance matrix is based on comparing populations; if FALSE, on individuals [default TRUE].	
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].	

Details

The script takes a genlight object as input, creates a set of sequences from the trimmed sequence tags for each individual, calculates distances between the individuals and then optionally averages those distances between the populations defined in the genlight object (typically OTUs).

min.tag.length : Sequence tags can vary considerably in length, which results in large numbers of Ns in alignments. This can have an impact of distance measures depending on how missing values are managed. To minimize this effect, you might elect to filter on tag length using this parameter.

subst.model : Use this parameter to specify the substitution model, selecting from the list used by package ape.

- raw: This is simply the proportion or the number of sites that differ between each pair of sequences. This may be useful to draw "saturation plots". The options variance and gamma have no effect, but pairwise.deletion can.
- TS, TV: These are the numbers of transitions and transversions, respectively.
- JC69: This model was developed by Jukes and Cantor (1969). It assumes that all substitutions (i.e. a change of a base by another one) have the same probability. This probability is the same for all sites along the DNA sequence. This last assumption can be relaxed by assuming that the substition rate varies among site following a gamma distribution which parameter must be given by the user. By default, no gamma correction is applied. Another assumption is that the base frequencies are balanced and thus equal to 0.25.
- K80: The distance derived by Kimura (1980), sometimes referred to as "Kimura's 2-parameters distance", has the same underlying assumptions than the Jukes–Cantor distance except that two kinds of substitutions are considered: transitions (A <-> G, C <-> T), and transversions (A <-> C, A <-> T, C <-> G, G <-> T). They are assumed to have different probabilities. A transition is the substitution of a purine (C, T) by another one, or the substitution of a pyrimidine (A, G) by another one. A transversion is the substitution of a purine by a pyrimidine, or vice-versa. Both transition and transversion rates are the same for all sites along the DNA sequence. Jin and Nei (1990) modified the Kimura model to allow for variation among sites following a gamma distribution. Like for the Jukes–Cantor model, the gamma parameter must be given by the user. By default, no gamma correction is applied.
- F81: Felsenstein (1981) generalized the Jukes–Cantor model by relaxing the assumption of equal base frequencies. The formulae used in this function were taken from McGuire et al. (1999).

- K81: Kimura (1981) generalized his model (Kimura 1980) by assuming different rates for two kinds of transversions: A <-> C and G <-> T on one side, and A <-> T and C <-> G on the other. This is what Kimura called his "three substitution types model" (3ST), and is sometimes referred to as "Kimura's 3-parameters distance".
- F84: This model generalizes K80 by relaxing the assumption of equal base frequencies. It was first introduced by Felsenstein in 1984 in Phylip, and is fully described by Felsenstein and Churchill (1996). The formulae used in this function were taken from McGuire et al. (1999).
- BH87: Barry and Hartigan (1987) developed a distance based on the observed proportions of changes among the four bases. This distance is not symmetric.
- T92: Tamura (1992) generalized the Kimura model by relaxing the assumption of equal base frequencies. This is done by taking into account the bias in G+C content in the sequences. The substitution rates are assumed to be the same for all sites along the DNA sequence.
- TN93: Tamura and Nei (1993) developed a model which assumes distinct rates for both kinds of transition (A <-> G versus C <-> T), and transversions. The base frequencies are not assumed to be equal and are estimated from the data. A gamma correction of the inter-site variation in substitution rates is possible.
- GG95: Galtier and Gouy (1995) introduced a model where the G+C content may change through time. Different rates are assumed for transitons and transversions.
- logdet: The Log-Det distance, developed by Lockhart et al. (1994), is related to BH87. However, this distance is symmetric. Formulae from Gu and Li (1996) are used. dist.logdet in phangorn uses a different implementation that gives substantially different distances for lowdiverging sequences.
- paralin: Lake (1994) developed the paralinear distance which can be viewed as another variant of the Barry–Hartigan distance.
- pairwise.missing : If TRUE, then missing values in the sequence (NNNs) will be accommodated in the calculations pair of taxa at a time; otherwise, the deletion of data at positions in the sequence will be global (deleted if any missing data at the position in any individual).

Value

The distance matrix as an object of class dist.

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

```
Other phylogeny: gl.tree.fitch()
```

Examples

```
## Not run:
tmp <- gl.filter.monomorphs(testset.gl)
gl.dist.phylo(xx=tmp,subst.model="F80")
```

```
## End(Not run)
```

gl.dist.pop

Calculates a distance matrix for populations with SNP or Silicodart genotypes in a genlight object

Description

This script calculates various distances between populations based on allele frequencies (SNP genotypes) or frequency of presences in PA (SilicoDArT) data

Usage

```
gl.dist.pop(
    x,
    as.pop = NULL,
    method = "euclidean",
    scale = FALSE,
    type = "dist",
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
    plot.file = NULL,
    plot.dir = NULL,
    verbose = NULL
)
```

Arguments

х	Name of the genlight object [required].
as.pop	Temporarily assign another locus metric as the population for the purposes of deletions [default NULL].
method	Specify distance measure [default euclidean].
scale	If TRUE and method='Euclidean', the distance will be scaled to fall in the range [0,1] [default FALSE].
type	Specify the type of output, dist or matrix [default 'dist']
plot.display	If TRUE, resultant plots are displayed in the plot window [default TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.colors	List of two color names for the borders and fill of the plots [default c("#2171B5", "#6BAED6")].
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL]
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

gl.drop.ind

Details

For SNP data, the distance measure can be one of 'euclidean', 'fixed-diff', 'reynolds', 'nei' and 'chord'. For SilicoDArT data, the distance measure can be one of 'Refer to the documentation of functions in https://doi.org/10.1101/2023.03.22.533737 for algorithms and definitions.

Value

An object of class 'dist' giving distances between populations

Author(s)

author(s): Arthur Georges. Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

Other distance: gl.dist.ind(), gl.fdsim(), utils.dist.ind.snp()

Examples

```
# SNP genotypes
D <- gl.dist.pop(possums.gl, method='euclidean')
D <- gl.dist.pop(possums.gl, method='euclidean',scale=TRUE)
D <- gl.dist.pop(possums.gl, method='nei')
D <- gl.dist.pop(possums.gl, method='reynolds')
D <- gl.dist.pop(possums.gl, method='chord')
D <- gl.dist.pop(possums.gl, method='fixed-diff')
#Presence-Absence data [only 10 individuals due to speed]
D <- gl.dist.pop(testset.gs[1:10,], method='euclidean')</pre>
```

gl.drop.ind

Removes specified individuals from a dartR genlight object

Description

This function deletes individuals and their associated metadata. Monomorphic loci and loci that are scored all NA are optionally deleted (mono.rm=TRUE). The script also optionally recalculates locus metadata statistics to accommodate the deletion of individuals from the dataset (recalc=TRUE).

The script returns a dartR genlight object with the retained individuals and the recalculated locus metadata. The script works with both genlight objects containing SNP genotypes and Tag P/A data (SilicoDArT).

Usage

```
gl.drop.ind(x, ind.list, recalc = FALSE, mono.rm = FALSE, verbose = NULL)
```

Arguments

х	Name of the genlight object [required].
ind.list	List of individuals to be removed [required].
recalc	If TRUE, recalculate the locus metadata statistics [default FALSE].
mono.rm	If TRUE, remove monomorphic and all NA loci [default FALSE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress but not results; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

A reduced dartR genlight object

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

gl.keep.ind to keep rather than drop specified individuals

```
Other data manipulation: gl.define.pop(), gl.drop.loc(), gl.drop.pop(), gl.edit.recode.pop(),
gl.impute(), gl.join(), gl.keep.ind(), gl.keep.loc(), gl.keep.pop(), gl.make.recode.ind(),
gl.merge.pop(), gl.reassign.pop(), gl.recode.ind(), gl.recode.pop(), gl.rename.pop(),
gl.sample(), gl.sim.genotypes(), gl.sort(), gl.subsample.ind(), gl.subsample.loc()
```

Examples

```
# SNP data
gl2 <- gl.drop.ind(testset.gl,
ind.list=c('AA019073','AA004859'))
# Tag P/A data
gs2 <- gl.drop.ind(testset.gs,
ind.list=c('AA020656','AA19077','AA004859'))
gs2 <- gl.drop.ind(testset.gs, ind.list=c('AA020656', 'AA19077','AA004859'),mono.rm=TRUE, recalc=TRUE)</pre>
```

gl.drop.loc

Removes specified loci from a dartR genlight object

Description

This function deletes individuals and their associated metadata. The script returns a dartR genlight object with the retained loci. The script works with both genlight objects containing SNP genotypes and Tag P/A data (SilicoDArT).

gl.drop.pop

Usage

gl.drop.loc(x, loc.list = NULL, first = NULL, last = NULL, verbose = NULL)

Arguments

x	Name of the genlight object [required].
loc.list	A list of loci to be deleted [required, if loc.range not specified].
first	First of a range of loci to be deleted [required, if loc.list not specified].
last	Last of a range of loci to be deleted [if not specified, last locus in the dataset].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress but not results; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

A reduced dartR genlight object

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

gl.keep.loc to keep rather than drop specified loci

```
Other data manipulation: gl.define.pop(), gl.drop.ind(), gl.drop.pop(), gl.edit.recode.pop(),
gl.impute(), gl.join(), gl.keep.ind(), gl.keep.loc(), gl.keep.pop(), gl.make.recode.ind(),
gl.merge.pop(), gl.reassign.pop(), gl.recode.ind(), gl.recode.pop(), gl.rename.pop(),
gl.sample(), gl.sim.genotypes(), gl.sort(), gl.subsample.ind(), gl.subsample.loc()
```

Examples

```
# SNP data
gl2 <- gl.drop.loc(testset.gl, loc.list=c('100051468|42-A/T', '100049816-51-A/G'),verbose=3)
# Tag P/A data
gs2 <- gl.drop.loc(testset.gs, loc.list=c('20134188','19249144'),verbose=3)</pre>
```

gl.drop.pop

Removes specified populations from a dartR genlight object

Description

Individuals are assigned to populations based on associated specimen metadata stored in the dartR genlight object. This function deletes all individuals in the nominated populations (pop.list). Monomorphic loci and loci that are scored all NA are optionally deleted (mono.rm=TRUE). The script also optionally recalculates locus metadata statistics to accommodate the deletion of individuals from the dataset (recalc=TRUE). The script returns a dartR genlight object with the retained populations and the recalculated locus metadata. The script works with both genlight objects containing SNP genotypes and Tag P/A data (SilicoDArT).

Usage

```
gl.drop.pop(
    x,
    pop.list,
    as.pop = NULL,
    recalc = FALSE,
    mono.rm = FALSE,
    verbose = NULL
)
```

Arguments

x	Name of the genlight object [required].
pop.list	List of populations to be removed [required].
as.pop	Temporarily assign another locus metric as the population for the purposes of deletions [default NULL].
recalc	If TRUE, recalculate the locus metadata statistics [default FALSE].
mono.rm	If TRUE, remove monomorphic and all NA loci [default FALSE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress but not results; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

A reduced dartR genlight object

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

gl.keep.pop to keep rather than drop specified populations

Other data manipulation: gl.define.pop(), gl.drop.ind(), gl.drop.loc(), gl.edit.recode.pop(), gl.impute(), gl.join(), gl.keep.ind(), gl.keep.loc(), gl.keep.pop(), gl.make.recode.ind(), gl.merge.pop(), gl.reassign.pop(), gl.recode.ind(), gl.recode.pop(), gl.rename.pop(), gl.sample(), gl.sim.genotypes(), gl.sort(), gl.subsample.ind(), gl.subsample.loc()

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gl.edit.recode.ind

Examples

```
# SNP data
 gl2 <- gl.drop.pop(testset.gl,</pre>
 pop.list=c('EmsubRopeMata', 'EmvicVictJasp'), verbose=3)
 gl2 <- gl.drop.pop(testset.gl, pop.list=c('EmsubRopeMata','EmvicVictJasp'),</pre>
 mono.rm=TRUE,recalc=TRUE)
  gl2 <- gl.drop.pop(testset.gl,as.pop='sex',pop.list=c('Male','Unknown'),verbose=3)</pre>
# Tag P/A data
  gs2 <- gl.drop.pop(testset.gs, pop.list=c('EmsubRopeMata', 'EmvicVictJasp'))</pre>
```

gl.edit.recode.ind	Creates or edits individual (=specimen) names, creates a recode_ind
	file and applies the changes to a genlight object @family data manip-
	ulation

Description

A function to edit names of individual in a dartR genlight object, or to create a reassignment table taking the individual labels from a genlight object, or to edit existing individual labels in an existing recode_ind file. The amended recode table is then applied to the genlight object.

Usage

```
gl.edit.recode.ind(
  х,
  out.recode.file = NULL,
  outpath = NULL,
  recalc = FALSE,
 mono.rm = FALSE,
  verbose = NULL
)
```

Arguments

x	Name of the genlight object [required].
out.recode.file	
	Name of the file to output the new individual labels [optional].
outpath	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]
recalc	If TRUE, recalculate the locus metadata statistics [default TRUE].
mono.rm	If TRUE, remove monomorphic loci [default TRUE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress but not results; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Details

Renaming individuals may be required when there have been errors in labeling arising in the passage of samples to sequencing. There may be occasions where renaming individuals is required for preparation of figures. This function will input an existing recode table for editing and optionally save it as a new table, or if the name of an input table is not supplied, will generate a table using the individual labels in the parent genlight object. When caution needs to be exercised because of the potential for breaking the 'chain of evidence' associated with the samples, recoding individuals using a recode table (csv) can provide a durable record of the changes. For SNP genotype data, the function, having deleted individuals, optionally identifies resultant monomorphic loci or loci with all values missing and deletes them. The script also optionally recalculates the locus metadata as appropriate. The optional deletion of monomorphic loci and the optional recalculation of locus statistics is not available for Tag P/A data (SilicoDArT). Use outpath=getwd() when calling this function to direct output files to your working directory. The function returns a dartR genlight object with the new population assignments and the recalculated locus metadata.

Value

An object of class ('genlight') with the revised individual labels.

Author(s)

Custodian: Arthur Georges - Post to https://groups.google.com/d/forum/dartr

See Also

gl.recode.ind, gl.drop.ind, gl.keep.ind

Examples

```
#this is an interactive example
if(interactive()){
gl <- gl.edit.recode.ind(testset.gl)
gl <- gl.edit.recode.ind(testset.gl, out.recode.file='ind.recode.table.csv')
}</pre>
```

gl.edit.recode.pop Creates or edits and applies a population re-assignment table

Description

A function to edit population assignments in a dartR genlight object, or to create a reassignment table taking the population assignments from a genlight object, or to edit existing population assignments in a pop.recode.table. The amended recode table is then applied to the genlight object.

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gl.edit.recode.pop

Usage

```
gl.edit.recode.pop(
    x,
    pop.recode = NULL,
    out.recode.file = NULL,
    outpath = NULL,
    recalc = FALSE,
    mono.rm = FALSE,
    verbose = NULL
)
```

Arguments

x	Name of the genlight object [required].
pop.recode	Path to recode file [default NULL].
out.recode.fil	e
	Name of the file to output the new individual labels [default NULL].
outpath	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]
recalc	If TRUE, recalculate the locus metadata statistics [default TRUE].
mono.rm	If TRUE, remove monomorphic loci [default TRUE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress but not results; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].
	@details Genlight objects assign specimens to populations based on information in the ind.metadata file provided when the genlight object is first generated. Of- ten one wishes to subset the data by deleting populations or to amalgamate pop- ulations. This can be done with a pop.recode table with two columns. The first column is the population assignment in the genlight object, the second column provides the new assignment. This function will input an existing reassignment table for editing and optionally save it as a new table, or if the name of an input table is not supplied, will generate a table using the population assignments in the parent genlight object. It will then apply the recodings to the genlight ob- ject. When caution needs to be exercised because of the potential for breaking the 'chain of evidence' associated with the samples, recoding individuals us- ing a recode table (csv) can provide a durable record of the changes. For SNP genotype data, the function, having deleted populations, optionally identifies resultant monomorphic loci or loci with all values missing and deletes them. The script also optionally recalculates the locus metadata as appropriate. The optional deletion of monomorphic loci and the optional recalculation of locus statistics is not available for Tag P/A data (SilicoDArT). Use outpath=getwd() when calling this function to direct output files to your working directory. The function returns a dartR genlight object with the new population assignments and the recalculated locus metadata.

Value

A genlight object with the revised population assignments

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

gl.recode.pop,gl.drop.pop,gl.keep.pop,gl.merge.pop,gl.reassign.pop

```
Other data manipulation: gl.define.pop(), gl.drop.ind(), gl.drop.loc(), gl.drop.pop(),
gl.impute(), gl.join(), gl.keep.ind(), gl.keep.loc(), gl.keep.pop(), gl.make.recode.ind(),
gl.merge.pop(), gl.reassign.pop(), gl.recode.ind(), gl.recode.pop(), gl.rename.pop(),
gl.sample(), gl.sim.genotypes(), gl.sort(), gl.subsample.ind(), gl.subsample.loc()
```

Examples

```
#this is an interactive example
if(interactive()){
gl <- gl.edit.recode.pop(testset.gl)
gs <- gl.edit.recode.pop(testset.gs)
}
# See also -----</pre>
```

gl.fdsim

Estimates the rate of false positives in a fixed difference analysis

Description

This function takes two populations and generates allele frequency profiles for them. It then samples an allele frequency for each, at random, and estimates a sampling distribution for those two allele frequencies. Drawing two samples from those sampling distributions, it calculates whether or not they represent a fixed difference. This is applied to all loci, and the number of fixed differences so generated are counted, as an expectation. The script distinguished between true fixed differences (with a tolerance of delta), and false positives. The simulation is repeated a given number of times (default=1000) to provide an expectation of the number of false positives, given the observed allele frequency profiles and the sample sizes. The probability of the observed count of fixed differences is greater than the expected number of false positives is calculated.

Usage

```
gl.fdsim(
    x,
    poppair,
    obs = NULL,
    sympatric = FALSE,
```

gl.fdsim

```
reps = 1000,
delta = 0.02,
verbose = NULL
)
```

Arguments

x	Name of the genlight containing the SNP genotypes [required].
poppair	Labels of two populations for comparison in the form c(popA,popB) [required].
obs	Observed number of fixed differences between the two populations [default NULL].
sympatric	If TRUE, the two populations are sympatric, if FALSE then allopatric [default FALSE].
reps	Number of replications to undertake in the simulation [default 1000].
delta	The threshold value for the minor allele frequency to regard the difference be- tween two populations to be fixed [default 0.02].
verbose	Verbosity: 0, silent, fatal errors only; 1, flag function begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

A list containing the following square matrices [[1]] observed fixed differences; [[2]] mean expected number of false positives for each comparison; [[3]] standard deviation of the no. of false positives for each comparison; [[4]] probability the observed fixed differences arose by chance for each comparison.

Author(s)

Custodian: Arthur Georges (Post to https://groups.google.com/d/forum/dartr)

See Also

Other distance: gl.dist.ind(), gl.dist.pop(), utils.dist.ind.snp()

Examples

```
fd <- gl.fdsim(testset.gl[,1:100],poppair=c('EmsubRopeMata','EmmacBurnBara'),
sympatric=TRUE,verbose=3)</pre>
```

gl.filter.allna

Description

This script deletes loci or individuals with all calls missing (NA), from a genlight object.

A DArT dataset will not have loci for which the calls are scored all as missing (NA) for a particular individual, but such loci can arise rarely when populations or individuals are deleted. Similarly, a DArT dataset will not have individuals for which the calls are scored all as missing (NA) across all loci, but such individuals may sneak in to the dataset when loci are deleted. Retaining individual or loci with all NAs can cause issues for several functions.

Also, on occasions an analysis will require that there are some loci scored in each population. Setting by.pop=TRUE will result in removal of loci when they are all missing in any one population.

Note that loci that are missing for all individuals in a population are not imputed with method 'frequency' or 'HW'. Consider using the function gl.filter.allna with by.pop=TRUE.

Usage

gl.filter.allna(x, by.pop = FALSE, recalc = FALSE, verbose = NULL)

Arguments

х	Name of the input genlight object [required].
by.pop	If TRUE, loci that are all missing in any one population are deleted [default FALSE]
recalc	Recalculate the locus metadata statistics if any individuals are deleted in the filtering [default FALSE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Value

A genlight object having removed individuals that are scored NA across all loci, or loci that are scored NA across all individuals.

Author(s)

Author(s): Arthur Georges. Custodian: Arthur Georges – Post to https://groups.google.com/ d/forum/dartr

See Also

Other filter functions: gl.filter.hwe(), gl.report.allna()

gl.filter.callrate

Examples

```
# SNP data
result <- gl.filter.allna(testset.gl, verbose=3)
# Tag P/A data
result <- gl.filter.allna(testset.gs, verbose=3)</pre>
```

gl.filter.callrate Filters loci or specimens in a genlight {adegenet} object based on call rate

Description

SNP datasets generated by DArT have missing values primarily arising from failure to call a SNP because of a mutation at one or both of the restriction enzyme recognition sites. The script gl.filter.callrate() will filter out the loci with call rates below a specified threshold. Tag Presence/Absence datasets (SilicoDArT) have missing values where it is not possible to determine reliably if there the sequence tag can be called at a particular locus.

Usage

```
gl.filter.callrate(
    x,
    method = "loc",
    threshold = 0.95,
    mono.rm = FALSE,
    recalc = FALSE,
    recursive = FALSE,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
    plot.file = NULL,
    plot.dir = NULL,
    bins = 25,
    verbose = NULL
)
```

Arguments

x	Name of the genlight object containing the SNP data, or the genind object con- taining the SilocoDArT data [required].
method	Use method='loc' to specify that loci are to be filtered, 'ind' to specify that specimens are to be filtered, 'pop' to remove loci that fail to meet the specified threshold in any one population [default 'loc'].
threshold	Threshold value below which loci will be removed [default 0.95].
mono.rm	Remove monomorphic loci after analysis is complete [default FALSE].

recalc	Recalculate the locus metadata statistics if any individuals are deleted in the filtering [default FALSE].
recursive	Repeatedly filter individuals on call rate, each time removing monomorphic loci. Only applies if method='ind' and mono.rm=TRUE [default FALSE].
plot.display	If TRUE, histograms are displayed in the plot window [default TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.colors	Vector with two color names for the borders and fill [default c("#2171B5", "#6BAED6")].
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL]
plot.dir	Directory in which to save files [default = working directory]
bins	Number of bins to display in histograms [default 25].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Details

Because this filter operates on call rate, this function recalculates Call Rate, if necessary, before filtering. If individuals are removed using method='ind', then the call rate stored in the genlight object is, optionally, recalculated after filtering. Note that when filtering individuals on call rate, the initial call rate is calculated and compared against the threshold. After filtering, if mono.rm=TRUE, the removal of monomorphic loci will alter the call rates. Some individuals with a call rate initially greater than the nominated threshold, and so retained, may come to have a call rate lower than the threshold. If this is a problem, repeated iterations of this function will resolve the issue. This is done by setting mono.rm=TRUE and recursive=TRUE, or it can be done manually. Callrate is summarized by locus or by individual to allow sensible decisions on thresholds for filtering taking into consideration consequential loss of data. The summary is in the form of a tabulation and plots. Plot themes can be obtained from

- https://ggplot2.tidyverse.org/reference/ggtheme.html and
- https://yutannihilation.github.io/allYourFigureAreBelongToUs/ggthemes/

Resultant ggplot(s) and the tabulation(s) are saved to the session's temporary directory.

Value

The reduced genlight or genind object, plus a summary

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

gl.report.callrate

```
Other matched filter: gl.filter.hamming(), gl.filter.ld(), gl.filter.locmetric(), gl.filter.maf(),
gl.filter.monomorphs(), gl.filter.overshoot(), gl.filter.pa(), gl.filter.secondaries()
```

gl.filter.excess.het

Examples

```
# SNP data
result <- gl.filter.callrate(testset.gl[1:10], method='loc', threshold=0.8,
verbose=3)
result <- gl.filter.callrate(testset.gl[1:10], method='ind', threshold=0.8,
verbose=3)
# Tag P/A data
result <- gl.filter.callrate(testset.gl[1:10], method='pop', threshold=0.8,
verbose=3)
# Tag P/A data
result <- gl.filter.callrate(testset.gs[1:10], method='loc',
threshold=0.95, verbose=3)
result <- gl.filter.callrate(testset.gs[1:10], method='ind',
threshold=0.8, verbose=3)
result <- gl.filter.callrate(testset.gs[1:10], method='ind',
threshold=0.8, verbose=3)
result <- gl.filter.callrate(testset.gs[1:10], method='pop',
threshold=0.8, verbose=3)
result <- gl.filter.callrate(testset.gs[1:10], method='pop',
threshold=0.8, verbose=3)</pre>
```

gl.filter.excess.het Filters excessively-heterozygous loci from a genlight object

Description

Calculates excess of heterozygosity in a genlight object and remove those loci

Usage

```
gl.filter.excess.het(
    x,
    Yates = FALSE,
    mono.rm = FALSE,
    recalc = FALSE,
    verbose = NULL
)
```

Arguments

х	A genlight object containing the SNP genotypes [required].
Yates	Whether to use Yates's continuity correction [default FALSE].
mono.rm	Remove monomorphic loci after analysis is complete [default FALSE].
recalc	Recalculate the locus metadata statistics if any individuals are deleted in the filtering [default FALSE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Value

Returns unaltered genlight object

Author(s)

Author(s): Jesús Castrejón-Figueroa, Diana A Robledo-Ruiz (Custodian: Ching Ching Lau) – Post to https://groups.google.com/d/forum/dartr

References

- · https://github.com/drobledoruiz/conservation_genomics/tree/main/filter.excess.het
- Robledo-Ruiz, D. A., Austin, L., Amos, J. N., Castrejón-Figueroa, J., Harley, D. K., Magrath, M. J., Sunnucks, P. & Pavlova, A. (2023). Easy-to-use R functions to separate reducedrepresentation genomic datasets into sex-linked and autosomal loci, and conduct sex assignment. Molecular Ecology Resources.

See Also

gl.filter.callrate

```
Other matched report: gl.report.allna(), gl.report.callrate(), gl.report.hamming(),
gl.report.locmetric(), gl.report.maf(), gl.report.overshoot(), gl.report.pa(), gl.report.rdepth(),
gl.report.reproducibility(), gl.report.secondaries(), gl.report.taglength()
```

Examples

```
filtered.gl <- gl.filter.excess.het(x = LBP, Yates = TRUE)
# Use below function to output information of the loci with Yates's continuity correction specified
filtered.table <- gl.report.excess.het(x = LBP, Yates = TRUE)</pre>
```

gl.filter.factorloadings

Filters loci based on factor loadings for a PCA or PCoA

Description

Extracts the factor loadings from a glPCA object (generated by gl.pcoa) and filters loci based on a user specified threshold for the ABSOLUTE value of the factor loadings.

Usage

```
gl.filter.factorloadings(
    x,
    pca,
    axis = 1,
    threshold,
    retain = FALSE,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
    plot.file = NULL,
    plot.dir = NULL,
```
```
bins = 25,
verbose = NULL,
...
```

Arguments

)

X	Name of the genlight object containing the SNP data or the SilocoDArT data [required].
рса	Name of the glPCA object containing factor loadings [required].
axis	Axis in the ordination used to display the factor loadings [default 1]
threshold	Numeric value for the factor loadings. This value is the ABSOLUTE value of the factor loadings [required].
retain	If true, the resultant genlight object holds only the loci that load high on the specified axis; if FALSE, the resultant genlight object has the loci loading high on the specified axis filtered out [default FALSE].
plot.display	If TRUE, resultant plots are displayed in the plot window [default TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.colors	List of two color names for the borders and fill of the plots [default c("#2171B5", "#6BAED6")].
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL]
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]
bins	Number of bins to display in histograms [default 25].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity]
	Parameters passed to function ggsave, such as width and height, when the ggplot is to be saved.

Details

The function extracts the factor loadings for a given axis from a PCA object generated by gl.pcoa and then filters loci on the basis of a user specified threshold. The threshold value is decided using gl.report.factorloadings. The function can be used to filter out loci that load high with a particular axis or alternatively if retain=TRUE, to retain loci that load high on a specified axis.

Note that this function also removes monomorphic loci because PCA is performed only on polymorphic loci.

A color vector can be obtained with gl.select.colors() and then passed to the function with the plot.colors parameter.

Themes can be obtained from in

- https://ggplot2.tidyverse.org/reference/ggtheme.html and
- https://yutannihilation.github.io/allYourFigureAreBelongToUs/ggthemes/

If a plot.file is given, the ggplot arising from this function is saved as an "RDS" binary file using saveRDS(); can be reloaded with readRDS(). A file name must be specified for the plot to be saved. If a plot directory (plot.dir) is specified, the ggplot binary is saved to that directory; otherwise to the tempdir().

Value

The unchanged genlight object

Author(s)

Custodian: Arthur Georges - Post to https://groups.google.com/d/forum/dartr

Examples

```
pca <- gl.pcoa(testset.gl)
gl.report.factorloadings(pca = pca)
gl2 <- gl.filter.factorloadings(pca=pca,x=testset.gl,threshold=0.2)</pre>
```

gl.filter.hamming Filters loci based on pairwise Hamming distance between sequence tags

Description

Hamming distance is calculated as the number of base differences between two sequences which can be expressed as a count or a proportion. Typically, it is calculated between two sequences of equal length. In the context of DArT trimmed sequences, which differ in length but which are anchored to the left by the restriction enzyme recognition sequence, it is sensible to compare the two trimmed sequences starting from immediately after the common recognition sequence and terminating at the last base of the shorter sequence.

Usage

```
gl.filter.hamming(
    x,
    threshold = 0.2,
    rs = 5,
    tag.length = 69,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
    plot.file = NULL,
    plot.dir = NULL,
    pb = FALSE,
    verbose = NULL
)
```

gl.filter.hamming

Arguments

x	Name of the genlight object containing the SNP data [required].
threshold	A threshold Hamming distance for filtering loci [default threshold 0.2].
rs	Number of bases in the restriction enzyme recognition sequence [default 5].
tag.length	Typical length of the sequence tags [default 69].
plot.display	If TRUE, histograms are displayed in the plot window [default TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.colors	List of two color names for the borders and fill of the plots [default c("#2171B5", "#6BAED6")].
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL]
plot.dir	Directory in which to save files [default = working directory]
pb	If TRUE, a progress bar will be displayed [default FALSE]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Details

Hamming distance can be computed by exploiting the fact that the dot product of two binary vectors x and (1-y) counts the corresponding elements that are different between x and y. This approach can also be used for vectors that contain more than two possible values at each position (e.g. A, C, T or G). If a pair of DNA sequences are of differing length, the longer is truncated. The algorithm is that of Johann de Jong https://johanndejong.wordpress.com/2015/10/02/faster-hamming-distance-in-r-2/ as implemented in utils.hamming. Only one of two loci are retained if their Hamming distance is less that a specified percentage. 5 base differences out of 100 bases is a 20

Value

A genlight object filtered on Hamming distance.

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

```
Other matched filter: gl.filter.callrate(), gl.filter.ld(), gl.filter.locmetric(), gl.filter.maf(),
gl.filter.monomorphs(), gl.filter.overshoot(), gl.filter.pa(), gl.filter.secondaries()
```

Examples

```
# SNP data
test <- gl.subsample.loc(platypus.gl,n=50)
result <- gl.filter.hamming(test, threshold=0.6, verbose=3)</pre>
```

```
gl.filter.heterozygosity
```

Filters individuals with average heterozygosity greater than a specified upper threshold or less than a specified lower threshold @family matched filter

Description

Calculates the observed heterozygosity for each individual in a genlight object and filters individuals based on specified threshold values. Use gl.report.heterozygosity to determine the appropriate thresholds.

Usage

```
gl.filter.heterozygosity(x, t.upper = 0.7, t.lower = 0, verbose = NULL)
```

Arguments

х	A genlight object containing the SNP genotypes [required].
t.upper	Filter individuals > the threshold [default 0.7].
t.lower	Filter individuals < the threshold [default 0].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Value

The filtered genlight object.

Author(s)

Custodian: Luis Mijangos - Post to https://groups.google.com/d/forum/dartr

Examples

```
result <- gl.filter.heterozygosity(testset.gl,t.upper=0.06,verbose=3)
tmp <- gl.report.heterozygosity(result,method='ind')</pre>
```

gl.filter.hwe

Filters loci that show significant departure from Hardy-Weinberg Equilibrium @family matched filter

Description

This function filters out loci showing significant departure from H-W proportions based on observed frequencies of reference homozygotes, heterozygotes and alternate homozygotes. Loci are filtered out if they show HWE departure either in any one population (n.pop.threshold =1) or in at least X number of populations (n.pop.threshold > 1).

Usage

```
gl.filter.hwe(
    x,
    subset = "each",
    n.pop.threshold = 1,
    test.type = "Exact",
    mult.comp.adj = FALSE,
    mult.comp.adj.method = "BY",
    alpha = 0.05,
    pvalue.type = "midp",
    cc.val = 0.5,
    n.min = 5,
    verbose = NULL
)
```

Arguments

х	Name of the genlight object containing the SNP data [required].
subset	Whether to perform H-W tests within each population ("each"), or taking all individuals as one population ("all") (see details) [default 'each'].
n.pop.threshold	
	The minimum number of populations where the same locus has to be out of H-W proportions to be removed [default 1].
test.type	Method for determining statistical significance: 'ChiSquare' or 'Exact' [default 'Exact'].
mult.comp.adj Whether to adjust p-values for multiple comparisons [default FALSE]. mult.comp.adj.method	
	Method to adjust p-values for multiple comparisons: 'holm', 'hochberg', 'hom- mel', 'bonferroni', 'BH', 'BY', 'fdr' (see details) [default 'fdr'].
alpha	Level of significance for testing [default 0.05].
pvalue.type	Type of p-value to be used in the Exact method. Either 'dost', 'selome', 'midp' (see details) [default 'midp'].
cc.val	The continuity correction applied to the ChiSquare test [default 0.5].

n.min	Minimum number of individuals per population in which perform H-W tests [default 5].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Details

Several factors can cause deviations from Hardy-Weinberg equilibrium including: mutation, finite population size, selection, population structure, age structure, assortative mating, sex linkage, non-random sampling and genotyping errors. Refer to Waples (2015). Note that tests for departure from H-W equilibrium are only valid if there is no population substructure (assuming random mating) and have sufficient power only when there is sufficient sample size (n individuals > 15). Populations can be defined in three ways:

- Merging all populations in the dataset using subset = 'all'.
- Within each population separately using: subset = 'each'.
- Within selected populations using for example: subset = c('pop1','pop2').

Two different statistical methods to test for deviations from Hardy Weinberg proportions:

- The classical chi-square test (test.type='ChiSquare') based on the function HWChisq of the R package HardyWeinberg. By default a continuity correction is applied (cc.val=0.5). The continuity correction can be turned off (by specifying cc.val=0), for example when extreme allele frequencies occur continuity correction can lead to excessive Type I error rates.
- The exact test (test.type='Exact') based on the exact calculations contained in the function HWExactStats of the R package HardyWeinberg as described by Wigginton et al. (2005). The exact test is recommended in most cases. Three different methods to estimate p-values (pvalue.type) in the Exact test can be used:
 - 'dost' p-value is computed as twice the tail area of a one-sided test.
 - 'selome' p-value is computed as the sum of the probabilities of all samples less or equally likely as the current sample.
 - 'midp', p-value is computed as half the probability of the current sample + the probabilities of all samples that are more extreme.

The standard exact p-value is overly conservative, in particular for small minor allele frequencies. The mid p-value ameliorates this problem by bringing the rejection rate closer to the nominal level, at the price of occasionally exceeding the nominal level (Graffelman & Moreno, 2013).

Correction for multiple tests can be applied using the following methods based on the function p.adjust:

- 'holm' is also known as the sequential Bonferroni technique (Rice, 1989). This method has a greater statistical power than the standard Bonferroni test, however this method becomes very stringent when many tests are performed and many real deviations from the null hypothesis can go undetected (Waples, 2015).
- 'hochberg' based on Hochberg, 1988.
- 'hommel' based on Hommel, 1988. This method is more powerful than Hochberg's, but the difference is usually small.

gl.filter.hwe

- 'bonferroni' in which p-values are multiplied by the number of tests. This method is very stringent and therefore has reduced power to detect multiple departures from the null hypothesis.
- 'BH' based on Benjamini & Hochberg, 1995.
- 'BY' based on Benjamini & Yekutieli, 2001.

The first four methods are designed to give strong control of the family-wise error rate. The last two methods control the false discovery rate (FDR), the expected proportion of false discoveries among the rejected hypotheses. The false discovery rate is a less stringent condition than the family-wise error rate, so these methods are more powerful than the others, especially when number of tests is large. The number of tests on which the adjustment for multiple comparisons is the number of populations times the number of loci. From v2.1 gl.filter.hwe takes the argument n.pop.threshold. if n.pop.threshold > 1 loci will be removed only if they are concurrently significant (after adjustment if applied) out of hwe in >= n.pop.threshold > 1.

Value

A genlight object with the loci departing significantly from H-W proportions removed.

Author(s)

Custodian: Luis Mijangos – Post to https://groups.google.com/d/forum/dartr

References

- Benjamini, Y., and Yekutieli, D. (2001). The control of the false discovery rate in multiple testing under dependency. Annals of Statistics, 29, 1165–1188.
- Graffelman, J. (2015). Exploring Diallelic Genetic Markers: The Hardy Weinberg Package. Journal of Statistical Software 64:1-23.
- Graffelman, J. & Morales-Camarena, J. (2008). Graphical tests for Hardy-Weinberg equilibrium based on the ternary plot. Human Heredity 65:77-84.
- Graffelman, J., & Moreno, V. (2013). The mid p-value in exact tests for Hardy-Weinberg equilibrium. Statistical applications in genetics and molecular biology, 12(4), 433-448.
- Hochberg, Y. (1988). A sharper Bonferroni procedure for multiple tests of significance. Biometrika, 75, 800–803.
- Hommel, G. (1988). A stagewise rejective multiple test procedure based on a modified Bonferroni test. Biometrika, 75, 383–386.
- Rice, W. R. (1989). Analyzing tables of statistical tests. Evolution, 43(1), 223-225.
- Waples, R. S. (2015). Testing for Hardy–Weinberg proportions: have we lost the plot?. Journal of heredity, 106(1), 1-19.
- Wigginton, J.E., Cutler, D.J., & Abecasis, G.R. (2005). A Note on Exact Tests of Hardy-Weinberg Equilibrium. American Journal of Human Genetics 76:887-893.

See Also

gl.report.hwe

Other filter functions: gl.filter.allna(), gl.report.allna()

Examples

result <- gl.filter.hwe(x = bandicoot.gl)</pre>

gl.filter.ld Filters loci based on linkage disequilibrium (LD)

Description

This function uses the statistic set in the parameter stat.keep from function gl.report.ld.map to choose the SNP to keep when two SNPs are in LD. When a SNP is selected to be filtered out in each pairwise comparison, the function stores its name in a list. In subsequent pairwise comparisons, if the SNP is already in the list, the other SNP will be kept.

Usage

```
gl.filter.ld(
    x,
    ld.report,
    threshold = 0.2,
    pop.limit = ceiling(nPop(x)/2),
    verbose = NULL
)
```

Arguments

х	Name of the genlight object containing the SNP data [required].
ld.report	Output from function gl.report.ld.map [required].
threshold	Threshold value above which loci will be removed [default 0.2].
pop.limit	Minimum number of populations in which LD should be more than the threshold for a locus to be filtered out. The default value is half of the populations [default ceiling($nPop(x)/2$)].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Value

The reduced genlight object.

Author(s)

Custodian: Luis Mijangos - Post to https://groups.google.com/d/forum/dartr

See Also

```
gl.report.ld.map
Other matched filter: gl.filter.callrate(), gl.filter.hamming(), gl.filter.locmetric(),
gl.filter.maf(), gl.filter.monomorphs(), gl.filter.overshoot(), gl.filter.pa(), gl.filter.secondaries()
```

gl.filter.locmetric Filters loci on the basis of numeric information stored in other\$loc.metrics in a genlight {adegenet} object

Description

This script uses any field with numeric values stored in \$other\$loc.metrics to filter loci. The loci to keep can be within the upper and lower thresholds ('within') or outside of the upper and lower thresholds ('outside').

Usage

```
gl.filter.locmetric(x, metric, upper, lower, keep = "within", verbose = NULL)
```

Arguments

х	Name of the genlight object containing the SNP data [required].
metric	Name of the metric to be used for filtering [required].
upper	Filter upper threshold [required].
lower	Filter lower threshold [required].
keep	Whether keep loci within of upper and lower thresholds or keep loci outside of upper and lower thresholds [within].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Details

The fields that are included in dartR, and a short description, are found below. Optionally, the user can also set his/her own filter by adding a vector into \$other\$loc.metrics as shown in the example.

- 1. SnpPosition position (zero is position 1) in the sequence tag of the defined SNP variant base.
- 2. CallRate proportion of samples for which the genotype call is non-missing (that is, not '-').
- 3. OneRatioRef proportion of samples for which the genotype score is 0.
- 4. OneRatioSnp proportion of samples for which the genotype score is 2.
- 5. FreqHomRef proportion of samples homozygous for the Reference allele.
- 6. FreqHomSnp proportion of samples homozygous for the Alternate (SNP) allele.
- 7. FreqHets proportion of samples which score as heterozygous, that is, scored as 1.
- 8. PICRef polymorphism information content (PIC) for the Reference allele.
- 9. PICSnp polymorphism information content (PIC) for the SNP.
- 10. AvgPIC average of the polymorphism information content (PIC) of the Reference and SNP alleles.
- 11. AvgCountRef sum of the tag read counts for all samples, divided by the number of samples with non-zero tag read counts, for the Reference allele row.

- 12. AvgCountSnp sum of the tag read counts for all samples, divided by the number of samples with non-zero tag read counts, for the Alternate (SNP) allele row.
- 13. RepAvg proportion of technical replicate assay pairs for which the marker score is consistent.

Value

The reduced genlight dataset.

Author(s)

Luis Mijangos – Post to https://groups.google.com/d/forum/dartr

See Also

```
Other matched filter: gl.filter.callrate(), gl.filter.hamming(), gl.filter.ld(), gl.filter.maf(),
gl.filter.monomorphs(), gl.filter.overshoot(), gl.filter.pa(), gl.filter.secondaries()
```

Examples

```
# adding dummy data
test <- testset.gl
test$other$loc.metrics$test <- 1:nLoc(test)
result <- gl.filter.locmetric(x=test, metric= 'test', upper=255,
lower=200, keep= 'within', verbose=3)</pre>
```

gl.filter.maf	Filters loci on the basis of minor allele frequency (MAF) or minor
	allele count (MAC)

Description

This script calculates the minor allele frequency for each locus and updates the locus metadata for FreqHomRef, FreqHomSnp, FreqHets and MAF (if it exists). It then uses the updated metadata for MAF to filter loci.

Usage

```
gl.filter.maf(
    x,
    threshold = 0.01,
    by.pop = FALSE,
    pop.limit = ceiling(nPop(x)/2),
    ind.limit = 10,
    recalc = FALSE,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
```

gl.filter.maf

```
plot.file = NULL,
plot.dir = NULL,
bins = 25,
verbose = NULL
```

Arguments

)

х	Name of the genlight object containing the SNP data [required].
threshold	Threshold MAF – loci with a MAF less than the threshold will be removed. If a value > 1 is provided it will be interpreted as MAC (i.e. the minimum number of times an allele needs to be observed) [default 0.01].
by.pop	Whether MAF should be calculated by population [default FALSE].
pop.limit	Minimum number of populations in which MAF should be less than the threshold for a locus to be filtered out. Only used if by.pop = TRUE. The default value is half of the populations [default ceiling($nPop(x)/2$)].
ind.limit	Minimum number of individuals that a population should contain to calculate MAF. Only used if by.pop=TRUE [default 10].
recalc	Recalculate the locus metadata statistics [default FALSE].
plot.display	If TRUE, histograms of base composition are displayed in the plot window [de-fault TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.colors	List of two color names for the borders and fill of the plots [default c("#2171B5", "#6BAED6")].
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL].
plot.dir	Directory in which to save files [default = working directory].
bins	Number of bins to display in histograms [default 25].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Details

Careful consideration needs to be given to the settings to be used for this function. When the filter is applied globally (i.e. by.pop=FALSE) but the data include multiple population, there is the risk to remove markers because the allele frequencies is low (at global level) but the allele frequencies for the same markers may be high within some of the populations (especially if the perpopulation sample size is small). Similarly, not always it is a sensible choice to run this function using by.pop=TRUE because allele that are rare in a population may be very common in other, but the (possible) allele frequencies will depend on the sample size within each population. Where the purpose of filtering for MAF is to remove possible spurious alleles (i.e. sequencing errors), it is perhaps better to filter based on the number of times an allele is observed (MAC, Minimum Allele Count), under the assumption that if an allele is observed > MAC, it is fairly rare to be an error.

From v2.1 The threshold can take values > 1. In this case, these are interpreted as a threshold for MAC.

Value

The reduced genlight dataset

Author(s)

Custodian: Luis Mijangos – Post to https://groups.google.com/d/forum/dartr

See Also

```
Other matched filter: gl.filter.callrate(), gl.filter.hamming(), gl.filter.ld(), gl.filter.locmetric(),
gl.filter.monomorphs(), gl.filter.overshoot(), gl.filter.pa(), gl.filter.secondaries()
```

Examples

```
result <- gl.filter.maf(platypus.gl, threshold = 0.05, verbose = 3)
#result <- gl.filter.maf(platypus.gl, by.pop = TRUE, threshold = 0.05, verbose = 3)</pre>
```

gl.filter.monomorphs Filters monomorphic loci, including those with all NAs

Description

This script deletes monomorphic loci from a genlight {adegenet} object A DArT dataset will not have monomorphic loci, but they can arise, along with loci that are scored all NA, when populations or individuals are deleted. Retaining monomorphic loci unnecessarily increases the size of the dataset and will affect some calculations. Note that for SNP data, NAs likely represent null alleles; in tag presence/absence data, NAs represent missing values (presence/absence could not be reliably scored)

Usage

```
gl.filter.monomorphs(x, verbose = NULL)
```

Arguments

х	Name of the input genlight object [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress
	and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Value

A genlight object with monomorphic (and all NA) loci removed.

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

gl.filter.overshoot

See Also

```
Other matched filter: gl.filter.callrate(), gl.filter.hamming(), gl.filter.ld(), gl.filter.locmetric(),
gl.filter.maf(), gl.filter.overshoot(), gl.filter.pa(), gl.filter.secondaries()
```

Examples

```
# SNP data
  result <- gl.filter.monomorphs(testset.gl, verbose=3)
# Tag P/A data
  result <- gl.filter.monomorphs(testset.gs, verbose=3)</pre>
```

gl.filter.overshoot Filters loci for which the SNP has been trimmed from the sequence tag along with the adaptor

Description

This function checks the position of the SNP within the trimmed sequence tag and identifies those for which the SNP position is outside the trimmed sequence tag. This can happen, rarely, when the sequence containing the SNP resembles the adaptor. The SNP genotype can still be used in most analyses, but functions like gl2fasta() will present challenges if the SNP has been trimmed from the sequence tag. Not fatal, but should apply this filter before gl.filter.secondaries, for obvious reasons.

Usage

gl.filter.overshoot(x, verbose = NULL)

Arguments

х	Name of the genlight object [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress
	and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Value

A new genlight object with the recalcitrant loci deleted

Author(s)

Author(s): Arthur Georges; Custodian: Arthur Georges – Post to https://groups.google.com/ d/forum/dartr

See Also

```
Other matched filter: gl.filter.callrate(), gl.filter.hamming(), gl.filter.ld(), gl.filter.locmetric(),
gl.filter.maf(), gl.filter.monomorphs(), gl.filter.pa(), gl.filter.secondaries()
```

Examples

result <- gl.filter.overshoot(testset.gl, verbose=3)</pre>

gl.filter.pa

Filters loci that contain private (and fixed alleles) between two populations

Description

This script is meant to be used prior to gl.nhybrids to maximise the information content of the SNPs used to identify hybrids (currently newhybrids does allow only 200 SNPs). The idea is to use first all loci that have fixed alleles between the potential source populations and then 'fill up' to 200 loci using loci that have private alleles between those. The functions filters for those loci (if invers is set to TRUE, the opposite is returned (all loci that are not fixed and have no private alleles - not sure why yet, but maybe useful.)

Usage

gl.filter.pa(x, pop1, pop2, invers = FALSE, verbose = NULL)

Arguments

х	Name of the genlight object containing the SNP data [required].
pop1	Name of the first parental population (in quotes) [required].
pop2	Name of the second parental population (in quotes) [required].
invers	Switch to filter for all loci that have no private alleles and are not fixed [FALSE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress
	and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Value

The reduced genlight dataset, containing now only fixed and private alleles.

Author(s)

Authors: Bernd Gruber & Ella Kelly (University of Melbourne); Custodian: Luis Mijangos – Post to https://groups.google.com/d/forum/dartr

See Also

```
Other matched filter: gl.filter.callrate(), gl.filter.hamming(), gl.filter.ld(), gl.filter.locmetric(),
gl.filter.maf(), gl.filter.monomorphs(), gl.filter.overshoot(), gl.filter.secondaries()
```

Examples

```
result <- gl.filter.pa(testset.gl, pop1=pop(testset.gl)[1],
pop2=pop(testset.gl)[2],verbose=3)
```

gl.filter.rdepth

Filters loci based on counts of sequence tags scored at a locus (read depth) @family matched filter

Description

SNP datasets generated by DArT report AvgCountRef and AvgCountSnp as counts of sequence tags for the reference and alternate alleles respectively. These can be used to back calculate Read Depth. Fragment presence/absence datasets as provided by DArT (SilicoDArT) provide Average Read Depth and Standard Deviation of Read Depth as standard columns in their report. Filtering on Read Depth using the companion script gl.filter.rdepth can be on the basis of loci with exceptionally low counts, or loci with exceptionally high counts.

Usage

```
gl.filter.rdepth(
    x,
    lower = 5,
    upper = 1000,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
    plot.file = NULL,
    plot.dir = NULL,
    verbose = NULL
)
```

Arguments

x	Name of the genlight object containing the SNP or tag presence/absence data [required].
lower	Lower threshold value below which loci will be removed [default 5].
upper	Upper threshold value above which loci will be removed [default infinite=1000].
plot.display	If TRUE, histograms of base composition are displayed in the plot window [de-fault TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.colors	List of two color names for the borders and fill of the plots [default c("#2171B5", "#6BAED6")].
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL]
plot.dir	Directory in which to save files [default = working directory]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Details

For examples of themes, see:

- https://ggplot2.tidyverse.org/reference/ggtheme.html and
- https://yutannihilation.github.io/allYourFigureAreBelongToUs/ggthemes/

Value

Returns a genlight object retaining loci with a Read Depth in the range specified by the lower and upper threshold.

Author(s)

Custodian: Arthur Georges (Post to https://groups.google.com/d/forum/dartr)

See Also

gl.filter.rdepth

Examples

```
# SNP data
gl.report.rdepth(testset.gl)
result <- gl.filter.rdepth(testset.gl, lower=8, upper=50, verbose=3)
# Tag P/A data
result <- gl.filter.rdepth(testset.gs, lower=8, upper=50, verbose=3)
res <- gl.filter.rdepth(platypus.gl)</pre>
```

gl.filter.reproducibility

Filters loci in a genlight {adegenet} object based on average repeatability of alleles at a locus @family matched filter

Description

SNP datasets generated by DArT have an index, RepAvg, generated by reproducing the data independently for 30 of alleles that give a repeatable result, averaged over both alleles for each locus. SilicoDArT datasets generated by DArT have a similar index, Reproducibility. For these fragment presence/absence data, repeatability is the percentage of scores that are repeated in the technical replicate dataset.

Usage

```
gl.filter.reproducibility(
    x,
    threshold = 0.99,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
    plot.file = NULL,
    plot.dir = NULL,
    verbose = NULL
)
```

Arguments

х	Name of the genlight object containing the SNP data [required].
threshold	Threshold value below which loci will be removed [default 0.99].
plot.display	If TRUE, histograms of base composition are displayed in the plot window [de-fault TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.colors	List of two color names for the borders and fill of the plots [default c("#2171B5", "#6BAED6")].
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL]
plot.dir	Directory in which to save files [default = working directory]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Value

Returns a genlight object retaining loci with repeatability (Repavg or Reproducibility) greater than the specified threshold.

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

gl.report.reproducibility

Examples

```
# SNP data
gl.report.reproducibility(testset.gl)
result <- gl.filter.reproducibility(testset.gl, threshold=0.99, verbose=3)
# Tag P/A data
gl.report.reproducibility(testset.gs)</pre>
```

```
result <- gl.filter.reproducibility(testset.gs, threshold=0.99)
res <- gl.filter.reproducibility(testset.gl)</pre>
```

gl.filter.secondaries Filters loci that represent secondary SNPs in a genlight object

Description

SNP datasets generated by DArT include fragments with more than one SNP and record them separately with the same CloneID (=AlleleID). These multiple SNP loci within a fragment (secondaries) are likely to be linked, and so you may wish to remove secondaries. This script filters out all but the first sequence tag with the same CloneID after ordering the genlight object on based on repeatability, avgPIC in that order (method='best') or at random (method='random'). The filter has not been implemented for tag presence/absence data.

Usage

```
gl.filter.secondaries(x, method = "random", verbose = NULL)
```

Arguments

х	Name of the genlight object containing the SNP data [required].
method	Method of selecting SNP locus to retain, 'best' or 'random' [default 'random'].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Value

The genlight object, with the secondary SNP loci removed.

Author(s)

Custodian: Arthur Georges - Post to https://groups.google.com/d/forum/dartr

See Also

```
Other matched filter: gl.filter.callrate(), gl.filter.hamming(), gl.filter.ld(), gl.filter.locmetric(),
gl.filter.maf(), gl.filter.monomorphs(), gl.filter.overshoot(), gl.filter.pa()
```

Examples

```
gl.report.secondaries(testset.gl)
result <- gl.filter.secondaries(testset.gl)</pre>
```

gl.filter.taglength Filters loci in a genlight {adegenet} object based on sequence tag length @family matched filter

Description

SNP datasets generated by DArT typically have sequence tag lengths ranging from 20 to 69 base pairs.

Usage

```
gl.filter.taglength(x, lower = 20, upper = 69, verbose = NULL)
```

Arguments

х	Name of the genlight object containing the SNP data [required].
lower	Lower threshold value below which loci will be removed [default 20].
upper	Upper threshold value above which loci will be removed [default 69].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Value

Returns a genlight object retaining loci with a sequence tag length in the range specified by the lower and upper threshold.

Author(s)

Custodian: Arthur Georges - Post to https://groups.google.com/d/forum/dartr

Examples

```
# SNP data
gl.report.taglength(testset.gl)
result <- gl.filter.taglength(testset.gl,lower=60)
gl.report.taglength(result)
# Tag P/A data
gl.report.taglength(testset.gs)
result <- gl.filter.taglength(testset.gs,lower=60)
gl.report.taglength(result)</pre>
```

gl.fixed.diff

Description

This script takes SNP data or sequence tag P/A data grouped into populations in a genlight object (DArTSeq) and generates a matrix of fixed differences between populations taken pairwise

Usage

```
gl.fixed.diff(
    x,
    tloc = 0,
    test = FALSE,
    delta = 0.02,
    alpha = 0.05,
    reps = 1000,
    mono.rm = TRUE,
    pb = FALSE,
    verbose = NULL
)
```

Arguments

x	Name of the genlight object containing SNP genotypes or tag P/A data (Silico-DArT) or an object of class 'fd' [required].
tloc	Threshold defining a fixed difference (e.g. 0.05 implies 95:5 vs 5:95 is fixed) [default 0].
test	If TRUE, calculate p values for the observed fixed differences [default FALSE].
delta	Threshold value for the true population minor allele frequency (MAF) from which resultant sample fixed differences are considered true positives [default 0.02].
alpha	Level of significance used to display non-significant differences between populations as they are compared pairwise [default 0.05].
reps	Number of replications to undertake in the simulation to estimate probability of false positives [default 1000].
mono.rm	If TRUE, loci that are monomorphic across all individuals are removed before beginning computations [default TRUE].
pb	If TRUE, show a progress bar on time consuming loops [default FALSE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

gl.fixed.diff

Details

A fixed difference at a locus occurs when two populations share no alleles or where all members of one population has a sequence tag scored, and all members of the other population has the sequence tag absent. The challenge with this approach is that when sample sizes are finite, fixed differences will occur through sampling error, compounded when many loci are examined. Simulations suggest that sample sizes of $n_{1=5}$ and $n_{2=5}$ are adequate to reduce the probability of [experiment-wide] type 1 error to negligible levels [ploidy=2]. A warning is issued if comparison between two populations involves sample sizes less than 5, taking into account allele drop-out. Optionally, if test=TRUE, the script will test the fixed differences between final OTUs for statistical significance, using simulation, and then further amalgamate populations that for which there are no significant fixed differences at a specified level of significance (alpha). To avoid conflation of true fixed differences with false positives in the simulations, it is necessary to decide a threshold value (delta) for extreme true allele frequencies that will be considered fixed for practical purposes. That is, fixed differences in the sample set will be considered to be positives (not false positives) if they arise from true allele frequencies of less than 1-delta in one or both populations. The parameter delta is typically set to be small (e.g. delta = 0.02). NOTE: The above test will only be calculated if tloc=0, that is, for analyses of absolute fixed differences. The test applies in comparisons of allopatric populations only. For sympatric populations, use gl.pval.sympatry(). An absolute fixed difference is as defined above. However, one might wish to score fixed differences at some lower level of allele frequency difference, say where percent allele frequencies are 95,5 and 5,95 rather than 100:0 and 0:100. This adjustment can be done with the tloc parameter. For example, tloc=0.05 means that SNP allele frequencies of 95,5 and 5,95 percent will be regarded as fixed when comparing two populations at a locus.

Value

A list of Class 'fd' containing the gl object and square matrices, as follows:

- 1. \$gl the output genlight object;
- 2. \$fd raw fixed differences;
- 3. \$pcfd percent fixed differences;
- 4. \$nobs mean no. of individuals used in each comparison;
- 5. \$nloc total number of loci used in each comparison;
- \$expfpos if test=TRUE, the expected count of false positives for each comparison [by simulation];
- \$sdfpos if test=TRUE, the standard deviation of the count of false positives for each comparison [by simulation];
- 8. \$pval if test=TRUE, the significance of the count of fixed differences [by simulation])

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

utils.is.fixed

Examples

```
fd <- gl.fixed.diff(testset.gl, tloc=0, verbose=3 )
fd <- gl.fixed.diff(testset.gl, tloc=0, test=TRUE, delta=0.02, reps=100, verbose=3 )</pre>
```

gl.fst.pop	Calculates a pairwise Fst values for populations in a genlight object
	This script calculates pairwise Fst values based on the implementa-
	tion in the StAMPP package (?stamppFst). It allows to run bootstrap
	to estimate probability of Fst values to be different from zero. For de-
	tailed information please check the help pages (?stamppFst).

Description

Calculates a pairwise Fst values for populations in a genlight object This script calculates pairwise Fst values based on the implementation in the StAMPP package (?stamppFst). It allows to run bootstrap to estimate probability of Fst values to be different from zero. For detailed information please check the help pages (?stamppFst).

Usage

```
gl.fst.pop(x, nboots = 1, percent = 95, nclusters = 1, verbose = NULL)
```

Arguments

х	Name of the genlight containing the SNP genotypes [required].
nboots	Number of bootstraps to perform across loci to generate confidence intervals and p-values [default 1].
percent	Percentile to calculate the confidence interval around [default 95].
nclusters	Number of processor threads or cores to use during calculations [default 1].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Value

A matrix of distances between populations (class dist), if nboots =1, otherwise a list with Fsts (in a matrix), Pvalues (a matrix of pvalues), Bootstraps results (data frame of all runs). Hint: Use as.matrix(as.dist(fsts)) if you want to have a squared matrix with symmetric entries returned, instead of a dist object.

Author(s)

Bernd Gruber (bugs? Post to https://groups.google.com/d/forum/dartr)

gl.He

Examples

```
test <- gl.filter.callrate(platypus.gl,threshold = 1)
test <- gl.filter.monomorphs(test)
out <- gl.fst.pop(test, nboots=1)</pre>
```

gl.He

Estimates expected Heterozygosity

Description

Estimates expected Heterozygosity

Usage

gl.He(gl)

Arguments

gl A genlight object [required]

Value

A simple vector whit Ho for each loci

Author(s)

Bernd Gruber (Post to https://groups.google.com/d/forum/dartr)

gl.Ho

Estimates observed Heterozygosity

Description

Estimates observed Heterozygosity

Usage

gl.Ho(gl)

Arguments

gl A genlight object [required]

Value

A simple vector whit Ho for each loci

Author(s)

Bernd Gruber (bugs? Post to https://groups.google.com/d/forum/dartr)

gl.hwe.pop Performs Hardy-Weinberg tests over loci and populations

Description

Hardy-Weinberg tests are performed for each loci in each of the populations as defined by the pop slot in a genlight object.

Usage

```
gl.hwe.pop(
    x,
    alpha_val = 0.05,
    plot.out = TRUE,
    plot_theme = theme_dartR(),
    plot_colors = c("gray90", "deeppink"),
    HWformat = FALSE,
    verbose = NULL
)
```

Arguments

x	A genlight object with a population defined $[pop(x) \text{ does not return NULL}]$.
alpha_val	Level of significance for testing [default 0.05].
plot.out	If TRUE, returns a plot object compatible with ggplot, otherwise returns a dataframe [default TRUE].
plot_theme	User specified theme [default theme_dartR()].
plot_colors	Vector with two color names for the borders and fill [default gl.colors(2)]. [default gl.colors("dis")].
HWformat	Switch if data should be returned in HWformat (counts of Genotypes to be used in package HardyWeinberg)
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity].

Details

This function employs the HardyWeinberg package, which needs to be installed. The function that is used is HWExactStats, but there are several other great functions implemented in the package regarding HWE. Therefore, this function can return the data in the format expected by the HWE package expects, via HWformat=TRUE and then use this to run other functions of the package. This functions performs a HWE test for every population (rows) and loci (columns) and returns a true false matrix. True is reported if the p-value of an HWE-test for a particular loci and population

gl.impute

was below the specified threshold (alpha_val, default=0.05). The thinking behind this approach is that loci that are not in HWE in several populations have most likely to be treated (e.g. filtered if loci under selection are of interest). If plot=TRUE a barplot on the loci and the sum of deviation over all population is returned. Loci that deviate in the majority of populations can be identified via colSums on the resulting matrix. Plot themes can be obtained from

- https://ggplot2.tidyverse.org/reference/ggtheme.html and
- https://yutannihilation.github.io/allYourFigureAreBelongToUs/ggthemes/

Resultant ggplots and the tabulation are saved to the session's temporary directory.

Value

The function returns a list with up to three components:

- 'HWE' is the matrix over loci and populations
- 'plot' is a plot (ggplot) which shows the significant results for population and loci (can be amended further using ggplot syntax)
- 'HWEformat=TRUE' the 'HWformat' entails SNP data for each population in 'HardyWeinberg'format to be used with other functions of the package (e.g HWPerm or HWExactPrevious).

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

Examples

out <- gl.hwe.pop(bandicoot.gl[,1:33], alpha_val=0.05, plot.out=TRUE, HWformat=FALSE)</pre>

gl.impute

Imputes missing data

Description

This function imputes genotypes on a population-by-population basis, where populations can be considered panmictic, or imputes the state for presence-absence data.

Usage

```
gl.impute(
    x,
    method = "neighbour",
    fill.residual = TRUE,
    parallel = FALSE,
    verbose = NULL
)
```

Arguments

x	Name of the genlight object containing the SNP or presence-absence data [re-quired].
method	Imputation method, either "frequency" or "HW" or "neighbour" or "random" [default "neighbour"].
fill.residual	Should any residual missing values remaining after imputation be set to 0, 1, 2 at random, taking into account global allele frequencies at the particular locus [default TRUE].
parallel	A logical indicating whether multiple cores -if available- should be used for the computations (TRUE), or not (FALSE); requires the package parallel to be installed [default FALSE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Details

We recommend that imputation be performed on sampling locations, before any aggregation. Imputation is achieved by replacing missing values using either of four methods:

- If "frequency", genotypes scored as missing at a locus in an individual are imputed using the average allele frequencies at that locus in the population from which the individual was drawn.
- If "HW", genotypes scored as missing at a locus in an individual are imputed by sampling at random assuming Hardy-Weinberg equilibrium. Applies only to genotype data.
- If "neighbour", substitute the missing values for the focal individual with the values taken from the nearest neighbour. Repeat with next nearest and so on until all missing values are replaced.
- if "random", missing data are substituted by random values (0, 1 or 2).

The nearest neighbour is the one at the smallest Euclidean distancefrom the focal individual The advantage of this approach is that it works regardless of how many individuals are in the population to which the focal individual belongs, and the displacement of the individual is haphazard as opposed to: (a) Drawing the individual toward the population centroid (HW and Frequency). (b) Drawing the individual toward the global centroid (glPCA). Note that loci that are missing for all individuals in a population are not imputed with method 'frequency' or 'HW' and can give unpredictable results for particular individuals using 'neighbour'. Consider using the function gl.filter.allna with by.pop=TRUE to remove them first.

Value

A genlight object with the missing data imputed.

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

gl.join

See Also

```
Other data manipulation: gl.define.pop(), gl.drop.ind(), gl.drop.loc(), gl.drop.pop(),
gl.edit.recode.pop(), gl.join(), gl.keep.ind(), gl.keep.loc(), gl.keep.pop(), gl.make.recode.ind(),
gl.merge.pop(), gl.reassign.pop(), gl.recode.ind(), gl.recode.pop(), gl.rename.pop(),
gl.sample(), gl.sim.genotypes(), gl.sort(), gl.subsample.ind(), gl.subsample.loc()
```

Examples

```
require("dartR.data")
# SNP genotype data
gl <- gl.filter.callrate(platypus.gl,threshold=0.95)
gl <- gl.filter.allna(gl)
gl <- gl.impute(gl,method="neighbour")
# Sequence Tag presence-absence data
gs <- gl.filter.callrate(testset.gs,threshold=0.95)
gl <- gl.filter.allna(gl)
gs <- gl.impute(gs, method="neighbour")
gs <- gl.impute(platypus.gl,method ="random")</pre>
```

gl.join

Combines two dartR genlight objects

Description

This function combines two genlight objects and their associated metadata. The history associated with the two genlight objects is cleared from the new genlight object. The individuals/samples must be the same in each genlight object. The function is typically used to combine datasets from the same service where the files have been split because of size limitations. The data is read in from multiple csv files, then the resultant genlight objects are combined. This function works with both SNP and Tag P/A data.

Usage

```
gl.join(x1, x2, method = "sidebyside", verbose = NULL)
```

Arguments

x1	Name of the first genlight object [required].
x2	Name of the first genlight object [required].
method	If method='sidebyside' then combine the two by bringing the loci together against the same set of individuals; If method='end2end' then combine the two by bringing two sets of individuals together against the same set of loci [default 'sidebyside']
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Details

This script joins two genlight objects together along with the associated metadata. if method='sidebyside' (the default), the individuals in the two genlight objects must be the same and in the same order. The loci are combined.

If method='end2end', the loci in the two genlight objects must be the same and in the same order. The data for the two sets of individuals are combined. Note that if two individuals have the same names, they will be made unique.#'

Value

A new genlight object

Author(s)

Custodian: Arthur Georges - Post to https://groups.google.com/d/forum/dartr

See Also

```
Other data manipulation: gl.define.pop(), gl.drop.ind(), gl.drop.loc(), gl.drop.pop(),
gl.edit.recode.pop(), gl.impute(), gl.keep.ind(), gl.keep.loc(), gl.keep.pop(), gl.make.recode.ind(),
gl.merge.pop(), gl.reassign.pop(), gl.recode.ind(), gl.recode.pop(), gl.rename.pop(),
gl.sample(), gl.sim.genotypes(), gl.sort(), gl.subsample.ind(), gl.subsample.loc()
```

Examples

```
x1 <- testset.gl[,1:100]</pre>
x1@other$loc.metrics <- testset.gl@other$loc.metrics[1:100,]</pre>
nLoc(x1)
x2 <- testset.gl[,101:150]</pre>
x2@other$loc.metrics <- testset.gl@other$loc.metrics[101:150,]</pre>
nLoc(x2)
gl <- gl.join(x1, x2, verbose=2)</pre>
nLoc(gl)
x1 <- testset.gl[,1:100]</pre>
x1@other$loc.metrics <- testset.gl@other$loc.metrics[1:100,]</pre>
nLoc(x1)
x2 <- testset.gl[,1:100]</pre>
x2@other$loc.metrics <- testset.gl@other$loc.metrics[1:100,]</pre>
nLoc(x2)
gl <- gl.join(x1, x2, method="end2end", verbose=2)</pre>
nInd(gl)
```

gl.keep.ind

Description

This script deletes all individuals apart from those listed (ind.list). Monomorphic loci and loci that are scored all NA are optionally deleted (mono.rm=TRUE). The script also optionally recalculates locus metatdata statistics to accommodate the deletion of individuals from the dataset (recalc=TRUE). The script returns a dartR genlight object with the retained individuals and the recalculated locus metadata. The script works with both genlight objects containing SNP genotypes and Tag P/A data (SilicoDArT).

Usage

```
gl.keep.ind(x, ind.list, recalc = FALSE, mono.rm = FALSE, verbose = NULL)
```

Arguments

Х	Name of the genlight object [required].
ind.list	A list of individuals to be retained [required].
recalc	If TRUE, recalculate the locus metadata statistics [default FALSE].
mono.rm	If TRUE, remove monomorphic and all NA loci [default FALSE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress
	and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

A reduced dartR genlight object

Author(s)

Custodian: Arthur Georges - Post to https://groups.google.com/d/forum/dartr

See Also

gl.drop.pop to drop rather than keep specified populations

```
Other data manipulation: gl.define.pop(), gl.drop.ind(), gl.drop.loc(), gl.drop.pop(),
gl.edit.recode.pop(), gl.impute(), gl.join(), gl.keep.loc(), gl.keep.pop(), gl.make.recode.ind(),
gl.merge.pop(), gl.reassign.pop(), gl.recode.ind(), gl.recode.pop(), gl.rename.pop(),
gl.sample(), gl.sim.genotypes(), gl.sort(), gl.subsample.ind(), gl.subsample.loc()
```

Examples

```
# SNP data
gl2 <- gl.keep.ind(testset.gl, ind.list=c('AA019073','AA004859'))
# Tag P/A data
gs2 <- gl.keep.ind(testset.gs, ind.list=c('AA020656','AA19077','AA004859'))</pre>
```

```
gl.keep.loc
```

Description

This function deletes loci that are not specified to keep, and their associated metadata. The script returns a dartR genlight object with the retained loci. The script works with both genlight objects containing SNP genotypes and Tag P/A data (SilicoDArT).

Usage

```
gl.keep.loc(x, loc.list = NULL, first = NULL, last = NULL, verbose = NULL)
```

Arguments

х	Name of the genlight object [required].
loc.list	A list of loci to be kept [required, if loc.range not specified].
first	First of a range of loci to be kept [required, if loc.list not specified].
last	Last of a range of loci to be kept [if not specified, last locus in the dataset].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress but not results; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

A genlight object with the reduced data

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

gl.drop.loc to drop rather than keep specified loci

```
Other data manipulation: gl.define.pop(), gl.drop.ind(), gl.drop.loc(), gl.drop.pop(),
gl.edit.recode.pop(), gl.impute(), gl.join(), gl.keep.ind(), gl.keep.pop(), gl.make.recode.ind(),
gl.merge.pop(), gl.reassign.pop(), gl.recode.ind(), gl.recode.pop(), gl.rename.pop(),
gl.sample(), gl.sim.genotypes(), gl.sort(), gl.subsample.ind(), gl.subsample.loc()
```

Examples

```
# SNP data
gl2 <- gl.keep.loc(testset.gl, loc.list=c('100051468|42-A/T', '100049816-51-A/G'))
# Tag P/A data
gs2 <- gl.keep.loc(testset.gs, loc.list=c('20134188','19249144'))</pre>
```

gl.keep.pop

Description

Individuals are assigned to populations based on associated specimen metadata stored in the dartR genlight object. This script deletes all individuals apart from those in listed populations (pop.list). Monomorphic loci and loci that are scored all NA are optionally deleted (mono.rm=TRUE). The script also optionally recalculates locus metadata statistics to accommodate the deletion of individuals from the dataset (recalc=TRUE). The script returns a dartR genlight object with the retained populations and the recalculated locus metadata. The script works with both genlight objects containing SNP genotypes and Tag P/A data (SilicoDArT).

Usage

```
gl.keep.pop(
    x,
    pop.list,
    as.pop = NULL,
    recalc = FALSE,
    mono.rm = FALSE,
    verbose = NULL
)
```

Arguments

х	Name of the genlight object [required].
pop.list	List of populations to be retained [required].
as.pop	Temporarily assign another locus metric as the population for the purposes of deletions [default NULL].
recalc	If TRUE, recalculate the locus metadata statistics [default FALSE].
mono.rm	If TRUE, remove monomorphic and all NA loci [default FALSE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress but not results; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

A reduced dartR genlight object

Author(s)

Custodian: Arthur Georges - Post to https://groups.google.com/d/forum/dartr

See Also

gl.drop.pop to drop rather than keep specified populations

```
Other data manipulation: gl.define.pop(), gl.drop.ind(), gl.drop.loc(), gl.drop.pop(),
gl.edit.recode.pop(), gl.impute(), gl.join(), gl.keep.ind(), gl.keep.loc(), gl.make.recode.ind(),
gl.merge.pop(), gl.reassign.pop(), gl.recode.ind(), gl.recode.pop(), gl.rename.pop(),
gl.sample(), gl.sim.genotypes(), gl.sort(), gl.subsample.ind(), gl.subsample.loc()
```

Examples

```
# SNP data
gl2 <- gl.keep.pop(testset.gl, pop.list=c('EmsubRopeMata', 'EmvicVictJasp'))
gl2 <- gl.keep.pop(testset.gl, pop.list=c('EmsubRopeMata', 'EmvicVictJasp'),
mono.rm=TRUE,recalc=TRUE)
gl2 <- gl.keep.pop(testset.gl, pop.list=c('Female'),as.pop='sex')
# Tag P/A data
gs2 <- gl.keep.pop(testset.gs, pop.list=c('EmsubRopeMata','EmvicVictJasp'))</pre>
```

gl.load

Loads an object from compressed binary format produced by gl.save()

Description

This is a wrapper for readRDS() The function loads the object from the current workspace, checks if it is a dartR genlight object, converts it if it is not, and returns the gl object. A compliance check can be requested.

Usage

gl.load(file, compliance = FALSE, verbose = NULL)

Arguments

file	Name of the file to receive data [required].
compliance	Whether to undertake a compliance check [default FALSE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

The loaded object

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

gl.mahal.assign

See Also

```
gl.save
```

```
Other io: gl.read.csv(), gl.read.dart(), gl.read.fasta(), gl.read.silicodart(), gl.save(),
gl.write.csv(), utils.read.dart()
```

gl.mahal.assign Assigns individuals to populations with an associated probability

Description

Uses Mahalanobis Distances between individuals and group centroids to calculate probability of group membership

Usage

```
gl.mahal.assign(
    x,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
    plot.file = NULL,
    plot.dir = NULL,
    verbose = NULL
)
```

Arguments

х	Name of the genlight object [required].
plot.display	If TRUE, resultant plots are displayed in the plot window [default TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.colors	List of two color names for the borders and fill of the plots [default c("#2171B5", "#6BAED6")]
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL]
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity]

Details

The function generates 200 simulated individuals for each group (=population in the genlight object) drawing from the observed allele frequencies for each group. The group centroids and covariance matricies are calculated for these simulated groups. The covariance matrix is inverted using package MASS::ginv to overcome the singularities that would otherwise arise with typical SNP data. Mahanobilis Distances are calculated using stats::mahanalobis for each individual in the dataset

and associated Chi Square probabilities of group membership are calculated for each individual in the original genlight object. The resultant table can be used for decisions on group membership. A special group (=population in the genlight object) called 'unknowns' can be used to specifically identify individuals with unknown group membership.

A color vector can be obtained with gl.select.colors() and then passed to the function with the plot.colors parameter. Themes can be obtained from in

- https://ggplot2.tidyverse.org/reference/ggtheme.html and
- https://yutannihilation.github.io/allYourFigureAreBelongToUs/ggthemes/

If a plot.file is given, the ggplot arising from this function is saved as an "RDS" binary file using saveRDS(); can be reloaded with readRDS(). A file name must be specified for the plot to be saved. If a plot directory (plot.dir) is specified, the ggplot binary is saved to that directory; otherwise to the tempdir().

Value

The unchanged genlight object

Author(s)

Custodian: Arthur Georges - Post to https://groups.google.com/d/forum/dartr

See Also

```
Other matched reports: gl.report.bases(), gl.report.factorloadings(), gl.report.fstat(),
gl.report.monomorphs()
```

gl.make.recode.ind *Creates a proforma recode_ind file for reassigning individual (=spec-imen) names*

Description

Renaming individuals may be required when there have been errors in labeling arising in the process from sample to sequencing files. There may be occasions where renaming individuals is required for preparation of figures.

Usage

```
gl.make.recode.ind(
    x,
    out.recode.file = "default_recode_ind.csv",
    outpath = NULL,
    verbose = NULL
)
```

Arguments

х	Name of the genlight object [required].	
out.recode.file		
	File name of the output file (including extension) [default default_recode_ind.csv].	
outpath	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]	
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].	

Details

This function facilitates the construction of a recode table by producing a proforma file with current individual (=specimen) names in two identical columns. Edit the second column to reassign individual names. Use keyword 'Delete' to delete an individual. When caution needs to be exercised because of the potential for breaking the 'chain of evidence' associated with the samples, recoding individuals using a recode table (csv) can provide a clear record of the changes. Use outpath=getwd() or when calling this function to direct output files to your working directory. The function works with both genlight objects containing SNP genotypes and Tag P/A data (Silico-DArT). Apply the recoding using gl.recode.ind().

Value

A vector containing the new individual names.

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

```
Other data manipulation: gl.define.pop(), gl.drop.ind(), gl.drop.loc(), gl.drop.pop(),
gl.edit.recode.pop(), gl.impute(), gl.join(), gl.keep.ind(), gl.keep.loc(), gl.keep.pop(),
gl.merge.pop(), gl.reassign.pop(), gl.recode.ind(), gl.recode.pop(), gl.rename.pop(),
gl.sample(), gl.sim.genotypes(), gl.sort(), gl.subsample.ind(), gl.subsample.loc()
```

Examples

result <- gl.make.recode.ind(testset.gl, out.recode.file ='Emmac_recode_ind.csv',outpath=tempdir())</pre>

gl.make.recode.pop

Creates a proforma recode_pop_table file for reassigning population names @family data manipulation

Description

Renaming populations may be required when there have been errors in assignment arising in the process from sample to sequence files or when one wishes to amalgamate populations, or delete populations. Recoding populations can also be done with a recode table (csv).

Usage

```
gl.make.recode.pop(
    x,
    out.recode.file = "recode_pop_table.csv",
    outpath = NULL,
    verbose = NULL
)
```

Arguments

Х	Name of the genlight object [required].	
out.recode.file		
	File name of the output file (including extension) [default recode_pop_table.csv].	
outpath	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]	
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].	

Details

This function facilitates the construction of a recode table by producing a proforma file with current population names in two identical columns. Edit the second column to reassign populations. Use keyword 'Delete' to delete a population. When caution needs to be exercised because of the potential for breaking the 'chain of evidence' associated with the samples, recoding individuals using a recode table (csv) can provide a clear record of the changes. Use outpath=getwd() or when calling this function to direct output files to your working directory. The function works with both genlight objects containing SNP genotypes and Tag P/A data (SilicoDArT). Apply the recoding using gl.recode.pop().

Value

A vector containing the new population names.

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr
gl.map.interactive

Examples

result <- gl.make.recode.pop(testset.gl,out.recode.file='test.csv',outpath=tempdir(),verbose=2)</pre>

gl.map.interactive Creates an interactive map (based on latlon) from a genlight object

Description

Creates an interactive map (based on latlon) from a genlight object

Usage

```
gl.map.interactive(
  х,
 matrix = NULL,
  standard = TRUE,
  symmetric = TRUE,
  pop.labels = TRUE,
  pop.labels.cex = 12,
  ind.circles = TRUE,
  ind.circle.cols = rainbow,
  ind.circle.cex = 10,
  ind.circle.transparency = 0.8,
  palette.links = NULL,
  legend.title = NULL,
  provider = "Esri.NatGeoWorldMap",
  raster.image = NULL,
  raster.opacity = 0.5,
  raster.colors = (scales::viridis_pal(option = "D"))(255),
  verbose = NULL
)
```

Arguments

x	A genlight object (including coordinates within the latlon slot) [required].
matrix	A distance matrix between populations or individuals. The matrix is visualised as lines between individuals/populations. If matrix is asymmetric two lines with arrows are plotted [default NULL].
standard	If a matrix is provided line width will be standardised to be between 1 to 10, if set to true, otherwise taken as given [default TRUE].
symmetric	If a symmetric matrix is provided only one line is drawn based on the lower triangle of the matrix. If set to false arrows indicating the direction are used instead [default TRUE].
pop.labels	Population labels at the center of the individuals of populations [default TRUE].
<pre>pop.labels.cex</pre>	Size of population labels [default 12].

ind.circles	Should individuals plotted as circles [default TRUE].
ind.circle.cols	5
	Colors of circles. A color palette or a vectot with as many colors as there are populations in the dataset [default rainbow].
ind.circle.cex	Size or circles in pixels [default 10].
ind.circle.trar	isparency
	Transparency of circles between 0=invisible and 1=no transparency. Defaults to 0.8.
palette.links	Color palette for the links in case a matrix is provided [default NULL].
legend.title	Legend's title for the links in case a matrix is provided [default NULL].
provider	Passed to leaflet [default "Esri.NatGeoWorldMap"].
raster.image	Path to a georeferenced raster image to plot [default NULL].
raster.opacity	The opacity of the raster, expressed from 0 to 1 [default 0.5].
raster.colors	The color palette to use to color the raster values [default scales::viridis_pal(option = "D")(255)].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Details

A wrapper around the **leaflet** package. For possible background maps check as specified via the provider: http://leaflet-extras.github.io/leaflet-providers/preview/index.html

The palette.links argument can be any of the following: A character vector of RGB or named colors. Examples: palette(), c("#000000", "#0000FF", "#FFFFFF"), topo.colors(10)

The name of an RColorBrewer palette, e.g. "BuPu" or "Greens".

The full name of a viridis palette: "viridis", "magma", "inferno", or "plasma".

A function that receives a single value between 0 and 1 and returns a color. Examples: color-Ramp(c("#000000", "#FFFFFF"), interpolate = "spline").

Value

plots a map

Author(s)

Bernd Gruber - Post to https://groups.google.com/d/forum/dartr

See Also

```
Other graphics: gl.colors(), gl.plot.heatmap(), gl.report.ld.map(), gl.select.colors(),
gl.select.shapes(), gl.smearplot(), gl.tree.nj()
```

gl.merge.pop

Examples

```
require("dartR.data")
gl.map.interactive(bandicoot.gl)
cols <- c("red","blue","yellow")
gl.map.interactive(platypus.gl, ind.circle.cols=cols, ind.circle.cex=10,
ind.circle.transparency=0.5)</pre>
```

gl.merge.pop

Merges two or more populations in a dartR genlight object into one population

Description

Individuals are assigned to populations based on the specimen metadata data file (csv) used with gl.read.dart(). This function assigns individuals from two nominated populations into a new single population. It can also be used to rename populations. The function works with both SNP and Tag P/A (silicoDArT) data. The function returns a genlight object with the new population assignments.

Usage

gl.merge.pop(x, old = NULL, new = NULL, verbose = NULL)

Arguments

х	Name of the genlight object [required].
old	A list of populations to be merged [required].
new	Name of the new population [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

A genlight object with the new population assignments.

Author(s)

Custodian: Arthur Georges - Post to https://groups.google.com/d/forum/dartr

See Also

```
Other data manipulation: gl.define.pop(), gl.drop.ind(), gl.drop.loc(), gl.drop.pop(),
gl.edit.recode.pop(), gl.impute(), gl.join(), gl.keep.ind(), gl.keep.loc(), gl.keep.pop(),
gl.make.recode.ind(), gl.reassign.pop(), gl.recode.ind(), gl.recode.pop(), gl.rename.pop(),
gl.sample(), gl.sim.genotypes(), gl.sort(), gl.subsample.ind(), gl.subsample.loc()
```

Examples

gl <- gl.merge.pop(testset.gl, old=c('EmsubRopeMata', 'EmvicVictJasp'), new='Outgroup')</pre>

gl.	рсоа
-----	------

Ordination applied to genotypes in a genlight object (PCA), in an fd object, or to a distance matrix (PCoA)

Description

This function takes the genotypes for individuals and undertakes a Pearson Principal Component analysis (PCA) on SNP or Sequence tag P/A (SilicoDArT) data; it undertakes a Gower Principal Coordinate analysis (PCoA) if supplied with a distance matrix.

Usage

```
gl.pcoa(
    x,
    nfactors = 5,
    pc.select = "broken-stick",
    correction = NULL,
    mono.rm = TRUE,
    parallel = FALSE,
    n.cores = 1,
    plot.out = TRUE,
    plot.out = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = gl.colors(2),
    plot.file = NULL,
    plot.dir = NULL,
    verbose = NULL
)
```

Arguments

x	Name of the genlight object or fd object containing the SNP data, or a distance matrix of type dist [required].
nfactors	Number of axes to retain in the output of factor scores [default 5].
pc.select	Method for identifying substantial PC axes. One of Kaiser-Guttman, broken- stick, Tracy-Widom [default broken-stick]
correction	Method applied to correct for negative eigenvalues, either 'lingoes' or 'cailliez' [Default NULL].
mono.rm	If TRUE, remove monomorphic loci [default TRUE].
parallel	TRUE if parallel processing is required (does fail under Windows) [default FALSE].
n.cores	Number of cores to use if parallel processing is requested [default 16].

gl.pcoa

plot.out	If TRUE, a diagnostic plot is displayed showing a scree plot for the "informa- tive" axes and a histogram of eigenvalues of the remaining "noise" axes [Default TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.colors	List of two color names for the borders and fill of the plot [default gl.colors(2)].
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL]
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]
verbose	verbose= 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Details

The function is essentially a wrapper for glPca {adegenet} or pcoa {ape} with default settings apart from those specified as parameters in this function. **Sources of stress in the visual representation** While, technically, any distance matrix can be represented in an ordinated space, the representation will not typically be exact. There are three major sources of stress in a reduced-representation of distances or dissimilarities among entities using PCA or PCoA. By far the greatest source comes from the decision to select only the top two or three axes from the ordinated set of axes derived from the PCA or PCoA. The representation of the entities such a heavily reduced space will not faithfully represent the distances in the input distance matrix simply because of the loss of information in deeper informative dimensions. For this reason, it is not sensible to be too precious about managing the other two sources of stress in the visual representation. The measure of distance between entities in a PCA is the Pearson Correlation Coefficient, essentially a standardized Euclidean distance. This is both a metric distance and a Euclidean distance and so the distances in the final ordination are faithful to those between entities in the dataset. Note that missing values are imputed in PCA, and that this can be a source of disparity between the distances between the entities in the dataset and the distances in the ordinated space.

In PCoA, the second source of stress is the choice of distance measure or dissimilarity measure. While pretty much any distance or dissimilarity matrix can be represented in an ordinated space, the distances between entities can be faithfully represented in that space (that is, without stress) only if the distances are metric. Furthermore, for distances between entities to be faithfully represented in a rigid Cartesian space, the distance measure needs to be Euclidean. If this is not the case, the distances between the entities in the ordinated visualized space will not #' exactly represent the distances in the input matrix (stress will be non-zero). This source of stress will be evident as negative eigenvalues in the deeper dimensions.

A third source of stress arises from having a sparse dataset, one with missing values. This affects both PCA and PCoA. If the original data matrix is not fully populated, that is, if there are missing values, then even a Euclidean distance matrix will not necessarily be 'positive definite'. It follows that some of the eigenvalues may be negative, even though the distance metric is Euclidean. This issue is exacerbated when the number of loci greatly exceeds the number of individuals, as is typically the case when working with SNP data. The impact of missing values can be minimized by stringently filtering on Call Rate, albeit with loss of data. An alternative is given in a paper 'Honey, I shrunk the sample covariance matrix' and more recently by Ledoit and Wolf (2018), but their approach has not been implemented here.

Options for imputing missing values while minimizing distortion are provided in the function gl.impute(). The good news is that, unless the sum of the negative eigenvalues, arising from a non-Euclidean distance measure or from missing values, approaches those of the final PCA or PCoA axes to be displayed, the distortion is probably of no practical consequence and certainly not comparable to the stress arising from selecting only two or three final dimensions out of several informative dimensions for the visual representation. **Function's output** Two diagnostic plots are produced. The first is a Scree Plot, showing the percentage variation explained by each of the PCA or PCoA axes, for those axes are considered informative. The scree plot informs a decision on the number of dimensions to be retained in the visual summaries. Various approaches are available for identifying which axes are informative (in terms of containing biologically significant variation) and which are noise axes. The simplest method is to consider only those axes that explain more variance than the original variables on average as being informative (pc.select="Kaiser-Guttman"). A second method (the default) is the broken-stick method (pc.select="broken-stick"). A third method is the Tracy-Widom statistical approach (pc.select="Tracy-Widom").

Once you have the informative axes identified, a judgement call is made as to how many dimensions to retain and present as results. This requires a decision on how much information on structure in the data is to be discarded. Retaining at least those axes that explain 10 The second graph is for diagnostic purposes only. It shows the distribution of eigenvalues for the remaining uninformative (noise) axes, including those with negative eigenvalues. If a plot.file is given, the ggplot arising from this function is saved as an binary file using gl.save(); can be reloaded with gl.load(). A file name must be specified for the plot to be saved. If a plot directory (plot.dir) is specified, the ggplot binary is saved to that directory; otherwise to the tempdir(). Action is recommended (verbose ≥ 2) if the negative eigenvalues are dominant, their sum approaching in magnitude the eigenvalues for axes selected for the final visual solution. Output is a glPca object conforming to adegenet::glPca but with only the following retained.

- \$call The call that generated the PCA/PCoA
- \$eig Eigenvalues All eigenvalues (positive, null, negative).
- \$scores Scores (coefficients) for each individual
- · \$loadings Loadings of each SNP for each principal component

Examples of other themes that can be used can be consulted in

- https://ggplot2.tidyverse.org/reference/ggtheme.html and
- https://yutannihilation.github.io/allYourFigureAreBelongToUs/ggthemes/

PCA was developed by Pearson (1901) and Hotelling (1933), whilst the best modern reference is Jolliffe (2002). PCoA was developed by Gower (1966) while the best modern reference is Legendre & Legendre (1998).

Value

An object of class pcoa containing the eigenvalues and factor scores

Author(s)

Author(s): Arthur Georges and Jesus Castrejon. Custodian: Arthur Georges (Post to https://groups.google.com/d/forum/dartr)

gl.pcoa

References

- Cailliez, F. (1983) The analytical solution of the additive constant problem. Psychometrika, 48, 305-308.
- Gower, J. C. (1966) Some distance properties of latent root and vector methods used in multivariate analysis. Biometrika, 53, 325-338.
- Hotelling, H., 1933. Analysis of a complex of statistical variables into Principal Components. Journal of Educational Psychology 24:417-441, 498-520.
- Jolliffe, I. (2002) Principal Component Analysis. 2nd Edition, Springer, New York.
- Ledoit, O. and Wolf, M. (2018). Analytical nonlinear shrinkage of large-dimensional covariance matrices. University of Zurich, Department of Economics, Working Paper No. 264, Revised version. Available at SSRN: https://ssrn.com/abstract=3047302 or http://dx.doi.org/10.2139/ssrn.3047302
- Legendre, P. and Legendre, L. (1998). Numerical Ecology, Volume 24, 2nd Edition. Elsevier Science, NY.
- Lingoes, J. C. (1971) Some boundary conditions for a monotone analysis of symmetric matrices. Psychometrika, 36, 195-203.
- Pearson, K. (1901). On lines and planes of closest fit to systems of points in space. Philosophical Magazine. Series 6, vol. 2, no. 11, pp. 559-572.

See Also

gl.pcoa.plot

Examples

```
# PCA (using SNP genlight object)
gl <- possums.gl
pca <- gl.pcoa(possums.gl[1:50,],verbose=2)
gl.pcoa.plot(pca,gl)

gs <- testset.gs
levels(pop(gs))<-c(rep('Coast',5),rep('Cooper',3),rep('Coast',5),
rep('MDB',8),rep('Coast',6),'Em.subglobosa','Em.victoriae')
# PCA (using SilicoDArT genlight object)
pca <- gl.pcoa(gs)
gl.pcoa.plot(pca,gs)
# Using a distance matrix
D <- gl.dist.ind(testset.gs, method='jaccard')
pcoa <- gl.pcoa(D,correction="cailliez")
gl.pcoa.plot(pcoa,gs)</pre>
```

gl.pcoa.plot

Description

This script takes output from the ordination generated by gl.pcoa() and plots the individuals classified by population.

Usage

```
gl.pcoa.plot(
 glPca,
  х,
  scale = FALSE,
  ellipse = FALSE,
  plevel = 0.95,
  pop.labels = "pop",
  interactive = FALSE,
  as.pop = NULL,
 hadjust = 1.5,
  vadjust = 1,
  xaxis = 1,
 yaxis = 2,
  zaxis = NULL,
  pt.size = 2,
 pt.colors = NULL,
  pt.shapes = NULL,
  label.size = 1,
  axis.label.size = 1.5,
  plot.file = NULL,
 plot.dir = NULL,
  verbose = NULL
)
```

Arguments

glPca	Name of the PCA or PCoA object containing the factor scores and eigenvalues [required].
x	Name of the genlight object or fd object containing the SNP genotypes or Tag P/A (SilicoDArT) genotypes [required to gain access to metadata].
scale	If TRUE, scale the x and y axes in proportion to % variation explained [default FALSE].
ellipse	If TRUE, display ellipses to encapsulate points for each population [default FALSE].

gl.pcoa.plot

plevel	Value of the percentile for the ellipse to encapsulate points for each population [default 0.95].	
pop.labels	How labels will be added to the plot ['none'l'pop'l'legend', default = 'pop'].	
interactive	If TRUE then the populations are plotted without labels, mouse-over to identify points [default FALSE].	
as.pop	Assign another metric to represent populations for the plot [default NULL].	
hadjust	Horizontal adjustment of label position in 2D plots [default 1.5].	
vadjust	Vertical adjustment of label position in 2D plots [default 1].	
xaxis	Identify the x axis from those available in the ordination (xaxis <= nfactors) [default 1].	
yaxis	Identify the y axis from those available in the ordination (yaxis <= nfactors) [default 2].	
zaxis	Identify the z axis from those available in the ordination for a 3D plot (zaxis <= nfactors) [default NULL].	
pt.size	Specify the size of the displayed points [default 2].	
pt.colors	Optionally provide a vector of nPop colors (run gl.select.colors() for color options) [default NULL].	
pt.shapes	Optionally provide a vector of nPop shapes (run gl.select.shapes() for shape options) [default NULL].	
label.size	Specify the size of the point labels [default 1].	
axis.label.size		
	Specify the size of the displayed axis labels [default 1.5].	
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL].	
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()].	
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].	

Details

The factor scores are taken from the output of gl.pcoa() and the population assignments are taken from from the original data file. In the bivariate plots, the specimens are shown optionally with adjacent labels and enclosing ellipses. Population labels on the plot are shuffled so as not to overlap (using package {directlabels}). This can be a bit clunky, as the labels may be some distance from the points to which they refer, but it provides the opportunity for moving labels around using graphics software (e.g. Adobe Illustrator). 3D plotting is activated by specifying a zaxis. Any pair or trio of axes can be specified from the ordination, provided they are within the range of the nfactors value provided to gl.pcoa(). In the 2D plots, axes can be scaled to represent the proportion of variation explained. In any case, the proportion of variation explained by each axis is provided in the axis label. Colors and shapes of the points can be altered by passing a vector of shapes and/or a vector of colors. These vectors can be created with gl.select.shapes() and gl.select.colors() and passed to this script using the pt.shapes and pt.colors parameters. Points displayed in the ordination can be identified if the option interactive=TRUE is chosen, in which case the resultant plot is ggplotly()

friendly. Identification of points is by moving the mouse over them. Refer to the plotly package for further information. The interactive option is automatically enabled for 3D plotting.

If a plot.file is given, the ggplot arising from this function is saved as an "RDS" binary file using saveRDS(); can be reloaded with readRDS(). A file name must be specified for the plot to be saved. If a plot directory (plot.dir) is specified, the ggplot binary is saved to that directory; otherwise to the tempdir().

Value

```
returns no value (i.e. NULL)
```

Author(s)

Custodian: Arthur Georges - Post to https://groups.google.com/d/forum/dartr

See Also

gl.pcoa

Examples

```
test <- gl.pcoa(platypus.gl)</pre>
 gl.pcoa.plot(glPca = test, x = platypus.gl)
# SET UP DATASET
gl <- testset.gl
levels(pop(gl))<-c(rep('Coast',5),rep('Cooper',3),rep('Coast',5),</pre>
rep('MDB',8),rep('Coast',7),'Em.subglobosa','Em.victoriae')
# RUN PCA
pca<-gl.pcoa(gl,nfactors=5)</pre>
# VARIOUS EXAMPLES
gl.pcoa.plot(pca, gl, ellipse=TRUE, plevel=0.95, pop.labels='pop',
axis.label.size=1, hadjust=1.5,vadjust=1)
gl.pcoa.plot(pca, gl, ellipse=TRUE, plevel=0.99, pop.labels='legend',
axis.label.size=1)
gl.pcoa.plot(pca, gl, ellipse=TRUE, plevel=0.99, pop.labels='legend',
axis.label.size=1.5,scale=TRUE)
gl.pcoa.plot(pca, gl, ellipse=TRUE, axis.label.size=1.2, xaxis=1, yaxis=3,
scale=TRUE)
gl.pcoa.plot(pca, gl, pop.labels='none',scale=TRUE)
gl.pcoa.plot(pca, gl, axis.label.size=1.2, interactive=TRUE)
gl.pcoa.plot(pca, gl, ellipse=TRUE, plevel=0.99, xaxis=1, yaxis=2, zaxis=3)
# COLOR AND SHAPE ADJUSTMENTS
shp <- gl.select.shapes(select=c(16,17,17,0,2))</pre>
col <- gl.select.colors(library='brewer',palette='Spectral',ncolors=11,</pre>
select=c(1,9,3,11,11))
gl.pcoa.plot(pca, gl, ellipse=TRUE, plevel=0.95, pop.labels='pop',
pt.colors=col, pt.shapes=shp, axis.label.size=1, hadjust=1.5,vadjust=1)
gl.pcoa.plot(pca, gl, ellipse=TRUE, plevel=0.99, pop.labels='legend',
pt.colors=col, pt.shapes=shp, axis.label.size=1)
# DISTANCE MATRIX
D <- gl.dist.ind(gl)</pre>
```

```
pco <- gl.pcoa(D)
gl.pcoa.plot(pco,gl,ellipse=TRUE)</pre>
```

gl.plot.heatmap Represents a distance matrix as a heatmap

Description

The script plots a heat map to represent the distances in the distance or dissimilarity matrix. This function is a wrapper for heatmap.2 (package gplots).

Usage

```
gl.plot.heatmap(D, palette.divergent = gl.colors("div"), verbose = NULL, ...)
```

Arguments

D	Name of the distance matrix or class fd object [required].
palette.divergent	
	A divergent palette for the distance values [default gl.colors("div")].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity]
	Parameters passed to function heatmap.2 (package gplots)

Value

returns no value (i.e. NULL)

Author(s)

Custodian: Luis Mijangos – Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other graphics: gl.colors(), gl.map.interactive(), gl.report.ld.map(), gl.select.colors(),
gl.select.shapes(), gl.smearplot(), gl.tree.nj()
```

Examples

```
gl <- testset.gl[1:10,]
D <- dist(as.matrix(gl),upper=TRUE,diag=TRUE)
gl.plot.heatmap(D)
D2 <- gl.dist.pop(possums.gl)
gl.plot.heatmap(D2)
D3 <- gl.fixed.diff(testset.gl)
gl.plot.heatmap(D3)
```

```
if ((requireNamespace("gplots", quietly = TRUE))) {
D2 <- gl.dist.pop(possums.gl)
gl.plot.heatmap(D2)
}</pre>
```

gl.print.history Prints history of a genlight object

Description

Prints history of a genlight object

Usage

gl.print.history(x = NULL, history = NULL)

Arguments

х	A genlight object (with history) [optional].
history	Either a link to a history slot (gl\@other\$history), or a vector indicating which part of the history of x is used [c(1,3,4) uses the first, third and forth entry from x\@other\$history]. If no history is provided the complete history of x is used (recreating the identical object x) [optional].

Value

Prints a table with all history records. Currently the style cannot be changed.

Author(s)

Bernd Gruber (bugs? Post to https://groups.google.com/d/forum/dartr)

See Also

Other environment: gl.check.verbosity(), gl.check.wd(), gl.set.wd(), theme_dartR()

Examples

```
dartfile <- system.file('extdata','testset_SNPs_2Row.csv', package='dartR.data')
metadata <- system.file('extdata','testset_metadata.csv', package='dartR.data')
gl <- gl.read.dart(dartfile, ind.metafile = metadata, probar=FALSE)
gl2 <- gl.filter.callrate(gl, method='loc', threshold=0.9)
gl3 <- gl.filter.callrate(gl2, method='ind', threshold=0.95)
gl.print.history(gl3)</pre>
```

gl.prop.shared

Calculates a similarity (distance) matrix for individuals on the proportion of shared alleles @family distance

Description

This script calculates an individual based distance matrix. It uses an C++ implementation, so package Rcpp needs to be installed and it is therefore really fast (once it has compiled the function after the first run).

Usage

gl.propShared(x)

Arguments

Х

Name of the genlight containing the SNP genotypes [required].

Value

A similarity matrix

Author(s)

Bernd Gruber (Post to https://groups.google.com/d/forum/dartr)

Examples

#takes some time at the first run of the function...

res <- gl.propShared(bandicoot.gl)
res[1:5,1:7] #show only a small part of the matrix</pre>

gl.randomize.snps Randomly changes the allocation of 0's and 2's in a genlight object

Description

This function samples randomly half of the SNPs and re-codes, in the sampled SNP's, 0's by 2's. This function samples randomly half of the SNPs and re-codes, in the sampled SNP's, 0's by 2's.

Usage

```
gl.randomize.snps(
  х,
  plot.display = TRUE,
  plot.theme = theme_dartR(),
  plot.colors = NULL,
  plot.file = NULL,
  plot.dir = NULL,
  verbose = NULL
)
gl.randomize.snps(
  х,
  plot.display = TRUE,
  plot.theme = theme_dartR(),
  plot.colors = NULL,
  plot.file = NULL,
  plot.dir = NULL,
  verbose = NULL
)
```

Arguments

x	Name of the genlight object containing the SNP data [required].
plot.display	If TRUE, resultant plots are displayed in the plot window [default TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.colors	List of two color names for the borders and fill of the plots [default c("#2171B5", "#6BAED6")]
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL]
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log ; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity].

Details

DArT calls the most common allele as the reference allele. In a genlight object, homozygous for the reference allele are coded with a '0' and homozygous for the alternative allele are coded with a '2'. This causes some distortions in visuals from time to time. If plot.display = TRUE, two smear plots (pre-randomisation and post-randomisation) are presented using a random subset of individuals (10) and loci (100) to provide an overview of the changes. Resultant ggplots are saved to the session's temporary directory.

DArT calls the most common allele as the reference allele. In a genlight object, homozygous for the reference allele are coded with a '0' and homozygous for the alternative allele are coded with a '2'. This causes some distortions in visuals from time to time. If plot.display = TRUE, two smear plots (pre-randomisation and post-randomisation) are presented using a random subset of individuals (10)

gl.read.csv

and loci (100) to provide an overview of the changes. Resultant ggplots are saved to the session's temporary directory.

Value

Returns a genlight object with half of the loci re-coded.

Returns a genlight object with half of the loci re-coded.

Author(s)

Custodian: Luis Mijangos – Post to https://groups.google.com/d/forum/dartr

Examples

```
require("dartR.data")
res <- gl.randomize.snps(platypus.gl[1:5,1:5],verbose = 5)
gl <- gl.filter.monomorphs(testset.gl)
res <- gl.randomize.snps(gl,verbose = 5)</pre>
```

gl.read.csv

Reads SNP data from a csv file into a genlight object

Description

This script takes SNP genotypes from a csv file, combines them with individual and locus metrics and creates a genlight object. The SNP data need to be in one of two forms. SNPs can be coded 0 for homozygous reference, 2 for homozygous alternate, 1 for heterozygous, and NA for missing values; or the SNP data can be coded A/A, A/C, C/T, G/A etc, and -/- for missing data. In this format, the reference allele is the most frequent allele, as used by DArT. Other formats will throw an error. The SNP data need to be individuals as rows, labeled, and loci as columns, also labeled. If the orientation is individuals as columns and loci by rows, then set transpose=TRUE. The individual metrics need to be in a csv file, with headings, with a mandatory id column corresponding exactly to the individual identity labels provided with the SNP data and in the same order. The locus metadata needs to be in a csv file with headings, with a mandatory column headed AlleleID corresponding exactly to the locus identity labels provided with the SNP data and in the same order. Note that the locus metadata will be complemented by calculable statistics corresponding to those that would be provided by Diversity Arrays Technology (e.g. CallRate).

Usage

```
gl.read.csv(
   filename,
   transpose = FALSE,
   ind.metafile = NULL,
   loc.metafile = NULL,
   verbose = NULL
)
```

Arguments

filename	Name of the csv file containing the SNP genotypes [required].
transpose	If TRUE, rows are loci and columns are individuals [default FALSE].
ind.metafile	Name of the csv file containing the metrics for individuals [optional].
loc.metafile	Name of the csv file containing the metrics for loci [optional].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

A genlight object with the SNP data and associated metadata included.

Author(s)

Custodian: Luis Mijangos – Post to https://groups.google.com/d/forum/dartr

See Also

```
Other io: gl.load(), gl.read.dart(), gl.read.fasta(), gl.read.silicodart(), gl.save(),
gl.write.csv(), utils.read.dart()
```

gl.read.dart	Imports DArT data into dartR and converts it into a dartR genlight
	object

Description

This function is a wrapper function that allows you to convert your DArT file into a genlight object of class dartR.

Usage

```
gl.read.dart(
   filename,
   ind.metafile = NULL,
   recalc = TRUE,
   mono.rm = FALSE,
   nas = "-",
   topskip = NULL,
   lastmetric = NULL,
   covfilename = NULL,
   service.row = 1,
   plate.row = 3,
   probar = FALSE,
   verbose = NULL
)
```

gl.read.dart

Arguments

filename	File containing the SNP data (csv file) [required].
ind.metafile	File that contains additional information on individuals [required].
recalc	If TRUE, force the recalculation of locus metrics [default TRUE].
mono.rm	If TRUE, force the removal of monomorphic loci (including all NAs. [default FALSE].
nas	A character specifying NAs [default '-'].
topskip	A number specifying the number of initial rows to be skipped [default NULL].
lastmetric	Deprecated, specifies the last column of locus metadata. Can be specified as a column number [default NULL].
covfilename	Deprecated, sse ind.metafile parameter [NULL].
service.row	The row number for the DArT service is contained [default 1].
plate.row	The row number the plate well [default 3].
probar	Show progress bar [default FALSE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, or as set by gl.set.verbose()].

Details

The function will determine automatically if the data are in Diversity Arrays one-row csv format or two-row csv format. The first row of data is determined from the number of rows with an * in the first column. This can be alternatively specified with the topskip parameter. The DArT service code is added to the ind.metrics of the genlight object. The row containing the service code for each individual can be specified with the service.row parameter. #'The DArT plate well is added to the ind.metrics of the genlight object. The row containing the plate well for each individual can be specified with the plate.row parameter. If individuals have been deleted from the input file manually, then the locus metrics supplied by DArT will no longer be correct and some loci may be monomorphic. To accommodate this, set mono.rm and recalc to TRUE.

Value

A dartR genlight object that contains individual and locus metrics [if data were provided] and locus metrics [from a DArT report].

Author(s)

Custodian: Bernd Gruber (Post to https://groups.google.com/d/forum/dartr)

See Also

```
utils.read.dart
```

```
Other io: gl.load(), gl.read.csv(), gl.read.fasta(), gl.read.silicodart(), gl.save(),
gl.write.csv(), utils.read.dart()
```

Examples

```
dartfile <- system.file('extdata','testset_SNPs_2Row.csv', package='dartR.data')
metadata <- system.file('extdata','testset_metadata.csv', package='dartR.data')
gl <- gl.read.dart(dartfile, ind.metafile = metadata, probar=TRUE)</pre>
```

gl.read.fasta Read

Reads FASTA files and converts them to genlight object

Description

The following IUPAC Ambiguity Codes are taken as heterozygotes:

- M is heterozygote for AC and CA
- R is heterozygote for AG and GA
- W is heterozygote for AT and TA
- S is heterozygote for CG and GC
- Y is heterozygote for CT and TC
- K is heterozygote for GT and TG

The following IUPAC Ambiguity Codes are taken as missing data:

- V
- H
- D
- B
- N

The function can deal with missing data in individuals, e.g. when FASTA files have different number of individuals due to missing data. The allele with the highest frequency is taken as the reference allele. SNPs with more than two alleles are skipped.

Usage

```
gl.read.fasta(fasta.files, parallel = FALSE, n.cores = NULL, verbose = NULL)
```

Arguments

fasta.files	Fasta files to read [required].
parallel	A logical indicating whether multiple cores -if available- should be used for the computations (TRUE), or not (FALSE); requires the package parallel to be installed [default FALSE].
n.cores	If parallel is TRUE, the number of cores to be used in the computations; if NULL, then the maximum number of cores available on the computer is used [default NULL].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

gl.read.PLINK

Details

Ambiguity characters are often used to code heterozygotes. However, using heterozygotes as ambiguity characters may bias many estimates. See more information in the link below: https://evodify.com/heterozygotes-ambiguity-characters/

Value

A genlight object.

Author(s)

Custodian: Luis Mijangos – Post to https://groups.google.com/d/forum/dartr

See Also

```
Other io: gl.load(), gl.read.csv(), gl.read.dart(), gl.read.silicodart(), gl.save(),
gl.write.csv(), utils.read.dart()
```

gl.read.PLINK

Reads PLINK data file into a genlight object

Description

This function imports PLINK data into a genlight object and append available metadata.

Usage

```
gl.read.PLINK(
   filename,
   ind.metafile = NULL,
   loc.metafile = NULL,
   plink.cmd = "plink",
   plink.path = "path",
   plink.flags = NULL,
   verbose = NULL
)
```

Arguments

filename	Fully qualified path to PLINK input file (without including the extension)
ind.metafile	Name of the csv file containing the metrics for individuals [optional].
loc.metafile	Name of the csv file containing the metrics for loci [optional].
plink.cmd	The 'name' to call plink. This will depend on the file name (without the extension '.exe' if on windows) or the name of the PATH variable
plink.path	The path where the executable is. If plink is listed in the PATH then there is no need for this. This is what the option "path" means

plink.flags	additional possible parameters passed on to plink.
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress
	and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Details

This function handles .ped or .bed file (with the associate files - e.g. .fam, .bim). However, if a .ped file is provided, PLINK needs to be installed and it is used to convert the .ped into a .bed, which is then converted into a genlight.

Additional metadata can be included passing .csv files. These will be appended to the existing metadata present in the PLINK files.

The locus metadata needs to be in a csv file with headings, with a mandatory column headed AlleleID corresponding exactly to the locus identity labels provided with the SNP data

Value

A genlight object with the SNP data and associated metadata included.

Author(s)

Custodian: Carlo Pacioni – Post to https://groups.google.com/d/forum/dartr

gl.read.silicodart	Imports presence/absence data from SilicoDArT to genlight {age-
	genet} format (ploidy=1)

Description

DaRT provide the data as a matrix of entities (individual animals) across the top and attributes (P/A of sequenced fragment) down the side in a format that is unique to DArT. This program reads the data in to adegenet format for consistency with other programming activity. The script may require modification as DArT modify their data formats from time to time.

Usage

```
gl.read.silicodart(
  filename,
  ind.metafile = NULL,
  nas = "-",
  topskip = NULL,
  lastmetric = "Reproducibility",
  probar = TRUE,
  verbose = NULL
)
```

gl.read.silicodart

Arguments

filename	Name of csv file containing the SilicoDArT data [required].
ind.metafile	Name of csv file containing metadata assigned to each entity (individual) [de-fault NULL].
nas	Missing data character [default '-'].
topskip	Number of rows to skip before the header row (containing the specimen identi- ties) [optional].
lastmetric	Specifies the last non genetic column (Default is 'Reproducibility'). Be sure to check if that is true, otherwise the number of individuals will not match. You can also specify the last column by a number [default "Reproducibility"].
probar	Show progress bar [default TRUE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, or as set by gl.set.verbose()].

Details

gl.read.silicodart() opens the data file (csv comma delimited) and skips the first n=topskip lines. The script assumes that the next line contains the entity labels (specimen ids) followed immediately by the SNP data for the first locus. It reads the presence/absence data into a matrix of 1s and 0s, and inputs the locus metadata and specimen metadata. The locus metadata comprises a series of columns of values for each locus including the essential columns of CloneID and the desirable variables Reproducibility and PIC. Refer to documentation provide by DArT for an explanation of these columns. The specimen metadata provides the opportunity to reassign specimens to populations, and to add other data relevant to the specimen. The key variables are id (specimen identity which must be the same and in the same order as the SilicoDArT file, each unique), pop (population assignment), lat (latitude, optional) and lon (longitude, optional). id, pop, lat, lon are the column headers in the csv file. Other optional columns can be added. The data matrix, locus names (forced to be unique), locus metadata, specimen names, specimen metadata are combined into a genind object. Refer to the documentation for {adegenet} for further details.

Value

An object of class genlight with ploidy set to 1, containing the presence/absence data, and locus and individual metadata.

Author(s)

Custodian: Bernd Gruber - Post to https://groups.google.com/d/forum/dartr

See Also

```
gl.read.dart
```

Other io: gl.load(), gl.read.csv(), gl.read.dart(), gl.read.fasta(), gl.save(), gl.write.csv(), utils.read.dart()

Examples

```
silicodartfile <- system.file('extdata','testset_SilicoDArT.csv', package='dartR.data')
metadata <- system.file('extdata',ind.metafile ='testset_metadata_silicodart.csv',
package='dartR.data')
testset.gs <- gl.read.silicodart(filename = silicodartfile, ind.metafile = metadata)</pre>
```

gl.read.vcf

Converts a vcf file into a genlight object

Description

This function needs package vcfR, please install it.

Usage

```
gl.read.vcf(vcffile, ind.metafile = NULL, mode = "genotype", verbose = NULL)
```

Arguments

vcffile	A vcf file (works only for diploid data) [required].
ind.metafile	Optional file in csv format with metadata for each individual (see details for explanation) [default NULL].
mode	"genotype" all heterozygous sites will be coded as 1 regardless ploidy level, dosage: sites will be codes as copy number of alternate allele [default genotype]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Details

The ind.metadata file needs to have very specific headings. First a heading called id. Here the ids have to match the ids in the dartR object. The following column headings are optional. pop: specifies the population membership of each individual. lat and lon specify spatial coordinates (in decimal degrees WGS1984 format). Additional columns with individual metadata can be imported (e.g. age, gender). Note also that this function checks to see if there are input of mode, missing input of mode will issue the user with an error. "Dosage" mode of this function assign ploidy levels as maximum copy number of alternate alleles. Please carefully check the data if "dosage" mode is used.

Value

A genlight object.

Author(s)

Bernd Gruber, Ching Lau (Post to https://groups.google.com/d/forum/dartr)

gl.reassign.pop

Examples

End(Not run)

gl.reassign.pop Assigns an individual metric as pop in a genlight {adegenet} object

Description

Individuals are assigned to populations based on the individual/sample/specimen metrics file (csv) used with gl.read.dart(). One might want to define the population structure in accordance with another classification, such as using an individual metric (e.g. sex, male or female). This script discards the current population assignments and replaces them with new population assignments defined by a specified individual metric. The function returns a genlight object with the new population assignments. Note that the original population assignments are lost.

Usage

gl.reassign.pop(x, as.pop, verbose = NULL)

Arguments

х	Name of the genlight object containing SNP genotypes [required].
as.pop	Specify the name of the individual metric to set as the pop variable [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

A genlight object with the reassigned populations.

Author(s)

Custodian: Arthur Georges - Post to https://groups.google.com/d/forum/dartr

See Also

Other data manipulation: gl.define.pop(), gl.drop.ind(), gl.drop.loc(), gl.drop.pop(), gl.edit.recode.pop(), gl.impute(), gl.join(), gl.keep.ind(), gl.keep.loc(), gl.keep.pop(), gl.make.recode.ind(), gl.merge.pop(), gl.recode.ind(), gl.recode.pop(), gl.rename.pop(), gl.sample(), gl.sim.genotypes(), gl.sort(), gl.subsample.ind(), gl.subsample.loc()

Examples

```
# SNP data
popNames(testset.gl)
gl <- gl.reassign.pop(testset.gl, as.pop='sex',verbose=3)
popNames(gl)
# Tag P/A data
popNames(testset.gs)
gs <- gl.reassign.pop(testset.gs, as.pop='sex',verbose=3)
popNames(gs)</pre>
```

gl.recalc.metrics	Recalculates	locus	metrics	when	individuals	or	populations	are
	deleted from a	a genli	ght {adeg	enet} o	object @fami	ily e	nvironment	

Description

When individuals, or populations, are deleted from a genlight object, the locus metrics no longer apply. For example, the Call Rate may be different considering the subset of individuals, compared with the full set. This script recalculates those affected locus metrics, namely, avgPIC, CallRate, freqHets, freqHomRef, freqHomSnp, OneRatioRef, OneRatioSnp, PICRef and PICSnp. Metrics that remain unaltered are RepAvg and TrimmedSeq as they are unaffected by the removal of individuals. The script optionally removes resultant monomorphic loci or loci with all values missing and deletes them (using gl.filter.monomorphs.r). The script returns a genlight object with the recalculated locus metadata.

Usage

gl.recalc.metrics(x, mono.rm = FALSE, verbose = NULL)

Arguments

х	Name of the genlight object containing SNP genotypes [required].
mono.rm	If TRUE, removes monomorphic loci [default FALSE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress
	and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

A genlight object with the recalculated locus metadata.

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

See Also

gl.filter.monomorphs

gl.recode.ind

Examples

gl <- gl.recalc.metrics(testset.gl, verbose=2)</pre>

gl.recode.ind Recodes individual (=specimen = sample) labels in a genlight object

Description

This function recodes individual labels and/or deletes individuals from a DaRT genlight SNP file based on a lookup table provided as a csv file.

Usage

```
gl.recode.ind(x, ind.recode, recalc = FALSE, mono.rm = FALSE, verbose = NULL)
```

Arguments

х	Name of the genlight object [required].
ind.recode	Name of the csv file containing the individual relabelling [required].
recalc	If TRUE, recalculate the locus metadata statistics if any individuals are deleted in the filtering [default FALSE].
mono.rm	If TRUE, remove monomorphic loci [default FALSE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Details

Renaming individuals may be required when there have been errors in labeling arising in the process from sample to sequence files. There may be occasions where renaming individuals is required for preparation of figures. When caution needs to be exercised because of the potential for breaking the 'chain of evidence' associated with the samples, recoding individuals using a recode table (csv) can provide a durable record of the changes. The function works with genlight objects containing SNP genotypes and Tag P/A data (SilicoDArT). For SNP genotype data, the function, having deleted individuals, optionally identifies resultant monomorphic loci or loci with all values missing and deletes them. The script also optionally recalculates the locus metadata as appropriate. The optional deletion of monomorphic loci and the optional recalculation of locus statistics is not available for Tag P/A data (SilicoDArT). The script returns a dartR genlight object with the new individual names and the recalculated locus metadata.

Value

A genlight or genind object with the recoded and reduced data.

Author(s)

Custodian: Arthur Georges - Post to https://groups.google.com/d/forum/dartr

See Also

gl.filter.monomorphs for filtering monomorphs, gl.recalc.metrics for recalculating locus metrics, gl.recode.pop for recoding populations

Other data manipulation: gl.define.pop(), gl.drop.ind(), gl.drop.loc(), gl.drop.pop(), gl.edit.recode.pop(), gl.impute(), gl.join(), gl.keep.ind(), gl.keep.loc(), gl.keep.pop(), gl.make.recode.ind(), gl.merge.pop(), gl.reassign.pop(), gl.recode.pop(), gl.rename.pop(), gl.sample(), gl.sim.genotypes(), gl.sort(), gl.subsample.ind(), gl.subsample.loc()

Examples

```
file <- system.file('extdata','testset_ind_recode.csv', package='dartR.data')
gl <- gl.recode.ind(testset.gl, ind.recode=file, verbose=3)</pre>
```

gl.recode.pop Recodes population assignments in a genlight object

Description

This function recodes population assignments and/or deletes populations from a DaRT genlight object based on information provided in a csv population recode file.

Usage

```
gl.recode.pop(x, pop.recode, recalc = FALSE, mono.rm = FALSE, verbose = NULL)
```

Arguments

Х	Name of the genlight object [required].
pop.recode	Name of the csv file containing the population reassignments [required].
recalc	If TRUE, recalculates the locus metadata statistics if any individuals are deleted in the filtering [default FALSE].
mono.rm	If TRUE, removes monomorphic loci [default FALSE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Details

Individuals are assigned to populations based on the specimen metadata data file (csv) used with gl.read.dart(). Recoding can be used to amalgamate populations or to selectively delete or retain populations. When caution needs to be exercised because of the potential for breaking the 'chain of evidence' associated with the samples, recoding individuals using a recode table (csv) can provide a durable record of the changes. The population recode file contains a list of populations taken from the genlight object as the first column of the csv file, and the new population assignments are located in the second column of the csv file. The keyword 'Delete' used as a new population assignment will result in the associated specimen being dropped from the dataset. The function works with genlight

gl.rename.pop

objects containing SNP genotypes and Tag P/A data (SilicoDArT). For SNP genotype data, the function, having deleted populations, optionally identifies resultant monomorphic loci or loci with all values missing and deletes them. The script also optionally recalculates the locus metadata as appropriate. The optional deletion of monomorphic loci and the optional recalculation of locus statistics is not available for Tag P/A data (SilicoDArT).

Value

A genlight object with the recoded and reduced data.

Author(s)

Custodian: Arthur Georges - Post to https://groups.google.com/d/forum/dartr

See Also

- gl.filter.monomorphs
- gl.recode.pop

```
Other data manipulation: gl.define.pop(), gl.drop.ind(), gl.drop.loc(), gl.drop.pop(),
gl.edit.recode.pop(), gl.impute(), gl.join(), gl.keep.ind(), gl.keep.loc(), gl.keep.pop(),
gl.make.recode.ind(), gl.merge.pop(), gl.reassign.pop(), gl.recode.ind(), gl.rename.pop(),
gl.sample(), gl.sim.genotypes(), gl.sort(), gl.subsample.ind(), gl.subsample.loc()
```

Examples

mfile <- system.file('extdata', 'testset_pop_recode.csv', package='dartR.data')
nPop(testset.gl)
gl <- gl.recode.pop(testset.gl, pop.recode=mfile, verbose=3)</pre>

gl.rename.pop

Renames a population in a genlight object

Description

Individuals are assigned to populations based on the specimen metadata data file (csv) used with gl.read.dart(). This script renames a nominated population. The script returns a genlight object with the new population name.

Usage

```
gl.rename.pop(x, old = NULL, new = NULL, verbose = NULL)
```

Arguments

x	Name of the genlight object containing SNP genotypes [required].
old	Name of population to be changed [required].
new	New name for the population [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

A genlight object with the new population name.

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

```
Other data manipulation: gl.define.pop(), gl.drop.ind(), gl.drop.loc(), gl.drop.pop(),
gl.edit.recode.pop(), gl.impute(), gl.join(), gl.keep.ind(), gl.keep.loc(), gl.keep.pop(),
gl.make.recode.ind(), gl.merge.pop(), gl.reassign.pop(), gl.recode.ind(), gl.recode.pop(),
gl.sample(), gl.sim.genotypes(), gl.sort(), gl.subsample.ind(), gl.subsample.loc()
```

Examples

gl <- gl.rename.pop(testset.gl, old='EmsubRopeMata', new='Outgroup')</pre>

gl.report.allelerich Report allelic richness per population from a genlight object

Description

This function needs package adegenet, please install it.

Usage

```
gl.report.allelerich(
    x,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.dir = NULL,
    plot.file = NULL,
    error.bar = "SD",
    verbose = 2
)
```

gl.report.allelerich

Arguments

x	A genlight file (works only for diploid data) [required].
plot.display	Specify if plot is to be produced [default TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()].
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL].
error.bar	Statistic to be plotted as error bar either "SD" (standard deviation) or "SE" (stan- dard error) or "CI" (confidence intervals) [default "SD"].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Details

details

- raw allele count
- allelic richness

Value

A dataframe containing richness per site, richness per population, raw reference allele count, raw alternate allele count.

Author(s)

Ching Ching Lau (Post to https://groups.google.com/d/forum/dartr)

References

• El Mousadik, A., & Petit, R. J. (1996). High level of genetic differentiation for allelic richness among populations of the argan tree [Argania spinosa (L.) Skeels] endemic to Morocco. Theoretical and applied genetics, 92, 832-839.

Examples

gl.report.allelerich(possums.gl)

gl.report.allna

Reports loci that are all NA across individuals and/or populations with all NA across loci

Description

This script reports loci or individuals with all calls missing (NA), from a genlight object.

A DArT dataset will not have loci for which the calls are scored all as missing (NA) for a particular individual, but such loci can arise rarely when populations or individuals are deleted. Similarly, a DArT dataset will not have individuals for which the calls are scored all as missing (NA) across all loci, but such individuals may sneak in to the dataset when loci are deleted. Retaining individual or loci with all NAs can cause issues for several functions.

Also, on occasions an analysis will require that there are some loci scored in each population. Setting by.pop=TRUE will result in removal of loci when they are all missing in any one population.

Note that loci that are missing for all individuals in a population are not imputed with method 'frequency' or 'HW'. Consider using the function gl.filter.allna with by.pop=TRUE.

Usage

gl.report.allna(x, by.pop = FALSE, verbose = NULL)

Arguments

х	Name of the input genlight object [required].
by.pop	If TRUE, loci that are all missing in any one population are reported [default FALSE]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Value

gl.report.allna

Author(s)

Author(s): Arthur Georges. Custodian: Arthur Georges – Post to https://groups.google.com/ d/forum/dartr

See Also

```
Other matched report: gl.filter.excess.het(), gl.report.callrate(), gl.report.hamming(),
gl.report.locmetric(), gl.report.maf(), gl.report.overshoot(), gl.report.pa(), gl.report.rdepth(),
gl.report.reproducibility(), gl.report.secondaries(), gl.report.taglength()
```

Other filter functions: gl.filter.allna(), gl.filter.hwe()

gl.report.bases

Examples

```
# SNP data
  result <- gl.report.allna(testset.gl, verbose=3)
# Tag P/A data
  result <- gl.report.allna(testset.gs, verbose=3)</pre>
```

gl.report.bases Reports summary of base pair frequencies

Description

Calculates the frequencies of the four DNA nucleotide bases: adenine (A), cytosine (C), 'guanine (G) and thymine (T), and the frequency of transitions (Ts) and transversions (Tv) in a DArT genlight object.

Usage

```
gl.report.bases(
    x,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
    plot.file = NULL,
    verbose = NULL,
    ...
)
```

Arguments

x	Name of the genlight object containing the SNP or presence/absence (Silico-DArT) data [required].
plot.display	If TRUE, resultant plots are displayed in the plot window [default TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.colors	List of two color names for the borders and fill of the plots [default c("#2171B5", "#6BAED6")]
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL].
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity].
	Parameters passed to function ggsave, such as width and height, when the ggplot is to be saved.

Details

The function checks first if trimmed sequences are included in the locus metadata (@other\$loc.metrics\$TrimmedSequence), and if so, tallies up the numbers of A, T, G and C bases. Only the reference state at the SNP locus is counted. Counts of transitions (Ts) and transversions (Tv) assume that there is no directionality, that is C->T is the same as T->C, because the reference state is arbitrary. For presence/absence data (SilicoDArT), it is not possible to count transversions or transitions or transversions/transitions ratio because the SNP data are not available, only a single sequence tag per locus. Only base frequencies are provided. A color vector can be obtained with gl.select.colors() and then passed to the function with the plot.colors parameter. If a plot.file is given, the ggplot arising from this function is saved as an "RDS" binary file using saveRDS(); can be reloaded with readRDS(). A file name must be specified for the plot to be saved. If a plot directory (plot.dir) is specified, the ggplot binary is saved to that directory; otherwise to the tempdir(). To avoid issues from inadvertent use of this function in an assignment statement, the function returns the genlight object unaltered.

Value

The unchanged genlight object

Author(s)

Author(s); Arthur Georges. Custodian: Arthur Georges – Post to https://groups.google.com/ d/forum/dartr

See Also

Other matched reports: gl.mahal.assign(), gl.report.factorloadings(), gl.report.fstat(), gl.report.monomorphs()

Examples

```
# SNP data
    out <- gl.report.bases(testset.gl)
    col <- gl.select.colors(select=c(6,1),palette=rainbow, verbose=0)
    out <- gl.report.bases(testset.gl,plot.colors=col)
# Tag P/A data
    out <- gl.report.bases(testset.gs)</pre>
```

gl.report.basics Basic statistics for a genlight object

Description

Calculates basic statistics for a genlight object.

Usage

gl.report.basics(x, verbose = NULL)

Arguments

x	Name of the genlight object [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Author(s)

Custodian: Arthur Georges (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other unmatched report: gl.allele.freq(), gl.report.diversity(), gl.report.excess.het(),
gl.report.heterozygosity(), gl.report.polyploid_heterozygosity()
```

Examples

```
gl.report.basics(platypus.gl)
```

gl.report.callrate Reports summary of Call Rate for loci or individuals

Description

SNP datasets generated by DArT have missing values primarily arising from failure to call a SNP because of a mutation at one or both of the restriction enzyme recognition sites. P/A datasets (SilicoDArT) have missing values because it was not possible to call whether a sequence tag was amplified or not.

Usage

```
gl.report.callrate(
    x,
    method = "loc",
    ind.to.list = 20,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
    plot.dir = NULL,
    plot.file = NULL,
    bins = 50,
    verbose = NULL,
    ...
)
```

Arguments

x	Name of the genlight object containing the SNP or presence/absence (Silico-DArT) data [required].
method	Specify the type of report by locus (method='loc') or individual (method='ind') [default 'loc'].
ind.to.list	Number of individuals to list for callrate [default 20]
plot.display	Specify if plot is to be displayed in the graphics window [default TRUE].
plot.theme	User specified theme [default theme_dartR()].
plot.colors	Vector with two color names for the borders and fill [default c("#2171B5", "#6BAED6")].
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]
plot.file	Filename (minus extension) for the RDS plot file [Required for plot save]
bins	Number of bins to display in histograms [default 25].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].
	Parameters passed to function ggsave, such as width and height, when the ggplot is to be saved.

Details

This function expects a genlight object, containing either SNP data or SilicoDArT (=presence/absence data).

Callrate is summarized by locus or by individual to allow sensible decisions on thresholds for filtering taking into consideration consequential loss of data. The summary is in the form of a tabulation and plots.

The table of quantiles is useful for deciding a threshold for subsequent filtering as it provides an indication of the percentages of loci that will be retained and lost.

In the case of method='ind', a list of individuals to be deleted is provided. To manage the screen output, this list is limited to ind.to.list individuals (or nInd(x)) whichever is the smaller.

To avoid issues from inadvertent use of this function in an assignment statement, the function returns the genlight object unaltered.

A color vector can be obtained with gl.select.colors() and then passed to the function with the plot.colors parameter.

If a plot.file is given, the ggplot arising from this function is saved as an "RDS" binary file using saveRDS(); can be reloaded with readRDS(). A file name must be specified for the plot to be saved.

If a plot directory (plot.dir) is specified, the ggplot binary is saved to that directory; otherwise to the tempdir().

Value

Returns unaltered genlight object

gl.report.diversity

Author(s)

Author(s): Arthur Georges. Custodian: Arthur Georges – Post to https://groups.google.com/ d/forum/dartr

See Also

gl.filter.callrate

```
Other matched report: gl.filter.excess.het(), gl.report.allna(), gl.report.hamming(),
gl.report.locmetric(), gl.report.maf(), gl.report.overshoot(), gl.report.pa(), gl.report.rdepth(),
gl.report.reproducibility(), gl.report.secondaries(), gl.report.taglength()
```

Examples

```
# SNP data
  test.gl <- testset.gl[1:20,]
  gl.report.callrate(test.gl)
  gl.report.callrate(test.gl,method='ind')
  gl.report.callrate(test.gl,method='ind',plot.file="test")
  gl.report.callrate(test.gl,method='loc',by.pop=TRUE)
  gl.report.callrate(test.gl,method='loc',by.pop=TRUE,plot.file="test")
# Tag P/A data
  test.gs <- testset.gs[1:20,]
  gl.report.callrate(test.gs,method='ind')
  test.gl <- testset.gl[1:20,]
  gl.report.callrate(test.gl)</pre>
```

gl.report.diversity Calculates diversity indexes for SNPs

Description

This script takes a genlight object and calculates alpha and beta diversity for q = 0.2. Formulas are taken from Sherwin et al. 2017. The paper describes nicely the relationship between the different q levels and how they relate to population genetic processes such as dispersal and selection. The citation below also includes a link to a 3-minute video that explains, q, D and H.

Usage

```
gl.report.diversity(
    x,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors.pop = gl.colors("dis"),
    plot.dir = NULL,
    plot.file = NULL,
```

```
table = "DH",
verbose = NULL
)
```

Arguments

x	Name of the genlight object containing the SNP or presence/absence (Silico-DArT) data [required].
plot.display	Specify if plot is to be displayed in the graphics window [default TRUE].
plot.theme	User specified theme [default theme_dartR()].
<pre>plot.colors.pop</pre>	
	A color palette for population plots or a list with as many colors as there are populations in the dataset [default gl.colors("dis")].
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()].
plot.file	If TRUE, saves any ggplots and listings to the session temporary directory (tem- pdir) [default FALSE].
table	Prints a tabular output to the console either 'D'=D values, or 'H'=H values or 'DH','HD'=both or 'N'=no table. [default 'DH'].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity].

Details

For all indexes, the entropies (H) and corresponding effective numbers, i.e. Hill numbers (D), which reflect the number of needed entities to get the observed values, are calculated. In a nutshell, the alpha indexes between the different q-values should be similar if there is no deviation from expected allele frequencies and occurrences (e.g. all loci in HWE & equilibrium). If there is a deviation of an index, this links to a process causing it, such as dispersal, selection or strong drift. For a detailed explanation of all the indexes, we recommend resorting to the literature provided below. Error bars are +/- 1 standard deviation.

Function's output

If the function's parameter "table" = "DH" (the default value) is used, the output of the function is 20 tables.

The first two show the number of loci used. The name of each of the rest of the tables starts with three terms separated by underscores.

The first term refers to the q value (0 to 2). The q values identify different ways of summarising diversity (H): q=0 is simply the number of alleles per locus, with no information about their relative proportions; q=2 is the expected heterozygosity, ie the chance of drawing two different alleles at random from the population; q=1 is the Shannon measure of 'surprise, relating to how likely it is that the next allele drawn will be one that has not been seen before (Sherwin et al 2017, 2021, and associated video).

The second term refers to whether it is the diversity measure (H) or its transformation to Hill numbers (D) The D value tells you how many equally-frequent alleles there would need to be to give the corresponding H-value (in the actual population) The D-values are all in units of numbers of
gl.report.diversity

alleles, so they can be plotted against the q-value to get a rich representation of the diversity (Box 1, Fig II in Sherwin et al 2017, 2021, and associated video).

The third term refers to whether the diversity is calculated within populations (alpha) or between populations (beta).

In the case of alpha diversity tables, standard deviations have their own table, which finishes with a fourth term: "sd".

In the case of beta diversity tables, standard deviations are in the upper triangle of the matrix and diversity values are in the lower triangle of the matrix.

Plotting

Plot colours can be set with gl.select.colors().

If plot.file is specified, plots are saved to the directory specified by the user, or the global default working directory set by gl.set.wd() or to the tempdir().

Examples of other themes that can be used can be consulted in

- https://ggplot2.tidyverse.org/reference/ggtheme.html and
- https://yutannihilation.github.io/allYourFigureAreBelongToUs/ggthemes/

Value

A list of entropy indexes for each level of q and equivalent numbers for alpha and beta diversity.

Author(s)

Bernd Gruber (Post to https://groups.google.com/d/forum/dartr), Contributors: William B. Sherwin, Alexander Sentinella

References

Sherwin, W.B., Chao, A., Johst, L., Smouse, P.E. (2017, 2021). Information Theory Broadens the Spectrum of Molecular Ecology and Evolution. TREE 32(12) 948-963. doi:10.1016/j.tree.2017.09.12 AND TREE 36:955-6 doi.org/10.1016/j.tree.2021.07.005 AND 3-Minute video: ars.els-cdn.com/content/image/1-s2.0-S0169534717302550-mmc2.mp4

See Also

Other unmatched report: gl.allele.freq(), gl.report.basics(), gl.report.excess.het(), gl.report.heterozygosity(), gl.report.polyploid_heterozygosity()

Examples

```
div <- gl.report.diversity(bandicoot.gl, table=FALSE)
div$zero_H_alpha
div$two_H_beta
names(div)</pre>
```

gl.report.excess.het Report loci with excess of heterozygosity

Description

Calculates excess of heterozygosity in a genlight object

Usage

```
gl.report.excess.het(
    x,
    Yates = FALSE,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
    plot.file = NULL,
    plot.dir = NULL,
    verbose = NULL
)
```

Arguments

x	Name of the genlight object containing the SNP data [required].
Yates	Boolean for Yates's continuity correction. [default FALSE]
plot.display	Specify if plot is to be produced [default TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.colors	Vector with two color names for the borders and fill [default c("#2171B5", "#6BAED6")].
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL].
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity].

Details

Loci with observed heterozygosity larger than 0.5 and expected heterozygosity would be indicated as excess (p-value <= 0.05). You can remove the loci with excess of heterozygosity from genlight object using gl.filter.excess.het

Function's output

If a plot.file is given, the ggplot arising from this function is saved as an "RDS" binary file using saveRDS(); can be reloaded with readRDS(). A file name must be specified for the plot to be saved.

If a plot directory (plot.dir) is specified, the ggplot binary is saved to that directory; otherwise to the tempdir().

Examples of other themes that can be used can be consulted in:

- https://ggplot2.tidyverse.org/reference/ggtheme.html and
- https://yutannihilation.github.io/allYourFigureAreBelongToUs/ggthemes/

A color vector can be obtained with gl.select.colors() and then passed to the function with the plot.colors parameter.

Value

1. Table with information of excessively-heterozygous loci

2. Two plots of heterozygosity of the loci before and after filtering (i.e. without excessively heterozygous loci).

3. A vector with the names of loci to be remove by gl.filter.excess.het

Author(s)

Jesús Castrejón-Figueroa, Diana A Robledo-Ruiz (Custodian: Ching Ching Lau) – Post to https: //groups.google.com/d/forum/dartr

References

- https://github.com/drobledoruiz/conservation_genomics/tree/main/filter.excess.het
- Robledo-Ruiz, D. A., Austin, L., Amos, J. N., Castrejón-Figueroa, J., Harley, D. K., Magrath, M. J., Sunnucks, P. & Pavlova, A. (2023). Easy-to-use R functions to separate reducedrepresentation genomic datasets into sex-linked and autosomal loci, and conduct sex assignment. Molecular Ecology Resources.

See Also

gl.filter.excess.het

Other unmatched report: gl.allele.freq(), gl.report.basics(), gl.report.diversity(), gl.report.heterozygosity(), gl.report.polyploid_heterozygosity()

Examples

filtered.table <- gl.report.excess.het(x = LBP, Yates = TRUE)</pre>

```
gl.report.factorloadings
```

Reports factor loadings for a PCA or PCoA

Description

Extracts the factor loadings from a glPCA object (generated by gl.pcoa) and plots their distribuion.

Usage

```
gl.report.factorloadings(
    pca,
    axis = 1,
    n.display = 15,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
    plot.file = NULL,
    plot.dir = NULL,
    bins = 25,
    verbose = NULL,
    ...
)
```

Arguments

рса	Name of the glPCA object containing factor loadings [required].
axis	Axis in the ordination used to display the factor loadings [default 1]
n.display	Number of loci for which to display factorloadings [default 15]
plot.display	If TRUE, resultant plots are displayed in the plot window [default TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.colors	List of two color names for the borders and fill of the plots [default c("#2171B5","#6BAED6")]
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL]
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]
bins	Number of bins to display in histograms [default 25].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity]
	Parameters passed to function ggsave, such as width and height, when the ggplot is to be saved.

gl.report.fstat

Details

The function extracts the factor loadings for a given axis from a PCA object generated by gl.pcoa and plots their magnitudes. Useful for identifying loci that load high for a given axis.

A color vector can be obtained with gl.select.colors() and then passed to the function with the plot.colors parameter.

Themes can be obtained from in

- https://ggplot2.tidyverse.org/reference/ggtheme.html and
- https://yutannihilation.github.io/allYourFigureAreBelongToUs/ggthemes/

If a plot.file is given, the ggplot arising from this function is saved as an "RDS" binary file using saveRDS(); can be reloaded with readRDS(). A file name must be specified for the plot to be saved. If a plot directory (plot.dir) is specified, the ggplot binary is saved to that directory; otherwise to the tempdir().

Value

The unchanged genlight object

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

Other matched reports: gl.mahal.assign(), gl.report.bases(), gl.report.fstat(), gl.report.monomorphs()

Examples

```
pca <- gl.pcoa(testset.gl)
gl.report.factorloadings(pca = pca)</pre>
```

gl.report.fstat Reports various statistics of genetic differentiation between populations with confidence intervals

Description

This function calculates four genetic differentiation between populations statistics (see the "Details" section for further information).

- Fst Measure of the degree of genetic differentiation of subpopulations (Nei, 1987).
- Fstp Unbiased (i.e. corrected for sampling error, see explanation below) Fst (Nei, 1987).
- Dest Jost's D (Jost, 2008).
- **Gst_H** Gst standardized by the maximum level that it can obtain for the observed amount of genetic variation (Hedrick 2005).

Sampling errors arise because allele frequencies in our samples differ from those in the subpopulations from which they were taken (Holsinger, 2012).

Confidence Intervals are obtained using bootstrapping.

Usage

```
gl.report.fstat(
    x,
    nboots = 0,
    conf = 0.95,
    CI.type = "bca",
    ncpus = 1,
    plot.stat = "Fstp",
    plot.display = TRUE,
    palette.divergent = gl.colors("div"),
    font.size = 0.5,
    plot.dir = NULL,
    plot.file = NULL,
    verbose = NULL,
    ...
)
```

Arguments

x	Name of the genlight object containing the SNP data [required].	
nboots	Number of bootstrap replicates to obtain confidence intervals [default 0].	
conf	The confidence level of the required interval [default 0.95].	
CI.type	Method to estimate confidence intervals. One of "norm", "basic", "perc" or "bca" [default "bca"].	
ncpus	Number of processes to be used in parallel operation. If ncpus > 1 parallel operation is activated, see "Details" section [default 1].	
plot.stat	Statistic to plot. One of "Fst", "Fstp", "Dest" or "Gst_H" [default "Fstp"].	
plot.display	If TRUE, a heatmap of the pairwise static chosen is displayed in the plot window [default TRUE].	
palette.divergent		
	A color palette function for the heatmap plot [default gl.colors("div")].	
font.size	Size of font for the labels of horizontal and vertical axes of the heatmap [default 0.5].	
plot.dir	Directory in which to save files [default working directory].	
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL].	
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity]	
	Parameters passed to function heatmap.2 (package gplots).	

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gl.report.fstat

Details

Even though Fst and its relatives can predict evolutionary processes (Holsinger & Weir, 2009), they are not true measures of genetic differentiation in the sense that they are dependent on the diversity within populations (Meirmans & Hedrick, 2011), the number of populations analysed (Alcala & Rosenberg, 2017) and are not monotonic (Sherwin et al., 2017). Recent approaches have been developed to accommodate these mathematical restrictions (G'ST; "Gst_H"; Hedrick, 2005, and Jost's D; "Dest"; Jost, 2008). More recently, novel approaches based on information theory (Mutual Information; Sherwin et al., 2017) and allele frequencies (Allele Frequency Difference; Berner, 2019) have distinct properties that make them valuable resources to interpret genetic differentiation between populations.

Note that each measure of genetic differentiation has advantages and drawbacks, and the decision of using a particular measure is usually based on the research question.

Statistics calculated

The equations used to calculate the statistics are shown below.

• *Ho* - Unbiased estimate of observed heterozygosity across subpopulations (Nei, 1987, pp. 164, eq. 7.38) is calculated as:

$$Ho = 1 - \sum_{k} \sum_{i} \frac{Pkii}{s}$$

where *Pkii* represents the proportion of homozygote *ii* for allele *i* in individual *k* and *s* represents the number of subpopulations.

• *Hs* - Unbiased estimate of the expected heterozygosity under Hardy-Weinberg equilibrium across subpopulations (Nei, 1987, pp. 164, eq. 7.39) is calculated as:

$$Hs = \frac{\tilde{n}}{\tilde{n} - 1} \left[1 - \sum_{i} \overline{p_{i}^{2}} - \frac{Ho}{2\tilde{n}} \right]$$

where $\tilde{n} = \frac{1}{mean(1/nk)}$ and $\overline{p_t^2} = \frac{\sum_k p_{kl}^2}{s}$

where \tilde{n} is the harmonic mean of nk (the number of individuals in each subpopulation), pki is the proportion (sometimes misleadingly called frequency) of allele *i* in subpopulation *k*.

• *Ht* - Heterozygosity for the total population (Nei, 1987, pp. 164, eq. 7.40) is calculated as:

$$Ht = 1 - \sum_{i} \bar{p_i}^2 + \frac{Hs}{\tilde{n}} - \frac{Ho}{2\tilde{n}s}$$

where $\bar{p_i} = \sum_k \frac{p_{ki}}{s}$

• *Dst* - The average allele frequency differentiation between populations (Nei, 1987, pp. 163) is calculated as:

$$Dst = Ht - Hs$$

• *Htp* - Unbiased estimate of Heterozygosity for the total population (Nei, 1987, pp. 165) is calculated as:

$$Ht' = Hs + Dst'$$

• *Dstp* - Unbiased estimate of the average allele frequency differentiation between populations (Nei, 1987, pp. 165) is calculated as:

$$Dst = Ht - Hs$$

• *Fst* - Measure of the extent of genetic differentiation of subpopulations (Nei, 1987, pp. 162, eq. 7.34) is calculated as:

$$F_{ST} = \frac{Dst}{Ht}$$

• *Fstp* - Unbiased measure of the extent of genetic differentiation of subpopulations (Nei, 1987, pp. 163, eq. 7.36) is calculated as:

$$F_{ST'} = \frac{Dst'}{Ht'}$$

• Dest - Jost's D (Jost, 2008, eq. 12) is calculated as:

$$Dst = Ht - Hs$$

• *Gst-max* - The maximum level that Gst can obtain for the observed amount of genetic variation (Hedrick 2005, eq. 4a) is calculated as:

$$G_{ST(max)} = \frac{(s-1)(1-Hs)}{s-1+Hs}$$

• *Gst-H* - Gst standardized by the maximum level that it can obtain for the observed amount of genetic variation (Hedrick 2005, eq. 4b) is calculated as:

$$G_{ST(H)} = \frac{F_{ST'}}{G_{ST(max)}}$$

Confidence Intervals

The uncertainty of a parameter, in this case the mean of the statistic, can be summarised by a confidence interval (CI) which includes the true parameter value with a specified probability (i.e. confidence level; the parameter "conf" in this function).

In this function, CI are obtained using Bootstrap which is an inference method that samples with replacement the data (i.e. loci) and calculates the statistics every time.

This function uses the function boot (package boot) to perform the bootstrap replicates and the function boot.ci (package boot) to perform the calculations for the CI.

Four different types of nonparametric CI can be calculated (parameter "CI.type" in this function):

- First order normal approximation interval ("norm").
- Basic bootstrap interval ("basic").
- Bootstrap percentile interval ("perc").
- Adjusted bootstrap percentile interval ("bca").

The studentized bootstrap interval ("stud") was not included in the CI types because it is computationally intensive, it may produce estimates outside the range of plausible values and it has been found to be erratic in practice, see for example the "Studentized (t) Intervals" section in:

https://www.r-bloggers.com/2019/09/understanding-bootstrap-confidence-interval-output-from-the-r-b

Nice tutorials about the different types of CI can be found in:

https://www.datacamp.com/tutorial/bootstrap-r

and

https://www.r-bloggers.com/2019/09/understanding-bootstrap-confidence-interval-output-from-the-r-bo Efron and Tibshirani (1993, p. 162) and Davison and Hinkley (1997, p. 194) suggest that the

number of bootstrap replicates should be between 1000 and 2000.

gl.report.fstat

It is important to note that unreliable confidence intervals will be obtained if too few number of bootstrap replicates are used. Therefore, the function boot.ci will throw warnings and errors if bootstrap replicates are too few. Consider increasing then number of bootstrap replicates to at least 200.

The "bca" interval is often cited as the best for theoretical reasons, however it may produce unstable results if the bootstrap distribution is skewed or has extreme values. For example, you might get the warning "extreme order statistics used as endpoints" or the error "estimated adjustment 'a' is NA". In this case, you may want to use more bootstrap replicates or a different method or check your data for outliers.

The error "estimated adjustment 'w' is infinite" means that the estimated adjustment 'w' for the "bca" interval is infinite, which can happen when the empirical influence values are zero or very close to zero. This can be caused by various reasons, such as:

The number of bootstrap replicates is too small, the statistic of interest is constant or nearly constant across the bootstrap samples, the data contains outliers or extreme values.

You can try some possible solutions, such as:

Increasing the number of bootstrap replicates, using a different type of bootstrap confidence interval or removing or transforming the outliers or extreme values.

Plotting

The plot can be customised by including any parameter(s) from the function heatmap.2 (package gplots).

For the color palette you could try for example:

```
>library(viridis)
```

> res <- gl.report.fstat(platypus.gl, palette.divergent = viridis)</pre>

If a plot.file is given, the plot arising from this function is saved as an "RDS" binary file using the function saveRDS (package base); can be reloaded with function readRDS (package base). A file name must be specified for the plot to be saved.

If a plot directory (plot.dir) is specified, the gplot binary is saved to that directory; otherwise to the tempdir().

Your plot might not shown in full because your 'Plots' pane is too small (in RStudio). Increase the size of the 'Plots' pane before running the function. Alternatively, use the parameter 'plot.file' to save the plot to a file.

Parallelisation

If the parameter ncpus > 1, parallelisation is enabled. In Windows, parallel computing employs a "socket" approach that starts new copies of R on each core. POSIX systems, on the other hand (Mac, Linux, Unix, and BSD), utilise a "forking" approach that replicates the whole current version of R and transfers it to a new core.

Opening and terminating R sessions in each core involves a significant amount of processing time, therefore parallelisation in Windows machines is only quicker than not using parallelisation when nboots > 1000-2000.

Value

Two lists, the first list contains matrices with genetic statistics taken pairwise by population, the second list contains tables with the genetic statistics for each pair of populations. If nboots > 0,

tables with the four statistics calculated with Low Confidence Intervals (LCI) and High Confidence Intervals (HCI).

Author(s)

Custodian: Luis Mijangos - Post to https://groups.google.com/d/forum/dartr

References

- Alcala, N., & Rosenberg, N. A. (2017). Mathematical constraints on FST: Biallelic markers in arbitrarily many populations. Genetics (206), 1581-1600.
- Berner, D. (2019). Allele frequency difference AFD-an intuitive alternative to FST for quantifying genetic population differentiation. Genes, 10(4), 308.
- Davison AC, Hinkley DV (1997). Bootstrap Methods and their Application. Cambridge University Press: Cambridge.
- Efron, B. (1979). Bootstrap methods: Another look at the jackknife. Annals of Statistics 7, 1–26.
- Efron B, Tibshirani RJ (1993). An Introduction to the Bootstrap. Chapman and Hall: London.
- Hedrick, P. W. (2005). A standardized genetic differentiation measure. Evolution, 59(8), 1633-1638.
- Holsinger, K. E. (2012). Lecture notes in population genetics.
- Holsinger, K. E., & Weir, B. S. (2009). Genetics in geographically structured populations: defining, estimating and interpreting FST. Nature Reviews Genetics, 10(9), 639-650.
- Jost, L. (2008). GST and its relatives do not measure differentiation. Molecular Ecology, 17(18), 4015-4026.
- Meirmans, P. G., & Hedrick, P. W. (2011). Assessing population structure: FST and related measures. Molecular Ecology Resources, 11(1), 5-18.
- Nei, M. (1987). Molecular evolutionary genetics: Columbia University Press.
- Sherwin, W. B., Chao, A., Jost, L., & Smouse, P. E. (2017). Information theory broadens the spectrum of molecular ecology and evolution. Trends in Ecology & Evolution, 32(12), 948-963.

See Also

Other matched reports: gl.mahal.assign(), gl.report.bases(), gl.report.factorloadings(), gl.report.monomorphs()

Examples

res <- gl.report.fstat(platypus.gl)</pre>

gl.report.hamming

Calculates the pairwise Hamming distance between DArT trimmed DNA sequences

Description

Hamming distance is calculated as the number of base differences between two sequences which can be expressed as a count or a proportion. Typically, it is calculated between two sequences of equal length. In the context of DArT trimmed sequences, which differ in length but which are anchored to the left by the restriction enzyme recognition sequence, it is sensible to compare the two trimmed sequences starting from immediately after the common recognition sequence and terminating at the last base of the shorter sequence.

Usage

```
gl.report.hamming(
    x,
    rs = 5,
    threshold = 3,
    tag.length = 69,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
    plot.dir = NULL,
    plot.file = NULL,
    probar = FALSE,
    verbose = NULL
)
```

Arguments

х	Name of the genlight object containing the SNP data [required].
rs	Number of bases in the restriction enzyme recognition sequence [default 5].
threshold	Minimum acceptable base pair difference for display on the boxplot and histogram [default 3].
tag.length	Typical length of the sequence tags [default 69].
plot.display	Specify if plot is to be produced [default TRUE].
plot.theme	User specified theme [default theme_dartR()].
plot.colors	Vector with two color names for the borders and fill [default c("#2171B5", "#6BAED6")].
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]
plot.file	Filename (minus extension) for the RDS plot file [Required for plot save]
probar	If TRUE, a progress bar is displayed during run [defalut FALSE]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Details

The function gl.filter.hamming will filter out one of two loci if their Hamming distance is less than a specified percentage Hamming distance can be computed by exploiting the fact that the dot product of two binary vectors x and (1-y) counts the corresponding elements that are different between x and y. This approach can also be used for vectors that contain more than two possible values at each position (e.g. A, C, T or G). If a pair of DNA sequences are of differing length, the longer is truncated. The algorithm is that of Johann de Jong https://johanndejong.wordpress.com/ 2015/10/02/faster-hamming-distance-in-r-2/ as implemented in utils.hamming If plot.file is specified, plots are saved to the directory specified by the user, or the global default working directory set by gl.set.wd() or to the tempdir(). Examples of other themes that can be used can be consulted in

- https://ggplot2.tidyverse.org/reference/ggtheme.html and
- https://yutannihilation.github.io/allYourFigureAreBelongToUs/ggthemes/

Value

Returns unaltered genlight object

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

gl.filter.hamming

```
Other matched report: gl.filter.excess.het(), gl.report.allna(), gl.report.callrate(),
gl.report.locmetric(), gl.report.maf(), gl.report.overshoot(), gl.report.pa(), gl.report.rdepth(),
gl.report.reproducibility(), gl.report.secondaries(), gl.report.taglength()
```

Examples

```
gl.report.hamming(testset.gl[,1:100])
gl.report.hamming(testset.gs[,1:100])
#' # SNP data
test <- platypus.gl
test <- gl.subsample.loc(platypus.gl,n=50)
result <- gl.report.hamming(test, verbose=3)
result <- gl.report.hamming(test, plot.file="ttest", verbose=3)</pre>
```

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```
gl.report.heterozygosity
```

Reports observed, expected and unbiased heterozygosities and FIS (inbreeding coefficient) by population or by individual from SNP data

Description

Calculates the observed, expected and unbiased expected (i.e. corrected for sample size) heterozygosities and FIS (inbreeding coefficient) for each population or the observed heterozygosity for each individual in a genlight object.

Usage

```
gl.report.heterozygosity(
 х,
 method = "pop",
 n.invariant = 0,
  subsample.pop = FALSE,
 n.limit = 10,
 nboots = 0,
 boot.method = "ind",
  conf = 0.95,
 CI.type = "bca",
  ncpus = 1,
  plot.display = TRUE,
  plot.theme = theme_dartR(),
 plot.colors.pop = gl.colors("dis"),
 plot.colors.ind = gl.colors(2),
 plot.file = NULL,
 plot.dir = NULL,
 error.bar = "SD",
  verbose = NULL
)
```

Arguments

x	Name of the genlight object containing the SNP data [required].
method	Calculate heterozygosity by population (method='pop') or by individual (method='ind') [default 'pop'].
n.invariant	An estimate of the number of invariant sequence tags used to adjust the het- erozygosity rate [default 0].
subsample.pop	Whether subsample populations to estimate observed heterozygosity (see De- tails) [default FALSE].
n.limit	Minimum number of individuals that should have a population to perform sub- sampling to estimate heterozygosity [default 10].

boot.methodboostraping across individuals ("ind") or across loci ("loc") [default "ind"].confThe confidence level of the required interval [default 0.95].CI.typeMethod to estimate confidence intervals. One of "norm", "basic", "perc" or "bca" [default "bca"].ncpusNumber of processes to be used in parallel operation. If ncpus > 1 parallel operation is activated, see "Details" section [default 1].plot.displaySpecify if plot is to be produced [default TRUE].plot.themeTheme for the plot. See Details for options [default theme_dartR()].plot.colors.popA color palette for population plots or a list with as many colors as there are populations in the dataset [default gl.colors("dis")].plot.colors.indList of two color names for the borders and fill of the plot by individual [default gl.colors(2)].plot.fileName for the RDS binary file to save (base name only, exclude extension) [de-
CI.type Method to estimate confidence intervals. One of "norm", "basic", "perc" or "bca" [default "bca"]. ncpus Number of processes to be used in parallel operation. If ncpus > 1 parallel operation is activated, see "Details" section [default 1]. plot.display Specify if plot is to be produced [default TRUE]. plot.theme Theme for the plot. See Details for options [default theme_dartR()]. plot.colors.pop A color palette for population plots or a list with as many colors as there are populations in the dataset [default gl.colors("dis")]. plot.colors.ind List of two color names for the borders and fill of the plot by individual [default gl.colors(2)]. plot.file Name for the RDS binary file to save (base name only, exclude extension) [de-
<pre>"bca" [default "bca"]. ncpus Number of processes to be used in parallel operation. If ncpus > 1 parallel operation is activated, see "Details" section [default 1]. plot.display Specify if plot is to be produced [default TRUE]. plot.theme Theme for the plot. See Details for options [default theme_dartR()]. plot.colors.pop</pre>
<pre>operation is activated, see "Details" section [default 1]. plot.display Specify if plot is to be produced [default TRUE]. plot.theme Theme for the plot. See Details for options [default theme_dartR()]. plot.colors.pop</pre>
<pre>plot.theme Theme for the plot. See Details for options [default theme_dartR()]. plot.colors.pop</pre>
<pre>plot.colors.pop</pre>
A color palette for population plots or a list with as many colors as there are populations in the dataset [default gl.colors("dis")]. plot.colors.ind List of two color names for the borders and fill of the plot by individual [default gl.colors(2)]. plot.file Name for the RDS binary file to save (base name only, exclude extension) [de-
populations in the dataset [default gl.colors("dis")]. plot.colors.ind List of two color names for the borders and fill of the plot by individual [default gl.colors(2)]. plot.file Name for the RDS binary file to save (base name only, exclude extension) [de-
List of two color names for the borders and fill of the plot by individual [default gl.colors(2)]. plot.file Name for the RDS binary file to save (base name only, exclude extension) [de-
gl.colors(2)]. plot.file Name for the RDS binary file to save (base name only, exclude extension) [de-
fault NULL].
plot.dir Directory to save the plot RDS files [default as specified by the global working directory or tempdir()].
error.bar Statistic to be plotted as error bar either "SD" (standard deviation) or "SE" (stan- dard error) or "CI" (confidence intervals) [default "SD"].
verbose Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity].

Details

Observed heterozygosity for a population takes the proportion of heterozygous loci for each individual and averages it over all individuals in that population. The calculations take into account missing values.

Expected heterozygosity for a population takes the expected proportion of heterozygotes, that is, expected under Hardy-Weinberg equilibrium, for each locus, then averages this across the loci for an average estimate for the population.

The unbiased expected heterozygosity is calculated using the correction for sample size following equation 2 from Nei 1978.

Accuracy of all heterozygosity estimates is affected by small sample sizes, and so is their comparison between populations or repeated analysis. Expected heterozygosities are less affected because their calculations are based on allele frequencies while observed heterozygosities are strongly susceptible to sampling effects when the sample size is small.

Observed heterozygosity for individuals is calculated as the proportion of loci that are heterozygous for that individual.

Finally, the loci that are invariant across all individuals in the dataset (that is, across populations), is typically unknown. This can render estimates of heterozygosity analysis specific, and so it is not valid to compare such estimates across species or even across different analyses (see Schimdt et al 2021). This is a similar problem faced by microsatellites. If you have an estimate of the number

of invariant sequence tags (loci) in your data, such as provided by gl.report.secondaries, you can specify it with the n.invariant parameter to standardize your estimates of heterozygosity. This is called autosomal heterozygosities by Schimddt et al (2021).

NOTE: It is important to realise that estimation of adjusted (autosomal) heterozygosity requires that secondaries not to be removed.

Heterozygosities and FIS (inbreeding coefficient) are calculated by locus within each population using the following equations, and then averaged across all loci:

- Observed heterozygosity (Ho) = number of heterozygotes / n_Ind, where n_Ind is the number of individuals without missing data for that locus.
- Observed heterozygosity adjusted (Ho.adj) <- Ho * n_Loc / (n_Loc + n.invariant), where n_Loc is the number of loci that do not have all missing data and n.invariant is an estimate of the number of invariant loci to adjust heterozygosity.
- Expected heterozygosity (He) = $1 (p^2 + q^2)$, where p is the frequency of the reference allele and q is the frequency of the alternative allele.
- Expected heterozygosity adjusted (He.adj) = He * n_Loc / (n_Loc + n.invariant)
- Unbiased expected heterozygosity (uHe) = He * (2 * n_Ind / (2 * n_Ind 1))
- Inbreeding coefficient (FIS) = 1 Ho / uHe

Function's output

Output for method='pop' is an ordered barchart of observed heterozygosity, unbiased expected heterozygosity and FIS (Inbreeding coefficient) across populations together with a table of mean observed and expected heterozygosities and FIS by population and their respective standard deviations (SD). In the output, it is also reported by population: the number of loci used to estimate heterozygosity (n.Loc), the number of polymorphic loci (polyLoc), the number of monomorphic loci (monoLoc) and loci with all missing data (all_NALoc).

Output for method='ind' is a histogram and a boxplot of heterozygosity across individuals.

If a plot.file is given, the ggplot arising from this function is saved as an "RDS" binary file using saveRDS(); can be reloaded with readRDS(). A file name must be specified for the plot to be saved.

If a plot directory (plot.dir) is specified, the ggplot binary is saved to that directory; otherwise to the tempdir().

Examples of other themes that can be used can be consulted in:

- https://ggplot2.tidyverse.org/reference/ggtheme.html and
- https://yutannihilation.github.io/allYourFigureAreBelongToUs/ggthemes/

Subsampling populations

To test the effect of five population sample sizes (n = 10, 5, 4, 3, 2) on observed heterozygosity estimates, the function subsamples individuals, without replacement. The subsampling is repeated 10 times for each sample size n. This approach is not an implementation of Schmidt et al (2021). Please refer to this paper for additional complexities in estimating heterozygosity using SNP data.

Error bars

The best method for presenting or assessing genetic statistics depends on the type of data you have and the specific questions you're trying to answer. Here's a brief overview of when you might use each method:

1. Confidence Intervals ("CI"):

- Usage: Often used to convey the precision of an estimate.

- Advantage: Confidence intervals give a range in which the true parameter (like a population mean) is likely to fall, given the data and a specified probability (like 95%).

- In Context: For genetic statistics, if you're estimating a parameter, a 95% CI gives you a range in which you're 95% confident the true parameter lies.

2. Standard Deviation ("SD"):

- Usage: Describes the amount of variation from the average in a set of data.

- Advantage: Allows for an understanding of the spread of individual data points around the mean.

- In Context: If you're looking at the distribution of a quantitative trait (like height) in a population with a particular genotype, the SD can describe how much individual heights vary around the average height.

3. Standard Error ("SE"):

- Usage: Describes the precision of the sample mean as an estimate of the population mean.

- Advantage: Smaller than the SD in large samples; it takes into account both the SD and the sample size.

- In Context: If you want to know how accurately your sample mean represents the population mean, you'd look at the SE.

Recommendation:

- If you're trying to convey the precision of an estimate, confidence intervals are very useful.
- For understanding variability within a sample, standard deviation is key.
- To see how well a sample mean might estimate a population mean, consider the standard error.

In practice, geneticists often use a combination of these methods to analyze and present their data, depending on their research questions and the nature of the data.

Confidence Intervals

The uncertainty of a parameter, in this case the mean of the statistic, can be summarised by a confidence interval (CI) which includes the true parameter value with a specified probability (i.e. confidence level; the parameter "conf" in this function).

In this function, CI are obtained using Bootstrap which is an inference method that samples with replacement the data (i.e. loci) and calculates the statistics every time.

This function uses the function boot (package boot) to perform the bootstrap replicates and the function boot.ci (package boot) to perform the calculations for the CI.

Four different types of nonparametric CI can be calculated (parameter "CI.type" in this function):

- First order normal approximation interval ("norm").
- Basic bootstrap interval ("basic").
- Bootstrap percentile interval ("perc").
- Adjusted bootstrap percentile interval ("bca").

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The studentized bootstrap interval ("stud") was not included in the CI types because it is computationally intensive, it may produce estimates outside the range of plausible values and it has been found to be erratic in practice, see for example the "Studentized (t) Intervals" section in:

https://www.r-bloggers.com/2019/09/understanding-bootstrap-confidence-interval-output-from-the-r-b

Nice tutorials about the different types of CI can be found in:

https://www.datacamp.com/tutorial/bootstrap-r

and

https://www.r-bloggers.com/2019/09/understanding-bootstrap-confidence-interval-output-from-the-r-b

Efron and Tibshirani (1993, p. 162) and Davison and Hinkley (1997, p. 194) suggest that the number of bootstrap replicates should be between 1000 and 2000.

It is important to note that unreliable confidence intervals will be obtained if too few number of bootstrap replicates are used. Therefore, the function boot.ci will throw warnings and errors if bootstrap replicates are too few. Consider increasing the number of bootstrap replicates to at least 200.

The "bca" interval is often cited as the best for theoretical reasons, however it may produce unstable results if the bootstrap distribution is skewed or has extreme values. For example, you might get the warning "extreme order statistics used as endpoints" or the error "estimated adjustment 'a' is NA". In this case, you may want to use more bootstrap replicates or a different method or check your data for outliers.

The error "estimated adjustment 'w' is infinite" means that the estimated adjustment 'w' for the "bca" interval is infinite, which can happen when the empirical influence values are zero or very close to zero. This can be caused by various reasons, such as:

The number of bootstrap replicates is too small, the statistic of interest is constant or nearly constant across the bootstrap samples, the data contains outliers or extreme values.

You can try some possible solutions, such as:

Increasing the number of bootstrap replicates, using a different type of bootstrap confidence interval or removing or transforming the outliers or extreme values.

Parallelisation

If the parameter ncpus > 1, parallelisation is enabled. In Windows, parallel computing employs a "socket" approach that starts new copies of R on each core. POSIX systems, on the other hand (Mac, Linux, Unix, and BSD), utilise a "forking" approach that replicates the whole current version of R and transfers it to a new core.

Opening and terminating R sessions in each core involves a significant amount of processing time, therefore parallelisation in Windows machines is only quicker than not using parallelisation when nboots > 1000-2000.

Value

A dataframe containing population labels, heterozygosities, FIS, their standard deviations and sample sizes.

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

References

- Nei, M. (1978). Estimation of average heterozygosity and genetic distance from a small number of individuals. Genetics, 89(3), 583-590.
- Schmidt, T. L., Jasper, M. E., Weeks, A. R., & Hoffmann, A. A. (2021). Unbiased population heterozygosity estimates from genome-wide sequence data. Methods in Ecology and Evolution, 12(10), 1888-1898.

See Also

```
gl.filter.heterozygosity
```

Other unmatched report: gl.allele.freq(), gl.report.basics(), gl.report.diversity(), gl.report.excess.het(), gl.report.polyploid_heterozygosity()

Examples

```
require("dartR.data")
df <- gl.report.heterozygosity(platypus.gl)
df <- gl.report.heterozygosity(platypus.gl,method='ind')
n.inv <- gl.report.secondaries(platypus.gl)
gl.report.heterozygosity(platypus.gl, n.invariant = n.inv[7, 2])
gl.report.heterozygosity(platypus.gl, subsample.pop = TRUE)</pre>
```

df <- gl.report.heterozygosity(platypus.gl)</pre>

gl.report.hwe Reports departure from Hardy-Weinberg proportions

Description

Calculates the probabilities of agreement with H-W proportions based on observed frequencies of reference homozygotes, heterozygotes and alternate homozygotes.

Usage

```
gl.report.hwe(
    x,
    subset = "each",
    method_sig = "Exact",
    multi_comp = FALSE,
    multi_comp_method = "BY",
    alpha_val = 0.05,
    pvalue_type = "midp",
    cc_val = 0.5,
    sig_only = TRUE,
    min_sample_size = 5,
    plot.out = TRUE,
    plot_colors = gl.colors("2c"),
```

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```
max_plots = 4,
save2tmp = FALSE,
verbose = NULL
```

Arguments

)

x	Name of the genlight object containing the SNP data [required].	
subset	Whether to perform H-W tests within each population ("each"), or taking all individuals as one population ("all") (see details) [default 'each'].	
method_sig	Method for determining statistical significance: 'ChiSquare' or 'Exact' [default 'Exact'].	
multi_comp	Whether to adjust p-values for multiple comparisons [default FALSE].	
multi_comp_meth	nod	
	Method to adjust p-values for multiple comparisons: 'holm', 'hochberg', 'hom- mel', 'bonferroni', 'BH', 'BY', 'fdr' (see details) [default 'fdr'].	
alpha_val	Level of significance for testing [default 0.05].	
pvalue_type	Type of p-value to be used in the Exact method. Either 'dost', 'selome', 'midp' (see details) [default 'midp'].	
cc_val	The continuity correction applied to the ChiSquare test [default 0.5].	
sig_only	Whether the returned table should include loci with a significant departure from Hardy-Weinberg proportions [default TRUE].	
min_sample_size		
	Minimum number of individuals per population in which perform H-W tests [default 5].	
plot.out	If TRUE, will produce Ternary Plot(s) [default TRUE].	
plot_colors	Vector with two color names for the significant and not-significant loci [default gl.colors("2c")].	
<pre>max_plots</pre>	Maximum number of plots to print per page [default 4].	
save2tmp	If TRUE, saves any ggplots and listings to the session temporary directory (tem- pdir) [default FALSE].	
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity].	

Details

There are several factors that can cause deviations from Hardy-Weinberg proportions including: mutation, finite population size, selection, population structure, age structure, assortative mating, sex linkage, nonrandom sampling and genotyping errors. Therefore, testing for Hardy-Weinberg proportions should be a process that involves a careful evaluation of the results, a good place to start is Waples (2015). Note that tests for H-W proportions are only valid if there is no population substructure (assuming random mating) and have sufficient power only when there is sufficient sample size (n individuals > 15). Populations can be defined in three ways:

• Merging all populations in the dataset using subset = 'all'.

- Within each population separately using: subset = 'each'.
- Within selected populations using for example: subset = c('pop1','pop2').

Two different statistical methods to test for deviations from Hardy Weinberg proportions:

- The classical chi-square test (method_sig='ChiSquare') based on the function HWChisq of the R package HardyWeinberg. By default a continuity correction is applied (cc_val=0.5). The continuity correction can be turned off (by specifying cc_val=0), for example in cases of extreme allele frequencies in which the continuity correction can lead to excessive type 1 error rates.
- The exact test (method_sig='Exact') based on the exact calculations contained in the function HWExactStats of the R package HardyWeinberg, and described in Wigginton et al. (2005). The exact test is recommended in most cases (Wigginton et al., 2005). Three different methods to estimate p-values (pvalue_type) in the Exact test can be used:
 - 'dost' p-value is computed as twice the tail area of a one-sided test.
 - 'selome' p-value is computed as the sum of the probabilities of all samples less or equally likely as the current sample.
 - 'midp', p-value is computed as half the probability of the current sample + the probabilities of all samples that are more extreme.

The standard exact p-value is overly conservative, in particular for small minor allele frequencies. The mid p-value ameliorates this problem by bringing the rejection rate closer to the nominal level, at the price of occasionally exceeding the nominal level (Graffelman & Moreno, 2013).

Correction for multiple tests can be applied using the following methods based on the function p.adjust:

- 'holm' is also known as the sequential Bonferroni technique (Rice, 1989). This method has a greater statistical power than the standard Bonferroni test, however this method becomes very stringent when many tests are performed and many real deviations from the null hypothesis can go undetected (Waples, 2015).
- 'hochberg' based on Hochberg, 1988.
- 'hommel' based on Hommel, 1988. This method is more powerful than Hochberg's, but the difference is usually small.
- 'bonferroni' in which p-values are multiplied by the number of tests. This method is very stringent and therefore has reduced power to detect multiple departures from the null hypothesis.
- 'BH' based on Benjamini & Hochberg, 1995.
- 'BY' based on Benjamini & Yekutieli, 2001.

The first four methods are designed to give strong control of the family-wise error rate. The last two methods control the false discovery rate (FDR), the expected proportion of false discoveries among the rejected hypotheses. The false discovery rate is a less stringent condition than the family-wise error rate, so these methods are more powerful than the others, especially when number of tests is large. The number of tests on which the adjustment for multiple comparisons is the number of populations times the number of loci. **Ternary plots** Ternary plots can be used to visualise patterns of H-W proportions (plot.out = TRUE). P-values and the statistical (non)significance

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of a large number of bi-allelic markers can be inferred from their position in a ternary plot. See Graffelman & Morales-Camarena (2008) for further details. Ternary plots are based on the function HWTernaryPlot from the package HardyWeinberg. Each vertex of the Ternary plot represents one of the three possible genotypes for SNP data: homozygous for the reference allele (AA), heterozygous (AB) and homozygous for the alternative allele (BB). Loci deviating significantly from Hardy-Weinberg proportions after correction for multiple tests are shown in pink. The blue parabola represents Hardy-Weinberg equilibrium, and the area between green lines represents the acceptance region. For these plots to work it is necessary to install the package ggtern.

Value

A dataframe containing loci, counts of reference SNP homozygotes, heterozygotes and alternate SNP homozygotes; probability of departure from H-W proportions, per locus significance with and without correction for multiple comparisons and the number of population where the same locus is significantly out of HWE.

Author(s)

Custodian: Luis Mijangos - Post to https://groups.google.com/d/forum/dartr

References

- Benjamini, Y., and Yekutieli, D. (2001). The control of the false discovery rate in multiple testing under dependency. Annals of Statistics, 29, 1165–1188.
- Graffelman, J. (2015). Exploring Diallelic Genetic Markers: The Hardy Weinberg Package. Journal of Statistical Software 64:1-23.
- Graffelman, J. & Morales-Camarena, J. (2008). Graphical tests for Hardy-Weinberg equilibrium based on the ternary plot. Human Heredity 65:77-84.
- Graffelman, J., & Moreno, V. (2013). The mid p-value in exact tests for Hardy-Weinberg equilibrium. Statistical applications in genetics and molecular biology, 12(4), 433-448.
- Hochberg, Y. (1988). A sharper Bonferroni procedure for multiple tests of significance. Biometrika, 75, 800–803.
- Hommel, G. (1988). A stagewise rejective multiple test procedure based on a modified Bonferroni test. Biometrika, 75, 383–386.
- Rice, W. R. (1989). Analyzing tables of statistical tests. Evolution, 43(1), 223-225.
- Waples, R. S. (2015). Testing for Hardy–Weinberg proportions: have we lost the plot?. Journal of heredity, 106(1), 1-19.
- Wigginton, J.E., Cutler, D.J., & Abecasis, G.R. (2005). A Note on Exact Tests of Hardy-Weinberg Equilibrium. American Journal of Human Genetics 76:887-893.

See Also

gl.filter.hwe

gl.report.ld

Calculates pairwise population based Linkage Disequilibrium across all loci using the specified number of cores @family matched report

Description

This function is implemented in a parallel fashion to speed up the process. There is also the ability to restart the function if crashed by specifying the chunk file names or restarting the function exactly in the same way as in the first run. This is implemented because sometimes, due to connectivity loss between cores, the function may crash half way. Before running the function, it is advisable to use the function gl.filter.allna to remove loci with all missing data.

Usage

```
gl.report.ld(
    x,
    name = NULL,
    save = TRUE,
    outpath = tempdir(),
    nchunks = 2,
    ncores = 1,
    chunkname = NULL,
    probar = FALSE,
    verbose = NULL
)
```

Arguments

x	A genlight or genind object created (genlight objects are internally converted via gl2gi to genind) [required].
name	Character string for rdata file. If not given genind object name is used [default NULL].
save	Switch if results are saved in a file [default TRUE].
outpath	Folder where chunks and results are saved (if save=TRUE) [default tempdir()].
nchunks	How many subchunks will be used (the less the faster, but if the routine crashes more bits are lost) [default 2].
ncores	How many cores should be used [default 1].
chunkname	The name of the chunks for saving [default NULL].
probar	if TRUE, a progress bar is displayed for long loops [default = TRUE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity]

Value

Returns calculation of pairwise LD across all loci between subpopulations. This functions uses if specified many cores on your computer to speed up. And if save is used can restart (if save=TRUE is used) with the same command starting where it crashed. The final output is a data frame that holds all statistics of pairwise LD between loci. (See ?LD in package genetics for details).

Author(s)

Bernd Gruber (Post to https://groups.google.com/d/forum/dartr)

gl.report.ld.map Calculates pairwise linkage disequilibrium by population

Description

This function calculates pairwise linkage disequilibrium (LD) by population using the function ld from package snpStats. If SNPs are not mapped to a reference genome, the parameter ld.max.pairwise should be set as NULL (the default). In this case, the function will assign the same chromosome ("1") to all the SNPs in the dataset and assign a sequence from 1 to n loci as the position of each SNP. The function will then calculate LD for all possible SNP pair combinations. If SNPs are mapped to a reference genome, the parameter ld.max.pairwise should be filled out (i.e. not NULL). In this case, the information for SNP's position should be stored in the genlight accessor "@position" and the SNP's chromosome name in the accessor "@chromosome" (see examples). The function will then calculate LD within each chromosome and for all possible SNP pair combinations within a distance of ld.max.pairwise.

Usage

```
gl.report.ld.map(
 х,
 ld.max.pairwise = 1e+06,
 maf = 0.05,
 ld.stat = "R.squared",
  ind.limit = 10,
  stat.keep = "AvgPIC",
  ld.threshold.pops = 0.2,
  plot.display = TRUE,
  plot.theme = theme_dartR(),
  plot.file = NULL,
  plot.dir = NULL,
  histogram.colors = NULL,
  boxplot.colors = NULL,
 bins = 50,
  verbose = NULL
)
```

Arguments

x	Name of the genlight object containing the SNP data [required].	
ld.max.pairwise		
	Maximum distance in number of base pairs at which LD should be calculated [default 1000000].	
maf	Minor allele frequency (by population) threshold to filter out loci. If a value > 1 is provided it will be interpreted as MAC (i.e. the minimum number of times an allele needs to be observed) [default 0.05].	
ld.stat	The LD measure to be calculated: "LLR", "OR", "Q", "Covar", "D.prime", "R.squared", and "R". See function ld from snpstats (package snpStats) for de- tails [default "R.squared"].	
ind.limit	Minimum number of individuals that a population should contain to take it in account to report loci in LD [default 10].	
stat.keep	Name of the column from the slot loc.metrics to be used to choose SNP to be kept [default "AvgPIC"].	
ld.threshold.pd	ops	
	LD threshold to report in the plot of "Number of populations in which the same SNP pair are in LD" [default 0.2].	
plot.display	If TRUE, histograms of base composition are displayed in the plot window [de-fault TRUE].	
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].	
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL]	
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]	
histogram.colors		
	Vector with two color names for the borders and fill [default NULL].	
boxplot.colors	A color palette for box plots by population or a list with as many colors as there are populations in the dataset [default NULL].	
bins	Number of bins to display in histograms [default 50].	
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].	

Details

This function reports LD between SNP pairs by population. The function gl.filter.ld filters out the SNPs in LD using as input the results of gl.report.ld.map. The actual number of SNPs to be filtered out depends on the parameters set in the function gl.filter.ld. Boxplots of LD by population and a histogram showing LD frequency are presented.

Value

A dataframe with information for each SNP pair in LD.

gl.report.locmetric

Author(s)

Custodian: Luis Mijangos - Post to https://groups.google.com/d/forum/dartr

See Also

gl.filter.ld

Other graphics: gl.colors(), gl.map.interactive(), gl.plot.heatmap(), gl.select.colors(), gl.select.shapes(), gl.smearplot(), gl.tree.nj()

Examples

```
require("dartR.data")
x <- platypus.gl
x <- gl.filter.callrate(x,threshold = 1)
x <- gl.filter.monomorphs(x)
x$position <- x$other$loc.metrics$ChromPos_Platypus_Chrom_NCBIv1
x$chromosome <- as.factor(x$other$loc.metrics$Chrom_Platypus_Chrom_NCBIv1)
ld_res <- gl.report.ld.map(x,ld.max.pairwise = 1000000)</pre>
```

gl.report.locmetric Reports summary of the slot \$other\$loc.metrics

Description

This function reports summary statistics (mean, minimum, average, quantiles), histograms and boxplots for any loc.metric with numeric values (stored in \$other\$loc.metrics) to assist the decision of choosing thresholds for the filter function gl.filter.locmetric.

Usage

```
gl.report.locmetric(
    x,
    metric,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
    plot.dir = NULL,
    verbose = NULL
)
```

Arguments

х	Name of the genlight object containing the SNP or presence/absence (Silico-
	DArT) data [required].
metric	Name of the metric to be used for filtering [required].
plot.display	Specify if plot is to be produced [default TRUE].

plot.theme	User specified theme [default theme_dartR()].
plot.colors	Vector with two color names for the borders and fill [default c("#2171B5", "#6BAED6")].
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]
plot.file	Filename (minus extension) for the RDS plot file [Required for plot save]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity].

Details

The function gl.filter.locmetric will filter out the loci with a locmetric value below a specified threshold. The fields that are included in dartR, and a short description, are found below. Optionally, the user can also set his/her own field by adding a vector into \$other\$loc.metrics as shown in the example. You can check the names of all available loc.metrics via: names(gl\$other\$loc.metrics).

- SnpPosition position (zero is position 1) in the sequence tag of the defined SNP variant base.
- CallRate proportion of samples for which the genotype call is non-missing (that is, not '-').
- OneRatioRef proportion of samples for which the genotype score is 0.
- OneRatioSnp proportion of samples for which the genotype score is 2.
- FreqHomRef proportion of samples homozygous for the Reference allele.
- FreqHomSnp proportion of samples homozygous for the Alternate (SNP) allele.
- FreqHets proportion of samples which score as heterozygous, that is, scored as 1.
- PICRef polymorphism information content (PIC) for the Reference allele.
- PICSnp polymorphism information content (PIC) for the SNP.
- AvgPIC average of the polymorphism information content (PIC) of the reference and SNP alleles.
- AvgCountRef sum of the tag read counts for all samples, divided by the number of samples with non-zero tag read counts, for the Reference allele row.
- AvgCountSnp sum of the tag read counts for all samples, divided by the number of samples with non-zero tag read counts, for the Alternate (SNP) allele row.
- RepAvg proportion of technical replicate assay pairs for which the marker score is consistent.
- rdepth read depth.

Function's output The minimum, maximum, mean and a tabulation of quantiles of the locmetric values against thresholds rate are provided. Output also includes a boxplot and a histogram. Quantiles are partitions of a finite set of values into q subsets of (nearly) equal sizes. In this function q = 20. Quantiles are useful measures because they are less susceptible to long-tailed distributions and outliers. Plot colours can be set with gl.select.colors(). If plot.file is specified, plots are saved to the directory specified by the user, or the global default working directory set by gl.set.wd() or to the tempdir(). Examples of other themes that can be used can be consulted in:

- https://ggplot2.tidyverse.org/reference/ggtheme.html and
- https://yutannihilation.github.io/allYourFigureAreBelongToUs/ggthemes/

gl.report.maf

Value

An unaltered genlight object.

Author(s)

Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

See Also

gl.filter.locmetric

```
Other matched report: gl.filter.excess.het(), gl.report.allna(), gl.report.callrate(),
gl.report.hamming(), gl.report.maf(), gl.report.overshoot(), gl.report.pa(), gl.report.rdepth(),
gl.report.reproducibility(), gl.report.secondaries(), gl.report.taglength()
```

Examples

```
# SNP data
out <- gl.report.locmetric(testset.gl,metric='SnpPosition')
# Tag P/A data
out <- gl.report.locmetric(testset.gs,metric='AvgReadDepth')</pre>
```

gl.report.maf Reports minor allele frequency (MAF) for each locus in a SNP dataset

Description

This script provides summary histograms of MAF for each population and an overall histogram to assist the decision of choosing thresholds for the filter function gl.filter.maf

Usage

```
gl.report.maf(
    x,
    as.pop = NULL,
    maf.limit = 0.5,
    ind.limit = 5,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
    plot.dir = NULL,
    plot.file = NULL,
    bins = 25,
    verbose = NULL
)
```

Arguments

х	Name of the genlight object containing the SNP data [required].
as.pop	Temporarily assign another locus metric as the population for the purposes of deletions [default NULL].
maf.limit	Show histograms MAF range <= maf.limit [default 0.5].
ind.limit	Show histograms only for populations of size greater than ind.limit [default 5].
plot.display	Specify if plot is to be displayed in the graphics window [default TRUE].
plot.theme	User specified theme [default theme_dartR()].
plot.colors	Vector with color names for the borders and fill [default c("#2171B5", "#6BAED6")].
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]
plot.file	Filename (minus extension) for the RDS plot file [Required for plot save].
bins	Number of bins to display in histograms [default 25].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity].

Details

The function gl.filter.maf will filter out the loci with MAF below a specified threshold.

Function's output

The minimum, maximum, mean and a tabulation of MAF quantiles against thresholds rate are provided. Output also includes a boxplot and a histogram.

This function reports the MAF for each of several quantiles. Quantiles are partitions of a finite set of values into q subsets of (nearly) equal sizes. In this function q = 20. Quantiles are useful measures because they are less susceptible to long-tailed distributions and outliers.

Plot colours can be set with gl.select.colors().

If plot.file is specified, plots are saved to the directory specified by the user, or the global default working directory set by gl.set.wd() or to the tempdir().

Examples of other themes that can be used can be consulted in

- https://ggplot2.tidyverse.org/reference/ggtheme.html and
- https://yutannihilation.github.io/allYourFigureAreBelongToUs/ggthemes/

Value

An unaltered genlight object

Author(s)

Custodian: Arthur Georges (Post to https://groups.google.com/d/forum/dartr)

gl.report.monomorphs

See Also

```
gl.filter.maf
```

```
Other matched report: gl.filter.excess.het(), gl.report.allna(), gl.report.callrate(),
gl.report.hamming(), gl.report.locmetric(), gl.report.overshoot(), gl.report.pa(),
gl.report.rdepth(), gl.report.reproducibility(), gl.report.secondaries(), gl.report.taglength()
```

Examples

```
gl <- gl.filter.allna(platypus.gl)
gl.report.maf(gl)</pre>
```

gl.report.monomorphs Reports monomorphic loci

Description

This script reports the number of monomorphic loci and those with all NAs in a genlight {adegenet} object

Usage

gl.report.monomorphs(x, verbose = NULL)

Arguments

х	Name of the input genlight object [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress
	and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity].

Details

A DArT dataset will not have monomorphic loci, but they can arise, along with loci that are scored all NA, when populations or individuals are deleted. Retaining monomorphic loci unnecessarily increases the size of the dataset and will affect some calculations. Note that for SNP data, NAs likely represent null alleles; in tag presence/absence data, NAs represent missing values (presence/absence could not be reliably scored)

Value

An unaltered genlight object

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

gl.filter.monomorphs

Other matched reports: gl.mahal.assign(), gl.report.bases(), gl.report.factorloadings(), gl.report.fstat()

Examples

```
# SNP data
gl.report.monomorphs(testset.gl)
# SilicoDArT data
gl.report.monomorphs(testset.gs)
```

gl.report.overshoot Reports loci for which the SNP has been trimmed from the sequence tag along with the adaptor

Description

This function checks the position of the SNP within the trimmed sequence tag and identifies those for which the SNP position is outside the trimmed sequence tag. This can happen, rarely, when the sequence containing the SNP resembles the adaptor.

Usage

gl.report.overshoot(x, verbose = NULL)

Arguments

х	Name of the genlight object [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress
	and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity].

Details

The SNP genotype can still be used in most analyses, but functions like gl2fasta() will present challenges if the SNP has been trimmed from the sequence tag. Resultant ggplot(s) and the tabulation(s) are saved to the session's temporary directory.

Value

An unaltered genlight object

Author(s)

Custodian: Arthur Georges - Post to https://groups.google.com/d/forum/dartr

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gl.report.pa

See Also

```
gl.filter.overshoot
```

```
Other matched report: gl.filter.excess.het(), gl.report.allna(), gl.report.callrate(),
gl.report.hamming(), gl.report.locmetric(), gl.report.maf(), gl.report.pa(), gl.report.rdepth(),
gl.report.reproducibility(), gl.report.secondaries(), gl.report.taglength()
```

Examples

gl.report.overshoot(testset.gl)

gl.report.pa

Reports private alleles (and fixed alleles) per pair of populations

Description

This function reports private alleles in one population compared with a second population, for all populations taken pairwise. It also reports a count of fixed allelic differences and the mean absolute allele frequency differences (AFD) between pairs of populations.

Usage

```
gl.report.pa(
  х,
  x2 = NULL,
  method = "pairwise",
  loc.names = FALSE,
  test.asym = FALSE,
  test.asym.boot = 100,
  plot.display = FALSE,
 matrix.pa = FALSE,
  plot.font = 14,
 map.interactive = FALSE,
  provider = "Esri.NatGeoWorldMap",
  palette.discrete = NULL,
  plot.file = NULL,
 plot.dir = NULL,
  verbose = NULL
)
```

Arguments

х	Name of the genlight object containing the SNP or SilicoDArT data [required].
x2	If two separate genlight objects are to be compared this can be provided here,
	but they must have the same number of SNPs [default NULL].

method	Method to calculate private alleles: 'pairwise' comparison or compare each population against the rest 'one2rest' [default 'pairwise'].
loc.names	Whether names of loci with private alleles and fixed differences should reported. If TRUE, loci names are reported using a list
test.asym	Bootstrap test for significant differences of private alleles (see details section) [default FALSE].
test.asym.boot	Number of bootstraps [default 100]. [default FALSE].
plot.display	Specify if Sankey plot is to be produced [default FALSE].
matrix.pa	Whether to generate a matrix of private alleles [default FALSE].
plot.font	Numeric font size in pixels for the node text labels [default 14].
map.interactive	
	Specify whether an interactive map showing private alleles between populations is to be produced [default FALSE].
provider	Passed to leaflet [default "Esri.NatGeoWorldMap"].
palette.discret	te
	A discrete palette for the color of populations or a list with as many colors as there are populations in the dataset [default gl.select.colors(x)].
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL]
plot.dir	Directory in which to save files [default = working directory]
verbose	Verbosity: 0, silent, fatal errors only; 1, flag function begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Details

Note that the number of paired alleles between two populations is not a symmetric dissimilarity measure.

If no x^2 is provided, the function uses the pop(gl) hierarchy to determine pairs of populations, otherwise it runs a single comparison between x and x2.

Hint: in case you want to run comparisons between individuals (assuming individual names are unique), you can simply redefine your population names with your individual names, as below:

pop(gl) <- indNames(gl)</pre>

Definition of fixed and private alleles

The table below shows the possible cases of allele frequencies between two populations (0 = ho-mozygote for Allele 1, x = both Alleles are present, 1 = homozygote for Allele 2).

- p: cases where there is a private allele in pop1 compared to pop2 (but not viceversa)
- f: cases where there is a fixed allele in pop1 (and pop2, as those cases are symmetric)

The absolute allele frequency difference (AFD) in this function is a simple differentiation metric displaying intuitive properties which provides a valuable alternative to FST. For details about its properties and how it is calculated see Berner (2019).

The Bootstrap test for significant differences of private alleles uses a bootstrap simulation by shuffling individuals between a pair of populations and drawing with replacement. For each bootstrap the ratio of private alleles is compared to the actual ratio and recorded how often it is larger than the simulated one. If number of individuals are different between population bootstrap is done using the smaller number of samples in both populations.

The function also reports an estimation of the lower bound of the number of undetected private alleles using the Good-Turing frequency formula, originally developed for cryptography, which estimates in an ecological context the true frequencies of rare species in a single assemblage based on an incomplete sample of individuals. The approach is described in Chao et al. (2017). For this function, the equation 2c is used. This estimate is reported in the output table as Chao1 and Chao2.

In this function a Sankey Diagram is used to visualize patterns of private alleles between populations. This diagram allows to display flows (private alleles) between nodes (populations). Their links are represented with arcs that have a width proportional to the importance of the flow (number of private alleles).

if save2temp=TRUE, resultant plot(s) and the tabulation(s) are saved to the session's temporary directory.

Value

A data.frame. Each row shows, for each pair of populations the number of individuals in each population, the number of loci with fixed differences (same for both populations) in pop1 (compared to pop2) and viceversa. Same for private alleles and finally the absolute mean allele frequency difference between loci (AFD). If loc.names = TRUE, loci names with private alleles and fixed differences are reported in a list in addition to the dataframe.

Author(s)

Custodian: Bernd Gruber – Post to https://groups.google.com/d/forum/dartr

References

- Berner, D. (2019). Allele frequency difference AFD an intuitive alternative to FST for quantifying genetic population differentiation. Genes, 10(4), 308.
- Chao, Anne, et al. "Deciphering the enigma of undetected species, phylogenetic, and functional diversity based on Good-Turing theory." Ecology 98.11 (2017): 2914-2929.

See Also

```
Other matched report: gl.filter.excess.het(), gl.report.allna(), gl.report.callrate(),
gl.report.hamming(), gl.report.locmetric(), gl.report.maf(), gl.report.overshoot(),
gl.report.rdepth(), gl.report.reproducibility(), gl.report.secondaries(), gl.report.taglength()
```

Other report functions: gl.report.replicates()

Examples

out <- gl.report.pa(platypus.gl)</pre>

out <- gl.report.pa(platypus.gl)</pre>

gl.report.polyploid_heterozygosity

Reports observed, expected and unbiased heterozygosities and FIS (inbreeding coefficient) by population or by individual from SNP data

Description

Calculates the observed, expected and unbiased expected (i.e. corrected for sample size) heterozygosities and FIS (inbreeding coefficient) for each population or the observed heterozygosity for each individual in a genlight object.

Usage

```
gl.report.polyploid_heterozygosity(
  х,
 method = "pop",
  n.invariant = 0,
  subsample.pop = FALSE,
  n.limit = 10,
  nboots = 0,
  conf = 0.95,
  CI.type = "bca",
  ncpus = 1,
  plot.display = TRUE,
  plot.theme = theme_dartR(),
  plot.colors.pop = gl.colors("dis"),
  plot.colors.ind = gl.colors(2),
  plot.file = NULL,
  plot.dir = NULL,
  error.bar = "SD",
  verbose = NULL
```

```
)
```

Arguments

х	Name of the genlight object containing the SNP data that converted to dosage mode [required].
method	Calculate heterozygosity by population (method='pop') or by individual (method='ind') [default 'pop'].

n.invariant	An estimate of the number of invariant sequence tags used to adjust the het- erozygosity rate [default 0].
subsample.pop	Whether subsample populations to estimate observed heterozygosity (see De- tails) [default FALSE].
n.limit	Minimum number of individuals that should have a population to perform sub- sampling to estimate heterozygosity [default 10].
nboots	Number of bootstrap replicates to obtain confidence intervals [default 0].
conf	The confidence level of the required interval [default 0.95].
CI.type	Method to estimate confidence intervals. One of "norm", "basic", "perc" or "bca" [default "bca"].
ncpus	Number of processes to be used in parallel operation. If ncpus > 1 parallel operation is activated, see "Details" section [default 1].
plot.display	Specify if plot is to be produced [default TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.colors.po	p
	A color palette for population plots or a list with as many colors as there are populations in the dataset [default gl.colors("dis")].
plot.colors.in	d
	List of two color names for the borders and fill of the plot by individual [default gl.colors(2)].
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL].
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()].
error.bar	Statistic to be plotted as error bar either "SD" (standard deviation) or "SE" (stan- dard error) or "CI" (confidence intervals) [default "SD"].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity].

Details

Observed heterozygosity for a population takes the proportion of heterozygous loci for each individual and averages it over all individuals in that population. The calculations take into account missing values.

Expected heterozygosity for a population takes the expected proportion of heterozygotes, that is, expected under Hardy-Weinberg equilibrium, for each locus, then averages this across the loci for an average estimate for the population.

The unbiased expected heterozygosity is calculated using the correction for sample size following equation 2 from Nei 1978.

Accuracy of all heterozygosity estimates is affected by small sample sizes, and so is their comparison between populations or repeated analysis. Expected heterozygosities are less affected because their calculations are based on allele frequencies while observed heterozygosities are strongly susceptible to sampling effects when the sample size is small. Observed heterozygosity for individuals is calculated as the proportion of heterozygous gametes that could be produced by that individual.

Finally, the loci that are invariant across all individuals in the dataset (that is, across populations), is typically unknown. This can render estimates of heterozygosity analysis specific, and so it is not valid to compare such estimates across species or even across different analyses (see Schimdt et al 2021). This is a similar problem faced by microsatellites. If you have an estimate of the number of invariant sequence tags (loci) in your data, such as provided by gl.report.secondaries, you can specify it with the n.invariant parameter to standardize your estimates of heterozygosity. This is called autosomal heterozygosities by Schimddt et al (2021).

NOTE: It is important to realise that estimation of adjusted (autosomal) heterozygosity requires that secondaries not to be removed.

Heterozygosities and FIS (inbreeding coefficient) are calculated by locus within each population using the following equations, and then averaged across all loci:

- Observed heterozygosity (Ho) = number of heterozygous gametes / all combinations of gametes, where n_Ind is the number of individuals without missing data for that locus.
- Observed heterozygosity adjusted (Ho.adj) <- Ho * n_Loc / (n_Loc + n.invariant), where n_Loc is the number of loci that do not have all missing data and n.invariant is an estimate of the number of invariant loci to adjust heterozygosity.
- Expected heterozygosity (He) = $1 (p^2 + q^2)$, where p is the frequency of the reference allele and q is the frequency of the alternative allele.
- Expected heterozygosity adjusted (He.adj) = He * n_Loc / (n_Loc + n.invariant)
- Unbiased expected heterozygosity (uHe) = He * (2 * n_Ind / (2 * n_Ind 1))
- Inbreeding coefficient (FIS) = 1 Ho / uHe

Function's output

Output for method='pop' is an ordered barchart of observed heterozygosity, unbiased expected heterozygosity and FIS (Inbreeding coefficient) across populations together with a table of mean observed and expected heterozygosities and FIS by population and their respective standard deviations (SD). In the output, it is also reported by population: the number of loci used to estimate heterozygosity (n.Loc), the number of polymorphic loci (polyLoc), the number of monomorphic loci (monoLoc) and loci with all missing data (all_NALoc).

Output for method='ind' is a histogram and a boxplot of heterozygosity across individuals.

If a plot.file is given, the ggplot arising from this function is saved as an "RDS" binary file using saveRDS(); can be reloaded with readRDS(). A file name must be specified for the plot to be saved.

If a plot directory (plot.dir) is specified, the ggplot binary is saved to that directory; otherwise to the tempdir().

Examples of other themes that can be used can be consulted in:

- https://ggplot2.tidyverse.org/reference/ggtheme.html and
- https://yutannihilation.github.io/allYourFigureAreBelongToUs/ggthemes/

Subsampling populations

To test the effect of five population sample sizes (n = 10, 5, 4, 3, 2) on observed heterozygosity estimates, the function subsamples individuals, without replacement. The subsampling is repeated 10 times for each sample size n. This approach is an implementation of Schmidt et al (2021).

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Error bars

The best method for presenting or assessing genetic statistics depends on the type of data you have and the specific questions you're trying to answer. Here's a brief overview of when you might use each method:

1. Confidence Intervals ("CI"):

- Usage: Often used to convey the precision of an estimate.

- Advantage: Confidence intervals give a range in which the true parameter (like a population mean) is likely to fall, given the data and a specified probability (like 95%).

- In Context: For genetic statistics, if you're estimating a parameter, a 95% CI gives you a range in which you're 95% confident the true parameter lies.

2. Standard Deviation ("SD"):

- Usage: Describes the amount of variation from the average in a set of data.

- Advantage: Allows for an understanding of the spread of individual data points around the mean.

- In Context: If you're looking at the distribution of a quantitative trait (like height) in a population with a particular genotype, the SD can describe how much individual heights vary around the average height.

3. Standard Error ("SE"):

- Usage: Describes the precision of the sample mean as an estimate of the population mean.

- Advantage: Smaller than the SD in large samples; it takes into account both the SD and the sample size.

- In Context: If you want to know how accurately your sample mean represents the population mean, you'd look at the SE.

Recommendation:

- If you're trying to convey the precision of an estimate, confidence intervals are very useful.

- For understanding variability within a sample, standard deviation is key.

- To see how well a sample mean might estimate a population mean, consider the standard error.

analyze and present their data, depending on their research questions and the nature of the data.

Confidence Intervals

The uncertainty of a parameter, in this case the mean of the statistic, can be summarised by a confidence interval (CI) which includes the true parameter value with a specified probability (i.e. confidence level; the parameter "conf" in this function).

In this function, CI are obtained using Bootstrap which is an inference method that samples with replacement the data (i.e. loci) and calculates the statistics every time.

This function uses the function boot (package boot) to perform the bootstrap replicates and the function boot.ci (package boot) to perform the calculations for the CI.

Four different types of nonparametric CI can be calculated (parameter "CI.type" in this function):

- First order normal approximation interval ("norm").
- Basic bootstrap interval ("basic").
- Bootstrap percentile interval ("perc").
- Adjusted bootstrap percentile interval ("bca").

The studentized bootstrap interval ("stud") was not included in the CI types because it is computationally intensive, it may produce estimates outside the range of plausible values and it has been found to be erratic in practice, see for example the "Studentized (t) Intervals" section in:

https://www.r-bloggers.com/2019/09/understanding-bootstrap-confidence-interval-output-from-the-r-b

Nice tutorials about the different types of CI can be found in:

https://www.datacamp.com/tutorial/bootstrap-r

and

https://www.r-bloggers.com/2019/09/understanding-bootstrap-confidence-interval-output-from-the-r-b

Efron and Tibshirani (1993, p. 162) and Davison and Hinkley (1997, p. 194) suggest that the number of bootstrap replicates should be between 1000 and 2000.

It is important to note that unreliable confidence intervals will be obtained if too few number of bootstrap replicates are used. Therefore, the function boot.ci will throw warnings and errors if bootstrap replicates are too few. Consider increasing the number of bootstrap replicates to at least 200.

The "bca" interval is often cited as the best for theoretical reasons, however it may produce unstable results if the bootstrap distribution is skewed or has extreme values. For example, you might get the warning "extreme order statistics used as endpoints" or the error "estimated adjustment 'a' is NA". In this case, you may want to use more bootstrap replicates or a different method or check your data for outliers.

The error "estimated adjustment 'w' is infinite" means that the estimated adjustment 'w' for the "bca" interval is infinite, which can happen when the empirical influence values are zero or very close to zero. This can be caused by various reasons, such as:

The number of bootstrap replicates is too small, the statistic of interest is constant or nearly constant across the bootstrap samples, the data contains outliers or extreme values.

You can try some possible solutions, such as:

Increasing the number of bootstrap replicates, using a different type of bootstrap confidence interval or removing or transforming the outliers or extreme values.

Parallelisation

If the parameter ncpus > 1, parallelisation is enabled. In Windows, parallel computing employs a "socket" approach that starts new copies of R on each core. POSIX systems, on the other hand (Mac, Linux, Unix, and BSD), utilise a "forking" approach that replicates the whole current version of R and transfers it to a new core.

Opening and terminating R sessions in each core involves a significant amount of processing time, therefore parallelisation in Windows machines is only quicker than not using parallelisation when nboots > 1000-2000.

Value

A dataframe containing population labels, heterozygosities, FIS, their standard deviations and sample sizes.

Author(s)

Custodian: Ching Lau (Post to https://groups.google.com/d/forum/dartr)

gl.report.rdepth

References

 Moody, M. E., Mueller, L. D., & Soltis, D. E. (1993). Genetic variation and random drift in autotetraploid populations. Genetics, 134(2), 649-657.

See Also

gl.filter.heterozygosity

```
Other unmatched report: gl.allele.freq(), gl.report.basics(), gl.report.diversity(),
gl.report.excess.het(), gl.report.heterozygosity()
```

gl.report.rdepth *Reports summary of Read Depth for each locus*

Description

SNP datasets generated by DArT report AvgCountRef and AvgCountSnp as counts of sequence tags for the reference and alternate alleles respectively. These can be used to back calculate Read Depth. Fragment presence/absence datasets as provided by DArT (SilicoDArT) provide Average Read Depth and Standard Deviation of Read Depth as standard columns in their report. This function reports the read depth by locus for each of several quantiles.

Usage

```
gl.report.rdepth(
    x,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
    plot.dir = NULL,
    plot.file = NULL,
    verbose = NULL
)
```

Arguments

x	Name of the genlight object containing the SNP or presence/absence (Silico-DArT) data [required].
plot.display	Specify if plot is to be produced [default TRUE].
plot.theme	User specified theme [default theme_dartR()].
plot.colors	Vector with two color names for the borders and fill [default c("#2171B5", "#6BAED6")].
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]
plot.file	Filename (minus extension) for the RDS plot file [Required for plot save]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Details

The function displays a table of minimum, maximum, mean and quantiles for read depth against possible thresholds that might subsequently be specified in gl.filter.rdepth. If plot.display=TRUE, display also includes a boxplot and a histogram to guide in the selection of a threshold for filtering on read depth. Plot colours can be set with gl.select.colors(). If plot.file is specified, plots are saved to the directory specified by the user, or the global default working directory set by gl.set.wd() or to the tempdir(). For examples of themes, see

- https://ggplot2.tidyverse.org/reference/ggtheme.html and
- https://yutannihilation.github.io/allYourFigureAreBelongToUs/ggthemes/

Value

An unaltered genlight object

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

gl.filter.rdepth

```
Other matched report: gl.filter.excess.het(), gl.report.allna(), gl.report.callrate(),
gl.report.hamming(), gl.report.locmetric(), gl.report.maf(), gl.report.overshoot(),
gl.report.pa(), gl.report.reproducibility(), gl.report.secondaries(), gl.report.taglength()
```

Examples

SNP data
df <- gl.report.rdepth(testset.gl)
df <- gl.report.rdepth(testset.gs)</pre>

gl.report.replicates Identify replicated individuals

Description

Identify replicated individuals

Usage

```
gl.report.replicates(
    x,
    loc_threshold = 100,
    perc_geno = 0.95,
    plot.out = TRUE,
```

```
plot_theme = theme_dartR(),
plot_colors = c("#2171B5", "#6BAED6"),
bins = 100,
verbose = NULL
```

Arguments

)

x	Name of the genlight object containing the SNP data [required].
loc_threshold	Minimum number of loci required to asses that two individuals are replicates [default 100].
perc_geno	Mimimum percentage of genotypes in which two individuals should be the same [default 0.95].
plot.out	Specify if plot is to be produced [default TRUE].
plot_theme	User specified theme [default theme_dartR()].
plot_colors	Vector with two color names for the borders and fill [default c("#2171B5", "#6BAED6")].
bins	Number of bins to display in histograms [default 100].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Details

This function uses an C++ implementation, so package Rcpp needs to be installed and it is therefore fast (once it has compiled the function after the first run).

Ideally, in a large dataset with related and unrelated individuals and several replicated individuals, such as in a capture/mark/recapture study, the first histogram should have four "peaks". The first peak should represent unrelated individuals, the second peak should correspond to second-degree relationships (such as cousins), the third peak should represent first-degree relationships (like parent/offspring and full siblings), and the fourth peak should represent replicated individuals.

In order to ensure that replicated individuals are properly identified, it's important to have a clear separation between the third and fourth peaks in the second histogram. This means that there should be bins with zero counts between these two peaks.

Value

A list with three elements:

- table.rep: A dataframe with pairwise results of percentage of same genotypes between two individuals, the number of loci used in the comparison and the missing data for each individual.
- ind.list.drop: A vector of replicated individuals to be dropped. Replicated individual with the least missing data is reported.
- ind.list.rep: A list of of each individual that has replicates in the dataset, the name of the replicates and the percentage of the same genotype.

Author(s)

Custodian: Luis Mijangos - Post to https://groups.google.com/d/forum/dartr

See Also

Other report functions: gl.report.pa()

Examples

```
res_rep <- gl.report.replicates(platypus.gl, loc_threshold = 500,
perc_geno = 0.85)
```

gl.report.reproducibility

Reports summary of RepAvg (repeatability averaged over both alleles for each locus) or reproducibility (repeatability of the scores for fragment presence/absence)

Description

SNP datasets generated by DArT have an index, RepAvg, generated by reproducing the data independently for 30 of alleles that give a repeatable result, averaged over both alleles for each locus. In the case of fragment presence/absence data (SilicoDArT), repeatability is the percentage of scores that are repeated in the technical replicate dataset.

Usage

```
gl.report.reproducibility(
    x,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
    plot.dir = NULL,
    plot.file = NULL,
    verbose = NULL
)
```

'

Arguments

x	Name of the genlight object containing the SNP or presence/absence (Silico-DArT) data [required].
plot.display	Specify if plot is to be produced [default TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.colors	Vector with two color names for the borders and fill [default c("#2171B5", "#6BAED6")].

plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]
plot.file	Filename (minus extension) for the RDS plot file [Required for plot save]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Details

The function displays a table of minimum, maximum, mean and quantiles for repeatbility against possible thresholds that might subsequently be specified in gl.filter.reproducibility. If plot.display=TRUE, display also includes a boxplot and a histogram to guide in the selection of a threshold for filtering on repeatability. If plot.file is specified, plots are saved to the directory specified by the user, or the global default working directory set by gl.set.wd() or to the tempdir(). For examples of themes, see:

- https://ggplot2.tidyverse.org/reference/ggtheme.html and
- https://yutannihilation.github.io/allYourFigureAreBelongToUs/ggthemes/

Value

An unaltered genlight object

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

gl.filter.reproducibility

```
Other matched report: gl.filter.excess.het(), gl.report.allna(), gl.report.callrate(),
gl.report.hamming(), gl.report.locmetric(), gl.report.maf(), gl.report.overshoot(),
gl.report.pa(), gl.report.rdepth(), gl.report.secondaries(), gl.report.taglength()
```

Examples

```
# SNP data
    out <- gl.report.reproducibility(testset.gl)</pre>
```

```
# Tag P/A data
out <- gl.report.reproducibility(testset.gs)</pre>
```

gl.report.secondaries Reports loci containing secondary SNPs in sequence tags and calculates number of invariant sites

Description

SNP datasets generated by DArT include fragments with more than one SNP (that is, with secondaries). They are recorded separately with the same CloneID (=AlleleID). These multiple SNP loci within a fragment are likely to be linked, and so you may wish to remove secondaries. This function reports statistics associated with secondaries, and the consequences of filtering them out, and provides three plots. The first is a boxplot, the second is a barplot of the frequency of secondaries per sequence tag, and the third is the Poisson expectation for those frequencies including an estimate of the zero class (no. of sequence tags with no SNP scored).

Usage

```
gl.report.secondaries(
    x,
    nsim = 1000,
    taglength = 69,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
    plot.dir = NULL,
    plot.file = NULL,
    verbose = NULL
)
```

Arguments

х	Name of the genlight object [required].
nsim	The number of simulations to estimate the mean of the Poisson distribution [de-fault 1000].
taglength	Typical length of the sequence tags [default 69].
plot.display	Specify if plot is to be produced [default TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.colors	Vector with two color names for the borders and fill [default c("#2171B5", "#6BAED6")].
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]
plot.file	Filename (minus extension) for the RDS plot file [Required for plot save]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Details

The function gl.filter.secondaries will filter out the loci with secondaries retaining only one sequence tag. Heterozygosity as estimated by the function gl.report.heterozygosity is in a sense relative, because it is calculated against a background of only those loci that are polymorphic somewhere in the dataset. To allow intercompatibility across studies and species, any measure of heterozygosity needs to accommodate loci that are invariant (autosomal heterozygosity. See Schmidt et al 2021). However, the number of invariant loci are unknown given the SNPs are detected as single point mutational variants and invariant sequences are discarded, and because of the particular additional filtering pre-analysis. Modelling the counts of SNPs per sequence tag as a Poisson distribution in this script allows estimate of the zero class, that is, the number of invariant loci. This is reported, and the veracity of the estimate can be assessed by the correspondence of the observed frequencies against those under Poisson expectation in the associated graphs. The number of invariant loci can then be optionally provided to the function gl.report.heterozygosity via the parameter n.invariants. In case the calculations for the Poisson expectation of the number of invariant sequence tags fail to converge, try to rerun the analysis with a larger nsim values. This function now also calculates the number of invariant sites (i.e. nucleotides) of the sequence tags (if TrimmedSequence is present in x\$other\$loc.metrics) or estimate these by assuming that the average length of the sequence tags is 69 nucleotides. Based on the Poisson expectation of the number of invariant sequence tags, it also estimates the number of invariant sites for these to eventually provide an estimate of the total number of invariant sites. Note, previous version of dartR would only return an estimate of the number of invariant sequence tags (not sites). If plot.file is specified, plots are saved to the directory specified by the user, or the global default working directory set by gl.set.wd() or to the tempdir(). Examples of other themes that can be used can be consulted in:

- https://ggplot2.tidyverse.org/reference/ggtheme.html and
- https://yutannihilation.github.io/allYourFigureAreBelongToUs/ggthemes/
- n.total.tags Number of sequence tags in total
- n.SNPs.secondaries Number of secondary SNP loci that would be removed on filtering
- n.invariant.tags Estimated number of invariant sequence tags
- n.tags.secondaries Number of sequence tags with secondaries
- n.inv.gen Number of invariant sites in sequenced tags
- mean.len.tag Mean length of sequence tags
- n.invariant Total Number of invariant sites (including invariant sequence tags)
- k Lambda: mean of the Poisson distribution of number of SNPs in the sequence tags

Value

A data.frame with the list of parameter values

Author(s)

Custodian: Arthur Georges (Post to https://groups.google.com/d/forum/dartr)

References

Schmidt, T.L., Jasper, M.-E., Weeks, A.R., Hoffmann, A.A., 2021. Unbiased population heterozygosity estimates from genome-wide sequence data. Methods in Ecology and Evolution n/a.

See Also

gl.filter.secondaries,gl.report.heterozygosity,utils.n.var.invariant

```
Other matched report: gl.filter.excess.het(), gl.report.allna(), gl.report.callrate(),
gl.report.hamming(), gl.report.locmetric(), gl.report.maf(), gl.report.overshoot(),
gl.report.pa(), gl.report.rdepth(), gl.report.reproducibility(), gl.report.taglength()
```

Examples

```
require("dartR.data")
test <- gl.filter.callrate(platypus.gl,threshold = 1)
n.inv <- gl.report.secondaries(test)
gl.report.heterozygosity(test, n.invariant = n.inv[7, 2])</pre>
```

gl.report.shannon	Report SNP diversity from a genlight object, with reference to Ma,
	Z., Li, L., & Zhang, Y. P. (2020). Defining individual-level genetic
	diversity and similarity profiles. Scientific reports, 10(1), 5805.

Description

This function needs package adegenet, please install it.

Usage

```
gl.report.shannon(
    x,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.dir = NULL,
    plot.file = NULL,
    level = "alpha",
    order = 5,
    verbose = 2
)
```

Arguments

x	A genlight file (works only for diploid data) [required].
plot.display	Specify if plot is to be produced [default TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()].
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL].

level	The types of SNP diversity to report. [default 'alpha', also accept 'beta', 'gamma'].
order	The number of order to report. Starts from 0. [default 5].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress
	and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Details

details

• SNP diversity per individual

Value

A dataframe containing SNP diversity per individual

Author(s)

Ching Ching Lau (Post to https://groups.google.com/d/forum/dartr)

References

• Ma, Z., Li, L., & Zhang, Y. P. (2020). Defining individual-level genetic diversity and similarity profiles. Scientific reports, 10(1), 5805.

Examples

```
## Not run:
obj <- gl.report.shannon(gl)
## End(Not run)
```

gl.report.taglength Reports summary of sequence tag length across loci

Description

SNP datasets generated by DArT typically have sequence tag lengths ranging from 20 to 69 base pairs. This function reports summary statistics of the tag lengths.

```
gl.report.taglength(
    x,
    plot.display = TRUE,
    plot.theme = theme_dartR(),
    plot.colors = NULL,
    plot.file = NULL,
    plot.dir = NULL,
    verbose = NULL
)
```

Arguments

x	Name of the genlight object containing the SNP [required].
plot.display	If TRUE, histograms of base composition are displayed in the plot window [de-fault TRUE].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].
plot.colors	List of two color names for the borders and fill of the plots [default c("#2171B5", "#6BAED6")].
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL]
plot.dir	Directory in which to save files [default = working directory]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity]

Details

This function reports sequence tag lengths for a genlight object. It is a companion function to gl.filter.taglength which can be used to filter out loci with a tag length less than a specified threshold. The table of quantiles is useful for deciding a threshold for subsequent filtering as it provides an indication of the percentages of loci that will be retained and lost. Function's output The minimum, maximum, mean and a tabulation of tag length quantiles against thresholds are output to the console. The output also includes a boxplot and a histogram to guide in the selection of a threshold for filtering on tag length.

If a plot.file is given, the ggplot arising from this function is saved as an "RDS" binary file using saveRDS(); can be reloaded with readRDS(). A file name must be specified for the plot to be saved. If a plot directory (plot.dir) is specified, the ggplot binary is saved to that directory; otherwise to the tempdir().

To avoid issues from inadvertent use of this function in an assignment statement, the function returns the genlight object unaltered.

Value

Returns unaltered genlight object

Author(s)

Author(s): Arthur Georges. Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

gl.filter.taglength

```
Other matched report: gl.filter.excess.het(), gl.report.allna(), gl.report.callrate(),
gl.report.hamming(), gl.report.locmetric(), gl.report.maf(), gl.report.overshoot(),
gl.report.pa(), gl.report.rdepth(), gl.report.reproducibility(), gl.report.secondaries()
```

gl.sample

Examples

out <- gl.report.taglength(testset.gl)</pre>

gl.sample

Samples individuals from populations

Description

A function to subsample individuals in a genlight object

Usage

```
gl.sample(
    x,
    nsample = min(table(pop(x))),
    replace = TRUE,
    onepop = FALSE,
    verbose = NULL
)
```

Arguments

х	genlight object containing SNP/silicodart genotypes
nsample	the number of individuals that should be sampled
replace	a switch to sample by replacement (default).
onepop	switch to ignore population settings of the genlight object and sample from all individuals disregarding the population definition. [default FALSE].
verbose	set verbosity

Details

This is convenience function to facilitate a bootstrap approach

This function is often used to support a bootstrap approach in dartR. For a bootstrap approach it is often desirable to sample a defined number of individuals for each of the populations in a genlight object and then calculate a certain quantity for that subset (redo a 1000 times)

Value

returns a genlight object with nsample samples from each populations.

Author(s)

Bernd Gruber (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other data manipulation: gl.define.pop(), gl.drop.ind(), gl.drop.loc(), gl.drop.pop(),
gl.edit.recode.pop(), gl.impute(), gl.join(), gl.keep.ind(), gl.keep.loc(), gl.keep.pop(),
gl.make.recode.ind(), gl.merge.pop(), gl.reassign.pop(), gl.recode.ind(), gl.recode.pop(),
gl.rename.pop(), gl.sim.genotypes(), gl.sort(), gl.subsample.ind(), gl.subsample.loc()
```

Examples

```
#bootstrap for 2 possums populations to check effect of sample size on fixed alleles
gl.set.verbosity(0)
pp <- possums.gl[c(1:30,91:120),]
nrep <- 1:10
nss <- seq(1,10,2)
res <- expand.grid(nrep=nrep, nss=nss)
for (i in 1:nrow(res)) {
  dummy <- gl.sample(pp, nsample=res$nss[i], replace=TRUE)
  dummy <- gl.compliance.check(dummy)
  pas <- gl.report.pa(dummy, plot.display= FALSE)
  res$fixed[i] <- pas$fixed[1]
  }
  boxplot(fixed ~ nss, data=res)</pre>
```

gl.save

Saves an object in compressed binary format for later rapid retrieval

Description

This is a wrapper for saveRDS(). The script saves the object in binary form to the current workspace and returns the input gl object.

Usage

gl.save(x, file, verbose = NULL)

Arguments

Х	Name of the genlight object containing SNP genotypes [required].
file	Name of the file to receive the binary version of the object [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

The input object

Author(s)

Custodian: Arthur Georges - Post to https://groups.google.com/d/forum/dartr

gl.select.colors

See Also

gl.load

```
Other io: gl.load(), gl.read.csv(), gl.read.dart(), gl.read.fasta(), gl.read.silicodart(),
gl.write.csv(), utils.read.dart()
```

Examples

gl.save(testset.gl,file.path(tempdir(),'testset.rds'))

gl.select.colors Selects colors from one of several palettes and outputs as a vector

Description

This function draws upon a number of specified color libraries to extract a vector of colors for plotting. For use where the function that follows has a color parameter expecting a vector of colors.

Usage

```
gl.select.colors(
  x = NULL,
  library = NULL,
  palette = NULL,
  ncolors = NULL,
  select = NULL,
  plot.display = TRUE,
  verbose = NULL
)
```

Arguments

Х	Optionally, provide a gl object from which to determine the number of popula- tions [default NULL].
library	Name of the color library to be used, one of 'brewer' 'gr.palette', 'r.hcl' or 'baseR' [default scales::hue_pl].
palette	Name of the color palette to be pulled from the specified library, refer function help [default is library specific].
ncolors	number of colors to be displayed and returned [default 9 or nPop(gl)].
select	select bu number the colors to retain in the output vector; can repeat colors. [default NULL].
plot.display	if TRUE, plot the colours in the plot window [default=TRUE]
verbose	- verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Details

Colors are chosen by specifying a library (one of 'brewer' 'gr.palette', 'r.hcl' or 'baseR') and a palette within that library. Each library has its own array of palettes, which can be listed as outlined below. Alternatively, if you specify an incorrect palette, the list of available palettes for the specified library will be listed.

The available color libraries and their palettes include:

- library 'brewer' and the palettes available can be listed by RColorBrewer::display.brewer.all() and RColorBrewer::brewer.pal.info.
- library 'gr.palette' and the palettes available can be listed by grDevices::palette.pals()
- library 'r.hcl' and the palettes available can be listed by grDevices::hcl.pals()
- library 'baseR' and the palettes available are: 'rainbow', 'heat', 'topo.colors', 'terrain.colors', 'cm.colors'.

If the library is not specified, then the default library 'scales' is set and the default palette of 'hue_pal is set.

If the library is set but the palette is not specified, all palettes for that library will be listed and a default palette will then be chosen. The color palette will be displayed in the graphics window for the requested number of colors (or 9 if not specified or nPop(gl) if a genlight object is specified), and the vector of colors returned by assignment for later use. The select parameter can be used to select colors from the specified ncolors. For example, select=c(1,1,3) will select color 1, 1 again and 3 to retain in the final vector. This can be useful for fine-tuning color selection, and matching colors and shapes.

Value

A vector with the required number of colors

Author(s)

Custodian: Arthur Georges - Post to https://groups.google.com/d/forum/dartr

See Also

gl.select.shapes

```
Other graphics: gl.colors(), gl.map.interactive(), gl.plot.heatmap(), gl.report.ld.map(),
gl.select.shapes(), gl.smearplot(), gl.tree.nj()
```

Examples

```
# SET UP DATASET
gl <- testset.gl
levels(pop(gl))<-c(rep('Coast',5),rep('Cooper',3),rep('Coast',5),
rep('MDB',8),rep('Coast',7),'Em.subglobosa','Em.victoriae')
# EXAMPLES -- SIMPLE
colors <- gl.select.colors()
colors <- gl.select.colors(library='brewer',palette='Spectral',ncolors=6)
colors <- gl.select.colors(library='baseR',palette='terrain.colors',ncolors=6)
colors <- gl.select.colors(library='baseR',palette='rainbow',ncolors=12)</pre>
```

gl.select.shapes

```
colors <- gl.select.colors(library='gr.hcl',palette='RdBu',ncolors=12)
colors <- gl.select.colors(library='gr.palette',palette='Pastel 1',ncolors=6)
# EXAMPLES -- SELECTING colorS
colors <- gl.select.colors(library='baseR',palette='rainbow',ncolors=12,select=c(1,1,1,5,8))
# EXAMPLES -- CROSS-CHECKING WITH A GENLIGHT OBJECT
colors <- gl.select.colors(x=gl,library='baseR',palette='rainbow',ncolors=12,select=c(1,1,1,5,8))</pre>
```

gl.select.shapes Selects shapes from the base R shape palette and outputs as a vector

Description

This script draws upon the standard R shape palette to extract a vector of shapes for plotting, where the script that follows has a shape parameter expecting a vector of shapes.

Usage

gl.select.shapes(x = NULL, select = NULL, verbose = NULL)

Arguments

x	Optionally, provide a gl object from which to determine the number of popula- tions [default NULL].
select	Select the shapes to retain in the output vector [default NULL, all shapes shown and returned].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Details

By default the shape palette will be displayed in full in the graphics window from which shapes can be selected in a subsequent run, and the vector of shapes returned for later use. The select parameter can be used to select shapes from the specified 26 shapes available (0-25). For example, select=c(1,1,3) will select shape 1, 1 again and 3 to retain in the final vector. This can be useful for fine-tuning shape selection, and matching colors and shapes.

Value

A vector with the required number of shapes

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

```
gl.select.colors
Other graphics: gl.colors(), gl.map.interactive(), gl.plot.heatmap(), gl.report.ld.map(),
gl.select.colors(), gl.smearplot(), gl.tree.nj()
```

Examples

```
# SET UP DATASET
gl <- testset.gl
levels(pop(gl))<-c(rep('Coast',5),rep('Cooper',3),rep('Coast',5),
rep('MDB',8),rep('Coast',7),'Em.subglobosa','Em.victoriae')
# EXAMPLES
shapes <- gl.select.shapes() # Select and display available shapes
# Select and display a restricted set of shapes
shapes <- gl.select.shapes(select=c(1,1,1,5,8))
# Select set of shapes and check with no. of pops.
shapes <- gl.select.shapes(x=gl,select=c(1,1,1,5,8))</pre>
```

gl.set.verbosity Sets the default verbosity level

Description

dartR functions have a verbosity parameter that sets the level of reporting during the execution of the function. The verbosity level, set by parameter 'verbose' can be one of verbose 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report. The default value for verbosity is stored in the r environment. This script sets the default value.

Usage

gl.set.verbosity(value = 2)

Arguments

value

Set the default verbosity to be this value: 0, silent only fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2]

Value

verbosity value [set for all functions]

Author(s)

Custodian: Arthur Georges (Post to https://groups.google.com/d/forum/dartr)

Examples

gl <- gl.set.verbosity(value=2)</pre>

gl.set.wd

Description

Many dartR functions have a plot.dir parameter which is used to save output to (e.g. ggplots as rds files) With this functions users can set the working directory globally so it is used in all functions, without setting is explicitly. The value for wd is stored in the r environment and if not set defaults to tempdir(). This script sets the default value.

Usage

gl.set.wd(wd = tempdir(), verbose = NULL)

Arguments

wd	Set the path to the wd directory globally to be used by all functions if not set explicitly in the function.
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Value

path the working directory [set for all functions]

Author(s)

Custodian: Bernd Gruber (Post to https://groups.google.com/d/forum/dartr)

See Also

Other environment: gl.check.verbosity(), gl.check.wd(), gl.print.history(), theme_dartR()

Examples

```
#set to current working directory
wd <- gl.set.wd(wd=getwd())</pre>
```

```
gl.sim.cross
```

Description

Generates random crosses between fathers (in one genlight object) and mothers (in a second genlight object) then randomly selects a specified number of offspring to retain.

Usage

```
gl.sim.crosses(
  fathers,
  mothers,
  broodsize = 10,
  sexratio = 0.5,
  n = 1000,
  error.check = TRUE,
  compliance.check = TRUE,
  verbose = NULL
)
```

Arguments

fathers	Genlight object of potential fathers [required].	
mothers	Genlight object of potential mothers simulated [required].	
broodsize	Number of offspring per mother [required].	
sexratio	Sex ratio of simulated offspring [default 0.5].	
n	Number of offspring to retain [default 1000 or mothers*broodsize whichever is the lesser]	
error.check	If TRUE, will perform error checks on the provided parameters [default TRUE]	
compliance.check		
	If TRUE, will perform a compliance check on the resultant genlight object be- fore returning it [default TRUE]	
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity]	

Details

This script is to be used in conjunction with gl.subsample.ind() applied initially to a base genlight object containing initial male and female genotypes. The workflow is to

(a) Select the males from the base genlight object using gl.keep.pop() with pop.list="male" and the as.pop parameter set to sex. (b) Select the females from the base genlight object using gl.keep.pop() with pop.list="female" and the as.pop parameter set to sex. (c) Subsample a cohort of males for breeding and a cohort of females for breeding using gl.subsample.ind() and the replace parameter as follows:

gl.sim.genotypes

To enforce monogamy – generate the fathers and mothers from the base genlight object using gl.subsample.ind() with replace=FALSE. To admit polygyny – generate the fathers from the base genlight object using gl.subsample.ind() with replace=FALSE and the mothers from the base genlight object using gl.subsample.ind() with replace=TRUE. To admit polyandry – generate the fathers from the base genlight object using gl.subsample.ind() with replace=TRUE. To admit polyandry – generate the fathers from the base genlight object using gl.subsample.ind() with replace=TRUE. To admit polyandry – generate the fathers from the base genlight object using gl.subsample.ind() with replace=TRUE and the mothers from the base genlight object using gl.subsample.ind() with replace=FALSE. To admit promiscuity – generate the fathers and mothers from the base genlight object using gl.subsample.ind() with replace=TRUE.

These are simple scenarios that leave the number of maternal mates per father (polygyny) and the number of paternal mates per mother (polyandry) to chance, depending on the random selection of males and females with replacement from the base genlight object.

(d) Cross the males with the females using gl.sim.crosses() retaining a subset of offspring at random.

So the input for this function is a genlight object with a sample of male individuals (fathers) selected from a larger set at random with or without replacement; a similar sample of female individuals in a second genlight object (mothers); specified broodsize; and desired offspring sex ratio.

Set check.error to FALSE if using this script in simulations

Value

A genlight object with n offspring of both sexes.

Author(s)

Custodian: Bernd Gruber (Post to https://groups.google.com/d/forum/dartr)

gl.sim.genotypes Generate random genotypes

Description

Generate random genotypes for a single population drawing upon the allele frequencies from that population.

Usage

```
gl.sim.genotypes(x, n.ind = 200, verbose = NULL)
```

Arguments

x	Name of the genlight object [required].
n.ind	Number of individuals to be simulated (should be less than the number of loci) [default 200]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity]

Value

Returns a genlight object with the simulated genotypes

Author(s)

Custodian: Arthur Georges (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other data manipulation: gl.define.pop(), gl.drop.ind(), gl.drop.loc(), gl.drop.pop(),
gl.edit.recode.pop(), gl.impute(), gl.join(), gl.keep.ind(), gl.keep.loc(), gl.keep.pop(),
gl.make.recode.ind(), gl.merge.pop(), gl.reassign.pop(), gl.recode.ind(), gl.recode.pop(),
gl.rename.pop(), gl.sample(), gl.sort(), gl.subsample.ind(), gl.subsample.loc()
```

gl.smearplot Smear plot

Description

Each locus is color coded for scores of 0, 1, 2 and NA for SNP data and 0, 1 and NA for presence/absence (SilicoDArT) data. Individual labels can be added. Plot may become cluttered if ind.labels If there are too many individuals, it is best to use ind.labels = FALSE.

Works with both SNP data and P/A data (SilicoDArT)

Usage

```
gl.smearplot(
    x,
    plot.display = TRUE,
    ind.labels = FALSE,
    label.size = 10,
    group.pop = FALSE,
    plot.theme = NULL,
    plot.colors = NULL,
    plot.file = NULL,
    het.only = FALSE,
    legend = "bottom",
    verbose = NULL
)
```

Arguments

х	Name of the genlight object [required].
plot.display	If TRUE, the plot is displayed in the plot window [default TRUE].
ind.labels	If TRUE, individual IDs are shown [default FALSE].

gl.sort

label.size	Size of the individual labels [default 10].
group.pop	If ind.labels is TRUE, group by population [default TRUE].
plot.theme	Theme for the plot. See Details for options [default NULL].
plot.colors	List of four color names for the column fill for homozygous reference, heterozy- gous, homozygous alternate, and missing value (NA) [default c("#0000FF", "#00FFFF", "#FF0000", "#e0et
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL]
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]#'
het.only	If TRUE, show only the heterozygous state [default FALSE]
legend	Position of the legend: "left", "top", "right", "bottom" or 'none' [default = 'bot-tom'].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity]

Value

Returns the ggplot object

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

Other graphics: gl.colors(), gl.map.interactive(), gl.plot.heatmap(), gl.report.ld.map(), gl.select.colors(), gl.select.shapes(), gl.tree.nj()

Examples

gl.smearplot(testset.gl,ind.labels=FALSE)
gl.smearplot(testset.gs,ind.labels=FALSE)
gl.smearplot(testset.gl[1:10,],ind.labels=TRUE)
gl.smearplot(testset.gs[1:10,],ind.labels=TRUE)

gl.sort

Sorts genlight objects

Description

This function provides the ability to sort genotypes in a genlight object by individual name or population name.

```
gl.sort(x, sort.by = "pop", order.by = NULL, verbose = NULL)
```

Arguments

х	Genlight object containing SNP/Silicodart genotypes [required].
sort.by	Specify to sort the genotypes by either 'ind', "pop" [default 'pop'].
order.by	Vector used to order genotypes [default NULL]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity]

Details

This is a function to sort genotypes in a genlight object by individual name or population name. This will be useful if you want to visualise the structure across populations (bands) in a gl.smearplot; order of genotypes is important.

The order.by parameter needs to be a vector upon which to effect the sort, of length of nPop(gl) if sort.by is 'pop' or nInd(gl) if sort.by is 'ind'. For sort.by='ind' order.by can be a vector such as a variable in gl@other\$ind.metrics.

If not specified by nominating a vector with order.by, alphabetical order of populations or individuals is used.

Value

Returns a reordered genlight object. Sorts also the ind/loc.metrics and coordinates accordingly

Author(s)

Bernd Gruber (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other data manipulation: gl.define.pop(), gl.drop.ind(), gl.drop.loc(), gl.drop.pop(),
gl.edit.recode.pop(), gl.impute(), gl.join(), gl.keep.ind(), gl.keep.loc(), gl.keep.pop(),
gl.make.recode.ind(), gl.merge.pop(), gl.reassign.pop(), gl.recode.ind(), gl.recode.pop(),
gl.rename.pop(), gl.sample(), gl.sim.genotypes(), gl.subsample.ind(), gl.subsample.loc()
```

Examples

```
#sort by populations
bc <- gl.sort(bandicoot.gl)
#sort from West to East
bc2 <- gl.sort(bandicoot.gl, sort.by="pop" ,
order.by=c("WA", "SA", "VIC", "NSW", "QLD"))
#sort by missing values
miss <- rowSums(is.na(as.matrix(bandicoot.gl)))
bc3 <- gl.sort(bandicoot.gl, sort.by="ind", order.by=miss)
gl.smearplot(bc3)
```

gl.subsample.ind Subsample individuals from a genlight object

Description

A function to subsample individuals at random in a genlight object with and without replacement.

Usage

```
gl.subsample.ind(
    x,
    n = NULL,
    replace = TRUE,
    by.pop = TRUE,
    error.check = TRUE,
    mono.rm = FALSE,
    verbose = NULL
)
```

Arguments

x	Name of the genlight object containing the SNP or presence/absence (Silico-DArT) data [required].
n	Number of individuals to include in the subsample [default NULL]
replace	If TRUE, sampling is with replacement [default TRUE]
by.pop	If FALSE, ignore population settings when subsampling; if TRUE, subsample each population to n individuals [default TRUE].
error.check	If TRUE, will undertake error checks on input paramaters [default TRUE]
mono.rm	If TRUE and error.check is TRUE, monomorphic loci arising from the deletion of individuals will be filtered from the resultant genlight object [default FALSE]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity]

Details

Retain a subset of individuals at random, with or without replacement. If subsampling globally, n must be less than or equal to nInd(x). If subsampling by population, then n must be less than the minimum sample size for any population.

Set error.check = FALSE for speedy execution in simulations

Value

Returns the subsampled genlight object

Author(s)

Custodian: Bernd Gruber (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other data manipulation: gl.define.pop(), gl.drop.ind(), gl.drop.loc(), gl.drop.pop(),
gl.edit.recode.pop(), gl.impute(), gl.join(), gl.keep.ind(), gl.keep.loc(), gl.keep.pop(),
gl.make.recode.ind(), gl.merge.pop(), gl.reassign.pop(), gl.recode.ind(), gl.recode.pop(),
gl.rename.pop(), gl.sample(), gl.sim.genotypes(), gl.sort(), gl.subsample.loc()
```

Examples

```
gl <- gl.subsample.ind(testset.gl, n=30, by.pop=FALSE, replace=TRUE)
gl <- gl.subsample.ind(platypus.gl, n=10, by.pop=TRUE, replace=TRUE)</pre>
```

gl.subsample.loc Subsample loci from a genlight object

Description

A function to subsample loci at random in a genlight object with and without replacement.

Usage

```
gl.subsample.loc(x, n, replace = TRUE, error.check = TRUE, verbose = NULL)
```

Arguments

x	Name of the genlight object containing the SNP or presence/absence (Silico-DArT) data [required].
n	Number of loci to include in the subsample [default NULL]
replace	If TRUE, sampling is with replacement [default TRUE]
error.check	If TRUE, will undertake error checks on input parameters [default TRUE]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity]

Details

Retain a subset of loci at random, with or without replacement. Parameter n must be less than or equal to nLoc(x).

#' Set error.check = FALSE for speedy execution in simulations

Value

Returns the subsampled genlight object

gl.subsample.loci

Author(s)

Custodian: Bernd Gruber (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other data manipulation: gl.define.pop(), gl.drop.ind(), gl.drop.loc(), gl.drop.pop(),
gl.edit.recode.pop(), gl.impute(), gl.join(), gl.keep.ind(), gl.keep.loc(), gl.keep.pop(),
gl.make.recode.ind(), gl.merge.pop(), gl.reassign.pop(), gl.recode.ind(), gl.recode.pop(),
gl.rename.pop(), gl.sample(), gl.sim.genotypes(), gl.sort(), gl.subsample.ind()
```

Examples

gl2 <- gl.subsample.loc(testset.gl, n=50, replace=TRUE, verbose=3)</pre>

gl.subsample.loci	Subsamples n loci from a genlight object and return it as a genlight
	object

Description

This is a support script, to subsample a genlight {adegenet} object based on loci. Two methods are used to subsample, random and based on information content.

Usage

```
gl.subsample.loci(x, n, method = "random", mono.rm = FALSE, verbose = NULL)
```

Arguments

x	Name of the genlight object containing the SNP or presence/absence (Silico-DArT) data [required].
n	Number of loci to include in the subsample [required].
method	Method: 'random', in which case the loci are sampled at random; or 'pic', in which case the top n loci ranked on information content are chosen. Information content is stored in AvgPIC in the case of SNP data and in PIC in the the case of presence/absence (SilicoDArT) data [default 'random'].
mono.rm	Delete monomorphic loci before sampling [default FALSE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

A genlight object with n loci

Author(s)

Custodian: Luis Mijangos - Post to https://groups.google.com/d/forum/dartr

Examples

```
# SNP data
gl2 <- gl.subsample.loci(testset.gl, n=200, method='pic')
# Tag P/A data
gl2 <- gl.subsample.loci(testset.gl, n=100, method='random')</pre>
```

gl.test.heterozygosity

Tests the difference in heterozygosity between populations taken pairwise

Description

Calculates the expected heterozygosities for each population in a genlight object, and uses rerandomization to test the statistical significance of differences in heterozygosity between populations taken pairwise.

Expected heterozygosity is calculated using the correction for sample size following equation 2 from Nei 1978.

Usage

```
gl.test.heterozygosity(
    x,
    nreps = 100,
    alpha1 = 0.05,
    alpha2 = 0.01,
    plot.out = TRUE,
    max_plots = 6,
    plot.theme = theme_dartR(),
    plot.colors = gl.select.colors(ncolors = 2, verbose = 0),
    plot.file = NULL,
    plot.dir = NULL,
    verbose = NULL
)
```

Arguments

х	A genlight object containing the SNP genotypes [required].
nreps	Number of replications of the re-randomization [default 1,000].
alpha1	First significance level for comparison with diff=0 on plot [default 0.05].
alpha2	Second significance level for comparison with diff=0 on plot [default 0.01].
plot.out	If TRUE, plots a sampling distribution of the differences for each comparison [default TRUE].
max_plots	Maximum number of plots to print per page [default 6].
plot.theme	Theme for the plot. See Details for options [default theme_dartR()].

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plot.colors	List of two color names for the borders and fill of the plots [default gl.colors(2)].
plot.file	Name for the RDS binary file to save (base name only, exclude extension) [de-fault NULL]
plot.dir	Directory to save the plot RDS files [default as specified by the global working directory or tempdir()]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity].

Details

Function's output If plot.out = TRUE, plots are created showing the sampling distribution for the difference between each pair of heterozygosities, marked with the critical limits alpha1 and alpha2, the observed heterozygosity, and the zero value (if in range). If a plot.file is given, the ggplot arising from this function is saved as an "RDS" binary file using saveRDS(); can be reloaded with readRDS(). A file name must be specified for the plot to be saved. If a plot directory (plot.dir) is specified, the ggplot binary is saved to that directory; otherwise to the tempdir(). Examples of other themes that can be used can be consulted in

- https://ggplot2.tidyverse.org/reference/ggtheme.html and
- https://yutannihilation.github.io/allYourFigureAreBelongToUs/ggthemes/

Value

A dataframe containing population labels, heterozygosities and sample sizes

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

References

Nei, M. (1978). Estimation of average heterozygosity and genetic distance from a small number of individuals. Genetics, 89(3), 583-590.

Examples

out <- gl.test.heterozygosity(platypus.gl, nreps=1, verbose=3, plot.out=TRUE)</pre>

gl.tree.fitch Generates a distance phylogeny

Description

Generates a distance phylogeny from a distance object using the Fitch-Margoliash algorithm in Phylip.

Usage

```
gl.tree.fitch(
 D,
 x = NULL,
 phylip.path,
 out.path = tempdir(),
  tree.method = "FM",
 outgroup = NULL,
 global.rearrange = FALSE,
 randomize = FALSE,
 n.jumble = 9,
 bstrap = 1,
 plot.type = "phylogram",
 bstrap.threshold = 0.8,
 branch.width = 2,
 branch.color = "blue",
  node.label.color = "red",
  terminal.label.cex = 0.8,
  node.label.cex = 0.8,
 offset = 1.2,
 verbose = NULL
)
```

Arguments

D	Name of the distance matrix for tree building [required]
x	Name of the genlight object containing the SNP data [required for bootstrapping, default NULL].
phylip.path	Path to the directory that holds the Phylip executables [required].
out.path	Path to the directory to save files produced by the analysis [default tempdir()]
tree.method	Algorithm used for constructing trees and selecting the best tree [default "FM"]
outgroup	Name of the outgroup taxon [default NULL, no outgroup, tree not rooted]
global.rearrang	ge
	If TRUE, undertake global rearrangements when generating the tree [default FALSE].
randomize	If TRUE, randomize the order of the input taxa [default FALSE].
n.jumble	Number of randomizations of the input order, must be odd [default 9]
bstrap	Number of bootstrap replicates [default 1000]
plot.type	One of 'phylogram', 'cladogram', 'unrooted', 'fan', 'tidy', 'radial' [default "phylo- gram"]
bstrap.threshold	
	Threshold for bootstrap values to be displayed on the tree [default 0.8]
branch.width	Width of the branches [default 2]
branch.color	Colour of the branches [default "blue"]

gl.tree.fitch

node.label.color	
	Colour of the node labels [default "red"]
terminal.label.cex	
	Height of the taxon label text [default 0.8]
node.label.cex	Height of the node label text [default 0.8]
offset	Horizontal offset of the node labels from the node [default 1.8]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Details

The script takes a distance object as input. This distance object is typically created with gl.dist.phylo(). The script then creates a file consistent with what is expected by program fitch in the Phylip suite of executables. It then runs fitch to generate the "best" phylogenetic tree. Program fitch is run again with bstrap replicates to generate bootstrap support for each node in the tree and plots these on the tree.

tree.method : Currently only Fitch-Margoliash is implemented.

outgroup : Name the taxon to be used as outgroup. Must be among the names of the populations defined in the genlight object.

Value

The tree file in newick format.

Author(s)

Custodian: Arthur Georges - Post to https://groups.google.com/d/forum/dartr

See Also

Other phylogeny: gl.dist.phylo()

Examples

```
## Not run:
tmp <- gl.filter.monomorphs(testset.gl)
D <- gl.dist.phylo(testset.gl,subst.model="F81")
gl.phylip(D=D,x=tmp,phylip.path="D:/workspace/R/phylip-3.695/exe",plot.type="unrooted",
node.label.cex=0.5,terminal.label.cex=0.6,global.rearrange = FALSE, bstrap=10)
```

End(Not run)

gl.tree.nj

Outputs a tree to summarize genetic similarity among populations (e.g. phenogram)

Description

This function is a wrapper for the nj function in package ape and hclust function in stats applied to Euclidean distances calculated from the genlight object.

Usage

```
gl.tree.nj(
    x,
    dist.matrix = NULL,
    method = "nj",
    by.pop = TRUE,
    as.pop = NULL,
    type = "phylogram",
    outgroup = NULL,
    labelsize = 0.7,
    treefile = NULL,
    verbose = NULL
)
```

Arguments

x	Name of the genlight object containing the SNP data [required].
dist.matrix	Distance matrix [default NULL].
method	Clustering method – nj, neighbor-joining tree; UGPMA, UGPMA tree [default 'nj'].
by.pop	If TRUE, populations are the terminal taxa; if FALSE, individuals are the termi- nal taxa [default TRUE]
as.pop	Assign another ind.metric as the population for the purposes of displaying more informative tip labels [default NULL].
type	Type of dendrogram "phylogram" "cladogram" "fan" "unrooted" [default "phylogram"].
outgroup	Vector containing the population names that are the outgroups [default NULL].
labelsize	Size of the labels as a proportion of the graphics default [default 0.7].
treefile	Name of the file for the tree topology using Newick format [default NULL].
verbose	Verbosity: 0, silent, fatal errors only; 1, flag function begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

gl.write.csv

Details

An euclidean distance matrix is calculated by default [dist.matrix = NULL]. Optionally the user can use as input for the tree any other distance matrix using this parameter, see for example the function gl.dist.pop.

Value

A tree file of class phylo.

Author(s)

Custodian: Arthur Georges (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other graphics: gl.colors(), gl.map.interactive(), gl.plot.heatmap(), gl.report.ld.map(),
gl.select.colors(), gl.select.shapes(), gl.smearplot()
```

Examples

```
# SNP data
gl.tree.nj(testset.gl,type='fan')
# Tag P/A data
gl.tree.nj(testset.gs,type='fan')
res <- gl.tree.nj(platypus.gl)</pre>
```

gl.write.csv

Writes out data from a genlight object to csv file

Description

This script writes to file the SNP genotypes with specimens as entities (columns) and loci as attributes (rows). Each row has associated locus metadata. Each column, with header of specimen id, has population in the first row. The data coding differs from the DArT 1row format in that 0 = reference homozygous, 2 = alternate homozygous, 1 = heterozygous, and NA = missing SNP assignment.

```
gl.write.csv(x, outfile = "outfile.csv", outpath = tempdir(), verbose = NULL)
```

Arguments

х	Name of the genlight object containing the SNP data [required].
outfile	File name of the output file (including extension) [default "outfile.csv"].
outpath	Path where to save the output file [default tempdir(), mandated by CRAN]. Use outpath=getwd() or outpath='.' when calling this function to direct output files to your working directory.
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

Saves a genlight object to csv, returns NULL.

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

```
Otherio:gl.load(),gl.read.csv(),gl.read.dart(),gl.read.fasta(),gl.read.silicodart(),
gl.save(),utils.read.dart()
```

Examples

```
# SNP data
gl.write.csv(testset.gl, outfile='SNP_1row.csv')
# Tag P/A data
gl.write.csv(testset.gs, outfile='PA_1row.csv')
```

gl2bayesAss

Converts a genlight object into bayesAss (BA3) input format

Description

This function exports a genlight object into bayesAss format and save it into a file. This function only caters for ploidy=2.

```
gl2bayesAss(
    x,
    ploidy = 2,
    outfile = "gl.BayesAss.txt",
    outpath = NULL,
    verbose = NULL
)
```

gl2bayescan

Arguments

х	Name of the genlight object containing the SNP data [required].
ploidy	Set the ploidy [defaults 2].
outfile	File name of the output file [default 'gl.BayesAss.txt'].
outpath	Path where to save the output file [default global working directory or if not specified, tempdir()].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

returns the input file as data.table

Author(s)

Custodian: Carlo Pacioni (Post to https://groups.google.com/d/forum/dartr)

References

Mussmann S. M., Douglas M. R., Chafin T. K. and Douglas M. E. (2019) BA3-SNPs: Contemporary migration reconfigured in BayesAss for next-generation sequence data. Methods in Ecology and Evolution 10, 1808-1813.

Wilson G. A. and Rannala B. (2003) Bayesian Inference of Recent Migration Rates Using Multilocus Genotypes. Genetics 163, 1177-1191.

See Also

Other linker: gl2bayescan(), gl2bpp(), gl2demerelate(), gl2eigenstrat(), gl2faststructure(), gl2gds(), gl2genalex(), gl2genepop(), gl2geno(), gl2gi(), gl2hiphop(), gl2phylip(), gl2plink(), gl2related(), gl2sa(), gl2structure(), gl2treemix(), gl2vcf()

Examples

```
require("dartR.data")
#only the first 100 due to check time
gl2bayesAss(platypus.gl[,1:100], outpath=tempdir())
```

gl2bayescan

Converts a genlight object into a format suitable for input to Bayescan

Description

The output text file contains the SNP data and relevant BAyescan command lines to guide input.

```
gl2bayescan(x, outfile = "bayescan.txt", outpath = NULL, verbose = NULL)
```

Arguments

x	Name of the genlight object containing the SNP data [required].
outfile	File name of the output file (including extension) [default bayescan.txt].
outpath	Path where to save the output file [default global working directory or if not specified, tempdir()].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

returns no value (i.e. NULL)

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

References

Foll M and OE Gaggiotti (2008) A genome scan method to identify selected loci appropriate for both dominant and codominant markers: A Bayesian perspective. Genetics 180: 977-993.

See Also

```
Other linker: gl2bayesAss(), gl2bpp(), gl2demerelate(), gl2eigenstrat(), gl2faststructure(),
gl2gds(), gl2genalex(), gl2genepop(), gl2geno(), gl2gi(), gl2hiphop(), gl2phylip(),
gl2plink(), gl2related(), gl2sa(), gl2structure(), gl2treemix(), gl2vcf()
```

Examples

out <- gl2bayescan(testset.gl, outpath = tempdir())</pre>

gl2bpp

Converts a genlight object into a format suitable for input to the BPP program

Description

This function generates the sequence alignment file and the Imap file. The control file should produced by the user. If method = 1, heterozygous positions are replaced by standard ambiguity codes. If method = 2, the heterozygous state is resolved by randomly assigning one or the other SNP variant to the individual. Trimmed sequences for which the SNP has been trimmed out, rarely, by adapter mis-identity are deleted. This function requires 'TrimmedSequence' to be among the locus metrics (@other\$loc.metrics) and information of the type of alleles (slot loc.all e.g. 'G/A') and the position of the SNP in slot position of the "genlight" object (see testset.gl@position and testset.gl@loc.all for how to format these slots.)
gl2bpp

Usage

```
gl2bpp(
    x,
    method = 1,
    merge.secondaries = FALSE,
    outfile = "output_bpp.txt",
    imap = "Imap.txt",
    outpath = NULL,
    verbose = NULL
)
```

Arguments

x	Name of the genlight object containing the SNP data [required].
method	One of $1 \mid 2$, see details [default = 1].
merge.secondari	ies
	Logical, if TRUE, secondary loci are merged into a single sequence [default = FALSE].
outfile	Name of the saved sequence alignment file ["output_bpp.txt"].
imap	Name of the saved Imap file ["Imap.txt"].
outpath	Path where to save the output file [default global working directory or if not specified, tempdir()].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Details

It's important to keep in mind that analyses based on coalescent theory, like those done by the programme BPP, are meant to be used with sequence data. In this type of data, large chunks of DNA are sequenced, so when we find polymorphic sites along the sequence, we know they are all on the same chromosome. This kind of data, in which we know which chromosome each allele comes from, is called "phased data." Most data from reduced representation genome-sequencing methods, like DArTseq, is unphased, which means that we don't know which chromosome each allele comes from. So, if we apply coalescence theory to data that is not phased, we will get biased results. As in Ellegren et al., one way to deal with this is to "haplodize" each genotype by randomly choosing one allele from heterozygous genotypes (2012) by using method = 2.

Be mindful that there is little information in the literature on the validity of this method.

Value

```
returns no value (i.e. NULL)
```

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

References

- Ellegren, Hans, et al. "The genomic landscape of species divergence in Ficedula flycatchers." Nature 491.7426 (2012): 756-760.
- Flouri T., Jiao X., Rannala B., Yang Z. (2018) Species Tree Inference with BPP using Genomic Sequences and the Multispecies Coalescent. Molecular Biology and Evolution, 35(10):2585-2593. doi:10.1093/molbev/msy147

See Also

```
Other linker: gl2bayesAss(), gl2bayescan(), gl2demerelate(), gl2eigenstrat(), gl2faststructure(),
gl2gds(), gl2genalex(), gl2genepop(), gl2geno(), gl2gi(), gl2hiphop(), gl2phylip(),
gl2plink(), gl2related(), gl2sa(), gl2structure(), gl2treemix(), gl2vcf()
```

Examples

```
require(dartR.data)
test <- gl.filter.callrate(platypus.gl,threshold = 1)
test <- gl.filter.monomorphs(test)
test <- gl.subsample.loc(test,n=25)
gl2bpp(x = test, outpath=tempdir())</pre>
```

gl2demerelate	Creates a dataframe suitable for input to package {Demerelate} from
	a genlight {adegenet} object

Description

Creates a dataframe suitable for input to package {Demerelate} from a genlight {adegenet} object

Usage

gl2demerelate(x, verbose = NULL)

Arguments

х	Name of the genlight object containing the SNP data [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress
	and results summary; 5, full report [default 2 or as specified using gl.set.verbosity]

Value

A dataframe suitable as input to package {Demerelate}

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

gl2eigenstrat

See Also

```
Other linker: gl2bayesAss(), gl2bayescan(), gl2bpp(), gl2eigenstrat(), gl2faststructure(),
gl2gds(), gl2genalex(), gl2genepop(), gl2geno(), gl2gi(), gl2hiphop(), gl2phylip(),
gl2plink(), gl2related(), gl2sa(), gl2structure(), gl2treemix(), gl2vcf()
```

Examples

```
df <- gl2demerelate(testset.gl)</pre>
```

gl2eigenstrat Converts a genlight object into eigenstrat format

Description

The output of this function are three files:

- genotype file: contains genotype data for each individual at each SNP with an extension 'eigenstratgeno.'
- snp file: contains information about each SNP with an extension 'snp.'
- indiv file: contains information about each individual with an extension 'ind.'

Usage

```
gl2eigenstrat(
    x,
    outfile = "gl_eigenstrat",
    outpath = NULL,
    snp.pos = 1,
    snp.chr = 1,
    pos.cM = 0,
    sex.code = "unknown",
    phen.value = "Case",
    verbose = NULL
)
```

Arguments

x	Name of the genlight object containing the SNP data [required].
outfile	File name of the output file [default 'gl_eigenstrat'].
outpath	Path where to save the output file [default global working directory or if not specified, tempdir()].
snp.pos	Field name from the slot loc.metrics where the SNP position is stored [default 1].
snp.chr	Field name from the slot loc.metrics where the chromosome of each is stored [default 1].

pos.cM	A vector, with as many elements as there are loci, containing the SNP position in morgans or centimorgans [default 1].
sex.code	A vector, with as many elements as there are individuals, containing the sex code ('male', 'female', 'unknown') [default 'unknown'].
phen.value	A vector, with as many elements as there are individuals, containing the pheno- type value ('Case', 'Control') [default 'Case'].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity]

Details

Eigenstrat only accepts chromosomes coded as numeric values, as follows: X chromosome is encoded as 23, Y is encoded as 24, mtDNA is encoded as 90, and XY is encoded as 91. SNPs with illegal chromosome values, such as 0, will be removed.

Value

```
returns no value (i.e. NULL)
```

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

References

- Patterson, N., Price, A. L., & Reich, D. (2006). Population structure and eigenanalysis. PLoS genetics, 2(12), e190.
- Price, A. L., Patterson, N. J., Plenge, R. M., Weinblatt, M. E., Shadick, N. A., & Reich, D. (2006). Principal components analysis corrects for stratification in genome-wide association studies. Nature genetics, 38(8), 904-909.

See Also

```
Other linker: gl2bayesAss(), gl2bayescan(), gl2bpp(), gl2demerelate(), gl2faststructure(),
gl2gds(), gl2genalex(), gl2genepop(), gl2geno(), gl2gi(), gl2hiphop(), gl2phylip(),
gl2plink(), gl2related(), gl2sa(), gl2structure(), gl2treemix(), gl2vcf()
```

Examples

```
require("dartR.data")
gl2eigenstrat(platypus.gl,snp.pos='ChromPos_Platypus_Chrom_NCBIv1',
snp.chr = 'Chrom_Platypus_Chrom_NCBIv1', outpath=tempdir())
```

gl2fasta

Description

Concatenated sequence tags are useful for phylogenetic methods where information on base frequencies and transition and transversion ratios are required (for example, Maximum Likelihood methods). Where relevant, heterozygous loci are resolved before concatenation by either assigning ambiguity codes or by random allele assignment.

Usage

```
gl2fasta(
    x,
    method = 1,
    outfile = "output.fasta",
    outpath = tempdir(),
    probar = FALSE,
    verbose = NULL
)
```

Arguments

х	Name of the genlight object containing the SNP data [required].
method	One of 1 2 3 4. Type method=0 for a list of options [method=1].
outfile	Name of the output file (fasta format) ["output.fasta"].
outpath	Path where to save the output file (set to tempdir by default)
probar	If TRUE, a progress bar will be displayed for long loops [default = TRUE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Details

Four methods are employed:

Method 1 – heterozygous positions are replaced by the standard ambiguity codes. The resultant sequence fragments are concatenated across loci to generate a single combined sequence to be used in subsequent ML phylogenetic analyses.

Method 2 – the heterozygous state is resolved by randomly assigning one or the other SNP variant to the individual. The resultant sequence fragments are concatenated across loci to generate a single composite haplotype to be used in subsequent ML phylogenetic analyses.

Method 3 – heterozygous positions are replaced by the standard ambiguity codes. The resultant SNP bases are concatenated across loci to generate a single combined sequence to be used in subsequent MP phylogenetic analyses.

Method 4 – the heterozygous state is resolved by randomly assigning one or the other SNP variant to the individual. The resultant SNP bases are concatenated across loci to generate a single composite haplotype to be used in subsequent MP phylogenetic analyses.

Trimmed sequences for which the SNP has been trimmed out, rarely, by adapter mis-identity are deleted.

The script writes out the composite haplotypes for each individual as a fastA file. Requires 'Trimmed-Sequence' to be among the locus metrics (@other\$loc.metrics) and information of the type of alleles (slot loc.all e.g. 'G/A') and the position of the SNP in slot position of the "'genlight" object (see testset.gl@position and testset.gl@loc.all for how to format these slots.)

Value

A new gl object with all loci rendered homozygous.

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

gl2faststructure	Converts a genlight object into faststructure format (to run faststruc-
	ture elsewhere)

Description

Recodes in the quite specific faststructure format (e.g first six columns need to be there, but are ignored...check faststructure documentation (if you find any :-())) The script writes out the a file in faststructure format.

Usage

```
gl2faststructure(
    x,
    outfile = "gl.str",
    outpath = NULL,
    probar = FALSE,
    verbose = NULL
)
```

Arguments

х	Name of the genlight object containing the SNP data [required].
outfile	File name of the output file (including extension) [default "gl.str"].
outpath	Path where to save the output file [default global working directory or if not specified, tempdir()].
probar	Switch to show/hide progress bar [default FALSE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

gl2gds

Value

returns no value (i.e. NULL)

Author(s)

Bernd Gruber (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other linker: gl2bayesAss(), gl2bayescan(), gl2bpp(), gl2demerelate(), gl2eigenstrat(),
gl2gds(), gl2genalex(), gl2genepop(), gl2geno(), gl2gi(), gl2hiphop(), gl2phylip(),
gl2plink(), gl2related(), gl2sa(), gl2structure(), gl2treemix(), gl2vcf()
```

gl2gds

Converts a genlight object into gds format

Description

Package SNPRelate relies on a bit-level representation of a SNP dataset that competes with {adegenet} genlight objects and associated files. This function converts a genlight object to a gds format file.

Usage

```
gl2gds(
    x,
    outfile = "gl_gds.gds",
    outpath = NULL,
    snp.pos = "0",
    snp.chr = "0",
    chr.format = "character",
    verbose = NULL
)
```

Arguments

x	Name of the genlight object containing the SNP data [required].
outfile	File name of the output file (including extension) [default 'gl_gds.gds'].
outpath	Path where to save the output file [default global working directory or if not specified, tempdir()].
snp.pos	Field name from the slot loc.metrics where the SNP position is stored [default '0'].
snp.chr	Field name from the slot loc.metrics where the chromosome of each is stored [default '0'].
chr.format	Whether chromosome information is stored as 'numeric' or as 'character', see details [default 'character'].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Details

This function orders the SNPS by chromosome and by position before converting to SNPRelate format, as required by this package. The chromosome of each SNP can be a character or numeric, as described in the vignette of SNPRelate: 'snp.chromosome, an integer or character mapping for each chromosome. Integer: numeric values 1-26, mapped in order from 1-22, 23=X, 24=XY (the pseudoautosomal region), 25=Y, 26=M (the mitochondrial probes), and 0 for probes with unknown positions; it does not allow NA. Character: "X", "XY", "Y" and "M" can be used here, and a blank string indicating unknown position.' When using some functions from package SNPRelate with datasets other than humans it might be necessary to use the option autosome.only=FALSE to avoid detecting chromosome coding. So, it is important to read the documentation of the function before using it. The chromosome information for unmapped SNPS is coded as 0, as required by SNPRelate. Remember to close the GDS file before working in a different GDS object with the function snpgdsClose (package SNPRelate).

Value

returns no value (i.e. NULL)

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other linker: gl2bayesAss(), gl2bayescan(), gl2bpp(), gl2demerelate(), gl2eigenstrat(),
gl2faststructure(), gl2genalex(), gl2genepop(), gl2geno(), gl2gi(), gl2hiphop(), gl2phylip(),
gl2plink(), gl2related(), gl2sa(), gl2structure(), gl2treemix(), gl2vcf()
```

Examples

```
require("dartR.data")
gl2gds(platypus.gl,snp.pos='ChromPos_Platypus_Chrom_NCBIv1',
snp.chr = 'Chrom_Platypus_Chrom_NCBIv1', outpath=tempdir())
```

gl2genalex

Converts a genlight object into a format suitable for input to genalex

Description

The output csv file contains the snp data and other relevant lines suitable for genalex. This function is a wrapper for genind2genalex (package poppr).

gl2genalex

Usage

```
gl2genalex(
    x,
    outfile = "genalex.csv",
    outpath = NULL,
    overwrite = FALSE,
    verbose = NULL
)
```

Arguments

х	Name of the genlight object containing SNP data [required].
outfile	Name of the output file (including extension) [default 'genalex.csv'].
outpath	Path where to save the output file [default global working directory or if not specified, tempdir()].
overwrite	If FALSE and filename exists, then the file will not be overwritten [default FALSE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

returns no value (i.e. NULL)

Author(s)

Custodian: Luis Mijangos, Author: Katrin Hohwieler, wrapper Arthur Georges (Post to https://groups.google.com/d/forum/dartr)

References

Peakall, R. and Smouse P.E. (2012) GenAlEx 6.5: genetic analysis in Excel. Population genetic software for teaching and research-an update. Bioinformatics 28, 2537-2539. http://bioinformatics.oxfordjournals.org/content

See Also

```
Other linker: gl2bayesAss(), gl2bayescan(), gl2bpp(), gl2demerelate(), gl2eigenstrat(),
gl2faststructure(), gl2gds(), gl2genepop(), gl2geno(), gl2gi(), gl2hiphop(), gl2phylip(),
gl2plink(), gl2related(), gl2sa(), gl2structure(), gl2treemix(), gl2vcf()
```

Examples

```
gl2genalex(testset.gl, outfile='testset.csv', outpath=tempdir())
```

gl2genepop

Description

The genepop format is used by several external applications (for example Neestimator2. Unfortunatelly, the software seems to be no longer easily available. To install use the gl.download.binary function. So the main idea is to create the genepop file and then run the other software externally. As a feature, the genepop file is also returned as an invisible data.frame by the function.

Usage

```
gl2genepop(
    x,
    outfile = "genepop.gen",
    outpath = NULL,
    pop.order = "alphabetic",
    output.format = "2_digits",
    verbose = NULL
)
```

Arguments

х	Name of the genlight object containing the SNP data [required].
outfile	File name of the output file [default 'genepop.gen'].
outpath	Path where to save the output file [default global working directory or if not specified, tempdir()].
pop.order	Order of the output populations either "alphabetic" or a vector of population names in the order required by the user (see examples) [default "alphabetic"].
output.format	Whether to use a 2-digit format ("2_digits") or 3-digits format ("3_digits") [default "2_digits"].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Value

Invisible data frame in genepop format

Author(s)

Custodian: Bernd Gruber (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other linker: gl2bayesAss(), gl2bayescan(), gl2bpp(), gl2demerelate(), gl2eigenstrat(),
gl2faststructure(), gl2gds(), gl2genalex(), gl2geno(), gl2gi(), gl2hiphop(), gl2phylip(),
gl2plink(), gl2related(), gl2sa(), gl2structure(), gl2treemix(), gl2vcf()
```

gl2geno

Examples

gl2geno

Converts a genlight object to geno format from package LEA

Description

The function converts a genlight object (SNP or presence/absence i.e. SilicoDArT data) into a file in the 'geno' and the 'lfmm' formats from (package LEA).

The function converts a genlight object (SNP or presence/absence i.e. SilicoDArT data) into a file in the 'geno' and the 'lfmm' formats from (package LEA).

Usage

```
gl2gapit(x, outfile = "gl_gapit", outpath = NULL, verbose = NULL)
gl2geno(x, outfile = "gl_geno", outpath = NULL, verbose = NULL)
```

Arguments

x	Name of the genlight object containing the SNP or presence/absence (Silico-DArT) data [required].
outfile	File name of the output file [default 'gl_geno'].
outpath	Path where to save the output file [default global working directory or if not specified, tempdir()].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Value

returns no value (i.e. NULL) returns no value (i.e. NULL)

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other linker: gl2bayesAss(), gl2bayescan(), gl2bpp(), gl2demerelate(), gl2eigenstrat(),
gl2faststructure(), gl2gds(), gl2genalex(), gl2genepop(), gl2gi(), gl2hiphop(), gl2phylip(),
gl2plink(), gl2related(), gl2sa(), gl2structure(), gl2treemix(), gl2vcf()
```

```
Other linker: gl2bayesAss(), gl2bayescan(), gl2bpp(), gl2demerelate(), gl2eigenstrat(),
gl2faststructure(), gl2gds(), gl2genalex(), gl2genepop(), gl2gi(), gl2hiphop(), gl2phylip(),
gl2plink(), gl2related(), gl2sa(), gl2structure(), gl2treemix(), gl2vcf()
```

Examples

```
# SNP data
t1 <- platypus.gl
# assigning chromosomet1
t1$chromosome <- t1$other$loc.metrics$Chrom_Platypus_Chrom_NCBIv1
# assigning SNP position
t1$position <- t1$other$loc.metrics$ChromPos_Platypus_Chrom_NCBIv1
res <- gl2gapit(t1)
# SNP data
gl2geno(testset.gl, outpath=tempdir())
# Tag P/A data
gl2geno(testset.gs, outpath=tempdir())</pre>
```

gl2gi

Converts a genind object into a genlight object

Description

Converts a genind object into a genlight object Converts a genlight object to genind object

Usage

```
gi2gl(gi, parallel = FALSE, verbose = NULL)
```

gl2gi(x, probar = FALSE, verbose = NULL)

Arguments

gi	A genind object [required].
parallel	Switch to deactivate parallel version. It might not be worth to run it parallel most of the times [default FALSE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].
х	A genlight object [required].
probar	If TRUE, a progress bar will be displayed for long loops [default TRUE].

gl2hiphop

Details

Be aware due to ambiguity which one is the reference allele a combination of gi2gl(gl2gi(gl)) does not return an identical object (but in terms of analysis this conversions are equivalent)

This function uses a faster version of df2genind (from the adegenet package)

Value

A genlight object, with all slots filled.

A genind object, with all slots filled.

Author(s)

Bernd Gruber (Post to https://groups.google.com/d/forum/dartr)

See Also

Other linker: gl2bayesAss(), gl2bayescan(), gl2bpp(), gl2demerelate(), gl2eigenstrat(), gl2faststructure(), gl2gds(), gl2genalex(), gl2genepop(), gl2geno(), gl2hiphop(), gl2phylip(), gl2plink(), gl2related(), gl2sa(), gl2structure(), gl2treemix(), gl2vcf()

Other linker: gl2bayesAss(), gl2bayescan(), gl2bpp(), gl2demerelate(), gl2eigenstrat(), gl2faststructure(), gl2gds(), gl2genalex(), gl2genepop(), gl2geno(), gl2hiphop(), gl2phylip(), gl2plink(), gl2related(), gl2sa(), gl2structure(), gl2treemix(), gl2vcf()

gl2hiphop

Converts a genlight objects into hiphop format

Description

This function exports genlight objects to the format used by the parentage assignment R package hiphop. Hiphop can be used for paternity and maternity assignment and outperforms conventional methods where closely related individuals occur in the pool of possible parents. The method compares the genotypes of offspring with any combination of potentials parents and scores the number of mismatches of these individuals at bi-allelic genetic markers (e.g. Single Nucleotide Polymorphisms).

Usage

```
gl2hiphop(x, verbose = NULL)
```

Arguments

Х	Name of the genlight object containing the SNP data [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress
	and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

Dataframe containing all the genotyped individuals (offspring and potential parents) and their genotypes scored using bi-allelic markers.

Author(s)

Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

References

Cockburn, A., Penalba, J.V., Jaccoud, D., Kilian, A., Brouwer, L., Double, M.C., Margraf, N., Osmond, H.L., van de Pol, M. and Kruuk, L.E.B. (in revision). HIPHOP: improved paternity assignment among close relatives using a simple exclusion method for bi-allelic markers. Molecular Ecology Resources, DOI to be added upon acceptance

See Also

```
Other linker: gl2bayesAss(), gl2bayescan(), gl2bpp(), gl2demerelate(), gl2eigenstrat(),
gl2faststructure(), gl2gds(), gl2genalex(), gl2genepop(), gl2geno(), gl2gi(), gl2phylip(),
gl2plink(), gl2related(), gl2sa(), gl2structure(), gl2treemix(), gl2vcf()
```

Examples

result <- gl2hiphop(testset.gl)</pre>

gl2paup.parsimony	Converts a genlight object to nexus format for parsimony phylogeny analysis in PAUP and, optionally produces accompanying files for parallel processing.
	paraitet processing.

Description

The output nexus file contains the SilicoDArT data as a single line per individual wrapped in the appropriate nexus commands. Pop Labels are used to define taxon partitions.

If out.type="bash", the function produces a series of files in support of an analysis taking advantage of multi-threading and parallel processing.

Usage

```
gl2paup.parsimony(
    x,
    outfileprefix = "parsimony",
    outpath = NULL,
    out.type = "standard",
    tip.labels = "ind",
```

gl2paup.parsimony

```
nreps = 100,
nbootstraps = 1000,
ncpus = 1,
mem = 4,
server = "gadi",
base.dir.name = NULL,
test = FALSE,
verbose = NULL
```

Arguments

х	Name of the genlight object containing the SilicoDArT data [required].
outfileprefix	A prefix to use for file names of the output files [default 'parsimony'].
outpath	Path where to save the output file [default global working directory or if not specified, tempdir()].
out.type	Specify the type of output file. Can be 'standard' (consensus tree) or 'newick' (newick) or 'bash' [default 'standard']
tip.labels	Specify whether the terminals should be labelled with the individual labels ('ind'), the population labels ('pop') or both ('indpop') [default 'ind']
nreps	Specify the number of replicate analyses to run in search of the shortest tree [default 100]
nbootstraps	Number of bootstrap replicates [default 1000]
ncpus	Number of cores to use for parallel processing [default 1]
mem	Memory to use for each process [default 4Gb per core]
server	If out.type='bash', provide the name of the linux server [default 'gadi']
base.dir.name	Name of the base directory on the server to act as the workspace [default NULL]
test	If TRUE, the analysis will run with a small subset of the data [default FALSE]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity]

Details

Additional details: This script only applies to SilicoDArT data. The output file is the name of the file PAUP will use to deliver the results of the analysis, in the directory specified by outpath.

The output type (out.type) can be 'standard' which uses default PAUP parameters to construct the boot.tre file. Or it can be 'newick' to add the parameter format=newick whereby the boot.tre file contains the final tree in newick format. This is useful for passing the results to a tree graphics program such as Mega 11 to format the tree for publication. Or it can be 'bash' which creates a number of files to facilitate parallel processing on a supercomputer.

The parameter nreps specifies the number of replicates to run in search of the shortest tree in each bootstrap iteration. The default is 100.

The parameter nbootstraps specifies the number of bootstrap replicates to run to generate a measure of node support. The default is 1000. The companion parameter ncpus specifies how many cpus to

use for parallel processing when out.type='bash'. The default is 1. Note that the number of cpus must divide evenly into the number of bootstrap replicates.

The parameter tip.labels specifies whether the terminals in the tree should be labelled with the individual names, or the population names (multiple tips will have the same label – which can cause problems at the point of generating a consensus tree), or a combination of the two. Including the population name in the terminal tip labels will assist in collapsing the tree to have populations as the terminals after checking fidelity of populations to supported clades. This can be done in Mega 11.

The parameter 'server' is to allow for future development as users modify the bash scripts to suit other multitasking environments. This script works only for the Gadi server on the Australian National Computing Infrastructure (NCI).

If test=TRUE, the data will be subsetted heavily on numbers of loci, numbers individuals, bootstrap replicates and number of replicates for branch swapping. This is used to test the job run without expenditure of the resources required for the full job.

Value

returns no value (i.e. NULL)

Author(s)

Custodian: Arthur Georges (Post to https://groups.google.com/d/forum/dartr)

See Also

Other linkers: gl2paup.svdquartets()

Examples

```
gg <- testset.gs[1:20,1:100]
gg@other$loc.metrics <- gg@other$loc.metrics[1:100,]
gl2paup.parsimony(gg,outfile="test.nex",outpath=tempdir(),nreps=1,nbootstraps=10)
gl2paup.parsimony(gg,outfile="test.nex",out.type="newick",outpath=tempdir(),nreps=1,nbootstraps=10)</pre>
```

gl2paup.svdquartets Converts a genlight object to nexus format PAUP SVDquartets

Description

The output nexus file contains the SNP data in one of two forms, depending upon what you regard as most appropriate. One form, that used by Chifman and Kubatko, has two lines per individual, one providing the reference SNP the second providing the alternate SNP (method=1). A second form, recommended by Dave Swofford, has a single line per individual, resolving heterozygous SNPs by replacing them with standard ambiguity codes (method=2). If the data are tag presence/absence, then method=2 is assumed.

gl2paup.svdquartets

Usage

```
gl2paup.svdquartets(
    x,
    outfile = "svd.nex",
    outpath = NULL,
    method = 2,
    nbootstraps = 10000,
    verbose = NULL
)
```

Arguments

х	Name of the genlight object containing the SNP data or tag P/A data [required].
outfile	File name of the output file (including extension) [default 'svd.nex'].
outpath	Path where to save the output file [default global working directory or if not specified, tempdir()].
method	Method = 1, nexus file with two lines per individual; method = 2, nexus file with one line per individual, ambiguity codes for SNP genotypes, 0 or 1 for presence/absence data [default 2].
nbootstraps	Number of bootstrap replicates [default 10000]
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity]

Value

returns no value (i.e. NULL)

Author(s)

Custodian: Arthur Georges (Post to https://groups.google.com/d/forum/dartr)

References

Chifman, J. and L. Kubatko. 2014. Quartet inference from SNP data under the coalescent. Bioinformatics 30: 3317-3324

See Also

Other linkers: gl2paup.parsimony()

Examples

```
gg <- testset.gl[1:20,1:100]
gg@other$loc.metrics <- gg@other$loc.metrics[1:100,]
gl2paup.svdquartets(gg, outpath=tempdir(),nbootstraps=100)</pre>
```

gl2phylip

Description

This function calculates and returns a matrix of Euclidean distances between populations and produces an input file for the phylogenetic program Phylip (Joe Felsenstein).

Usage

```
gl2phylip(
    x,
    outfile = "phyinput.txt",
    outpath = tempdir(),
    bstrap = 1,
    verbose = NULL
)
```

Arguments

х	Name of the genlight object containing the SNP or presence/absence (Silico-DArT) data [required].
outfile	Name of the file to become the input file for phylip [default "phyinput.txt"].
outpath	Path where to save the output file [default tempdir(), mandated by CRAN]. Use outpath=getwd() or outpath='.' when calling this function to direct output files to your working directory.
bstrap	Number of bootstrap replicates [default 1].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity]

Value

Matrix of Euclidean distances between populations.

Author(s)

Custodian: Arthur Georges (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other linker: gl2bayesAss(), gl2bayescan(), gl2bpp(), gl2demerelate(), gl2eigenstrat(),
gl2faststructure(), gl2gds(), gl2genalex(), gl2genepop(), gl2geno(), gl2gi(), gl2hiphop(),
gl2plink(), gl2related(), gl2sa(), gl2structure(), gl2treemix(), gl2vcf()
```

gl2plink

Examples

result <- gl2phylip(testset.gl, outfile='test.txt', bstrap=10)</pre>

gl2plink

Converts a genlight object into PLINK format

Description

This function exports a genlight object into PLINK format and save it into a file. This function produces the following PLINK files: bed, bim, fam, ped and map.

Usage

```
gl2plink(
    x,
    plink.bin.path = getwd(),
    bed.files = FALSE,
    outfile = "gl_plink",
    outpath = NULL,
    chr.format = "character",
    pos.cM = "0",
    ID.dad = "0",
    ID.mum = "0",
    sex.code = "unknown",
    phen.value = "0",
    verbose = NULL
)
```

Arguments

Х	Name of the genlight object containing the SNP data [required].
plink.bin.path	Path of PLINK binary file [default getwd()].
bed.files	Whether create PLINK files .bed, .bim and .fam [default FALSE].
outfile	File name of the output file [default 'gl_plink'].
outpath	Path where to save the output file [default global working directory or if not specified, tempdir()].
chr.format	Whether chromosome information is stored as 'numeric' or as 'character', see details [default 'character'].
pos.cM	A vector, with as many elements as there are loci, containing the SNP position in morgans or centimorgans [default '0'].
ID.dad	A vector, with as many elements as there are individuals, containing the ID of the father, '0' if father isn't in dataset [default '0'].

ID.mum	A vector, with as many elements as there are individuals, containing the ID of the mother, '0' if mother isn't in dataset [default '0'].
sex.code	A vector, with as many elements as there are individuals, containing the sex code ('male', 'female', 'unknown'). Sex information needs just to start with an "F" or "f" for females, with an "M" or "m" for males and with a "U", "u" or being empty if the sex is unknown [default 'unknown'].
phen.value	A vector, with as many elements as there are individuals, containing the pheno- type value. '1' = control, '2' = case, '0' = unknown [default '0'].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity]

Details

To create PLINK files .bed, .bim and .fam (bed.files = TRUE), it is necessary to download the binary file of PLINK 1.9 and provide its path (plink.bin.path). The binary file can be downloaded from: https://www.cog-genomics.org/plink/ After downloading, unzip the file, access the unzipped folder and move the binary file ("plink") to your working directory. If you are using a Mac, you might need to open the binary first to grant access to the binary. The chromosome of each SNP can be a character or numeric. The chromosome information for unmapped SNPS is coded as 0. Family ID is taken from x\$pop. Within-family ID (cannot be '0') is taken from indNames(x). Variant identifier is taken from locNames(x). SNP position is taken from the accessor x\$position. Chromosome name is taken from the accessor x\$chromosome Note that if names of populations or individuals contain spaces, they are replaced by an underscore "_". If you like to use chromosome information when converting to plink format and your chromosome names are not from human, you need to change the chromosome names as 'contig1', 'contig2', etc. as described in the section "Nonstandard chromosome IDs" in the following link: https://www.cog-genomics.org/plink/1.9/input

Value

```
returns no value (i.e. NULL)
```

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

References

Purcell, Shaun, et al. 'PLINK: a tool set for whole-genome association and population-based linkage analyses.' The American journal of human genetics 81.3 (2007): 559-575.

See Also

```
Other linker: gl2bayesAss(), gl2bayescan(), gl2bpp(), gl2demerelate(), gl2eigenstrat(),
gl2faststructure(), gl2gds(), gl2genalex(), gl2genepop(), gl2geno(), gl2gi(), gl2hiphop(),
gl2phylip(), gl2related(), gl2sa(), gl2structure(), gl2treemix(), gl2vcf()
```

gl2related

Examples

```
require("dartR.data")
test <- platypus.gl
# assigning SNP position
test$position <- test$other$loc.metrics$ChromPos_Platypus_Chrom_NCBIv1
# assigning a dummy name for chromosomes
test$chromosome <- as.factor("1")
gl2plink(test, outpath=tempdir())</pre>
```

gl2related

Converts a genlight object to format suitable to be run with Coancestry

Description

The output txt file contains the SNP data and an additional column with the names of the individual. The file then can be used and loaded into coancestry or - if installed - run with the related package. Be aware the related package was crashing in previous versions, but in general is using the same code as coancestry and therefore should have identical results. Also running coancestry with thousands of SNPs via the GUI seems to be not reliable and therefore for comparisons between coancestry and related we suggest to use the command line version of coancestry.

Usage

```
gl2related(
    x,
    outfile = "related.txt",
    outpath = NULL,
    save = TRUE,
    verbose = NULL
)
```

Arguments

x	Name of the genlight object containing the SNP data [required].
outfile	File name of the output file (including extension) [default 'related.txt'].
outpath	Path where to save the output file [default global working directory or if not specified, tempdir()].
save	A switch if you want to save the file or not. This might be useful for someone who wants to use the coancestry function to calculate relatedness and not export to coancestry. See the example below [default TRUE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2, unless specified using gl.set.verbosity].

Value

A data.frame that can be used to run with the related package

Author(s)

Bernd Gruber (bugs? Post to https://groups.google.com/d/forum/dartr)

References

Jack Pew, Jinliang Wang, Paul Muir and Tim Frasier (2014). related: related: an R package for analyzing pairwise relatedness data based on codominant molecular markers. R package version 0.8/r2. https://R-Forge.R-project.org/projects/related/

See Also

```
Other linker: gl2bayesAss(), gl2bayescan(), gl2bpp(), gl2demerelate(), gl2eigenstrat(),
gl2faststructure(), gl2gds(), gl2genalex(), gl2genepop(), gl2geno(), gl2gi(), gl2hiphop(),
gl2phylip(), gl2plink(), gl2sa(), gl2structure(), gl2treemix(), gl2vcf()
```

Examples

```
gtd <- gl2related(bandicoot.gl[1:10,1:20], save=FALSE, )
## Not run:
##running with the related package, use
#install.packages('related', repos='http://R-Forge.R-project.org')
library(related)
coan <- coancestry(gtd, wang=1)
head(coan$relatedness)
##check ?coancestry for information how to use the function.</pre>
```

End(Not run)

gl2sa

Converts genlight objects to the format used in the SNPassoc package

Description

This function exports a genlight object into a SNPassoc object. See package SNPassoc for details. This function needs package SNPassoc. At the time of writing (August 2020) the package was no longer available from CRAN. To install the package check their github repository. https://github.com/isglobal-brge/SNPassoc and/or use install_github('isglobal-brge/SNPassoc') to install the function and uncomment the function code.

Usage

gl2sa(x, installed = FALSE, verbose = NULL)

gl2snapper

Arguments

х	Name of the genlight object containing the SNP data [required].
installed	Switch to run the function once SNPassoc package i s installed [default FALSE].#' @param verbose Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

Returns an object of class 'snp' to be used with SNPassoc.

Author(s)

Bernd Guber (Post to https://groups.google.com/d/forum/dartr)

References

Gonzalez, J.R., Armengol, L., Sol?, X., Guin?, E., Mercader, J.M., Estivill, X. and Moreno, V. (2017). SNPassoc: an R package to perform whole genome association studies. Bioinformatics 23:654-655.

See Also

```
Other linker: gl2bayesAss(), gl2bayescan(), gl2bpp(), gl2demerelate(), gl2eigenstrat(),
gl2faststructure(), gl2gds(), gl2genalex(), gl2genepop(), gl2geno(), gl2gi(), gl2hiphop(),
gl2phylip(), gl2plink(), gl2related(), gl2structure(), gl2treemix(), gl2vcf()
```

gl2snapper	Converts a genlight object to nexus format suitable for phylogenetic
	analysis by Snapper (via BEAUti) @family linker

Description

Produces a nexus file contains the SNP calls and relevant PAUP command lines suitable for for the software package BEAUti.

Usage

```
gl2snapper(
    x,
    outfile = "snapper.nex",
    rm.autapomorphies = FALSE,
    nloc = NULL,
    outpath = NULL,
    verbose = NULL
)
```

Arguments

х	Name of the genlight object containing the SNP data [required].	
outfile	Name of the output file (including extension) [default "snapper.nex"].	
rm.autapomorphi	ies	
	Prune the loci by removing autapomorphies (not phylogentically informative), that is, SNP polymorphisms limited to only one population [default TRUE].	
nloc	Number of loci to subsample to bring down computational time [default NULL]	
outpath	Path where to save the output file [default global working directory or if not specified, tempdir()].	
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].	

Details

Snapper is a phylogenetic approach implemented in Beauti/BEAST that can handle larger SNP datasets than can SNAPP. This script produces a nexus file for Beauti which allows options to be set and creates the xml file for BEAST.

Although improved over SNAPP in terms of computational efficiency, Snapper remains constrained by computational times, even when implemented on high performance computers with parallel processing. Computation time of Snapper is not particularly sensitive to the number of individuals in each terminal taxon (= population), but is impacted by the number of populations and the numbers of loci.

Particular attention needs to be directed at amalgamating populations where their joint taxonomic identity is without question and reducing the number of SNP loci by prudent filtering. Removing monomorphic loci, increasing the reliability of loci (read depth, reproducibility), minimizing missing data (call rate), removing multiple SNPs in a single sequence tag (secondaries) are all good options. You may wish also to remove SNP loci that are polymorphic within only a single population (autapomorphies) are all options for reducing the number of loci (rm.autapomorphies=TRUE)

If computational time is still an issue (say requiring one month for a single run), following strategy is recommended. First, subsample the loci to say 100 and test the process to ensure there are no syntax or other issues (nloc=100). You do not want to wait several days or weeks running the full dataset to discover a simple syntax error or incorrectly specified parameter. Second consider running BEAST on a platform that allows multi-threading as this will dramatically reduce compute time. Note that adding threads does not always improve computational time. The optimal number of threads depends on the particular analysis. This means you have to experiment to find out how many threads give the best performance foa a computer cycle.

Also, there is an overheads cost in using many threads. Instead, you could run independent snapper analyses and combine resulting log and tree files. For example, if the optimal number of threads is 8 (adding more threads reduces speed), but 8 threads gives marginal improvement over 4 threads, you can run 2 chains with 4 threads each instead and (after getting through burn-in) then combine results and get a better result than running a single chain at 8 threads. A bonus benefit from running multiple chains is that you can verify that the MCMC ends up with the same posterior distribution each time.

If computational time is still an issue, run the #'analysis on a series of subsamples of loci (say nloc=300) to see if a consistent topology is obtained, then adopt that topology as the final result.

gl2snapper

Note that there is a cost to manipulating your data to achieve reasonable computation times. Omission of sequence tags that are invariant during the SNP calling process, removal of monomorphic loci generated during taxon selection, and removing autapomorphic loci will all affect branch lengths, perhaps differentially, and so compromise branch lengths and estimates of divergence times. Fortunately, the topology should be little affected.

Finally, the analysis relies on certain assumptions. First is that the structure is one of a bifurcating tree and not a network. One needs to assign individuals to populations in advance of the analysis, confident that they are discrete entities and free of horizontal transfer. A second assumption is that the loci scored for SNPs are assorting independently. This is probably a reasonable assumption for SNPs derived from sparse representational sampling (e.g. DArT), but if dense SNP arrays are being used, then some form of thinning will be required. Of course, multiple SNPs on a single sequence tag will be linked, so filtering all but one SNP per sequence tag is required (gl.filter.secondaries).

The workflow is

- "1" Execute gl2snapper()
- "2" Install beast2
- "3" Run beauti in the beast2 bin
- "4" Set the template to snapper [File | Template | Snapper]
- "5" Load the nexus file produced by gl2snapper()
- "6" Select and set the parameters you consider appropriate
- "7" Save the xml file [File | Save As]
- "8" Run beast
- "9" Load the xml file and execute
- "10" When beast is finished, examine the diagnostics with Tracer
- "11" Visualize the resultant trees using DensiTree and FigTree.

If using the command line to run beast, the command is beast -threads myxmlfile. Progress can be monitored with awk (awk '{print \$1, \$2}' snapper.log ltail). When beast is finished, transfer the log and tree files to a windows platform and use Tracer, DensiTree and FigTree as above.

gl2snapper does not work with SilicoDArT data.

Value

```
returns no value (i.e. NULL)
```

Author(s)

Custodian: Arthur Georges (Post to https://groups.google.com/d/forum/dartr)

References

Bryant, D., Bouckaert, R., Felsenstein, J., Rosenberg, N.A. and RoyChoudhury, A. (2012). Inferring species trees directly from biallelic genetic markers: bypassing gene trees in a full coalescent analysis. Molecular Biology and Evolution 29:1917-1932.

Rambaut A, Drummond AJ, Xie D, Baele G and Suchard MA (2018) Posterior summarisation in Bayesian phylogenetics using Tracer 1.7. Systematic Biology. syy032. doi:10.1093/sysbio/syy032

Examples

```
x <- gl.filter.monomorphs(testset.gl)
gl2snapper(x, outfile="test.nex", outpath=tempdir())</pre>
```

gl2structure

Converts a genlight object to STRUCTURE formatted files

Description

This function exports genlight objects to STRUCTURE formatted files (be aware there is a gl2faststructure version as well). It is based on the code provided by Lindsay Clark (see https://github.com/lvclark/R_genetics_conv) and this function is basically a wrapper around her numeric2structure function. See also: Lindsay Clark. (2017, August 22). lvclark/R_genetics_conv: R_genetics_conv 1.1 (Version v1.1). Zenodo: doi.org/10.5281/zenodo.846816.

Usage

```
gl2structure(
    x,
    ind.names = NULL,
    add.columns = NULL,
    ploidy = 2,
    export.marker.names = TRUE,
    outfile = "gl.str",
    outpath = NULL,
    verbose = NULL
)
```

Arguments

x	Name of the genlight object containing the SNP data and location data, lat longs [required].	
ind.names	Specify individuals names to be added [if NULL, defaults to ind.names(x)].	
add.columns	Additional columns to be added before genotypes [default NULL].	
ploidy	Set the ploidy [defaults 2].	
export.marker.names		
	If TRUE, locus names locNames(x) will be included [default TRUE].	
outfile	File name of the output file (including extension) [default "gl.str"].	
outpath	Path where to save the output file [default global working directory or if not specified, tempdir()].	
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].	

gl2treemix

Value

returns no value (i.e. NULL)

Author(s)

Bernd Gruber (wrapper) and Lindsay V. Clark [lvclark@illinois.edu]; Custodian Bernd Gruber

See Also

```
Other linker: gl2bayesAss(), gl2bayescan(), gl2bpp(), gl2demerelate(), gl2eigenstrat(),
gl2faststructure(), gl2gds(), gl2genalex(), gl2genepop(), gl2geno(), gl2gi(), gl2hiphop(),
gl2phylip(), gl2plink(), gl2related(), gl2sa(), gl2treemix(), gl2vcf()
```

Examples

gl2structure(testset.gl[1:10,1:50], outpath=tempdir())

gl2treemix

Converts a genlight object to a treemix input file

Description

The output file contains the SNP data in the format expected by treemix – see the treemix manual. The file will be gzipped before in order to be recognised by treemix. Plotting functions provided with treemix will need to be sourced from the treemix download page.

Usage

```
gl2treemix(x, outfile = "treemix_input.gz", outpath = NULL, verbose = NULL)
```

Arguments

х	Name of the genlight object [required].
outfile	File name of the output file (including gz extension) [default 'treemix_input.gz'].
outpath	Path where to save the output file [default global working directory or if not specified, tempdir()].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

returns no value (i.e. NULL)

gl2vcf

Author(s)

Custodian: Arthur Georges (Post to https://groups.google.com/d/forum/dartr)

References

Pickrell and Pritchard (2012). Inference of population splits and mixtures from genome-wide allele frequency data. PLoS Genetics https://doi.org/10.1371/journal.pgen.1002967

See Also

```
Other linker: gl2bayesAss(), gl2bayescan(), gl2bpp(), gl2demerelate(), gl2eigenstrat(),
gl2faststructure(), gl2gds(), gl2genalex(), gl2genepop(), gl2geno(), gl2gi(), gl2hiphop(),
gl2phylip(), gl2plink(), gl2related(), gl2sa(), gl2structure(), gl2vcf()
```

Examples

```
gl2treemix(testset.gl, outpath=tempdir())
```

gl2vcf

```
Converts a genlight object into vcf format
```

Description

This function exports a genlight object into VCF format and save it into a file.

Usage

```
gl2vcf(
    x,
    plink.bin.path = getwd(),
    outfile = "gl_vcf",
    outpath = NULL,
    snp.pos = "0",
    snp.chr = "0",
    chr.format = "character",
    pos.cM = "0",
    ID.dad = "0",
    ID.mum = "0",
    sex.code = "unknown",
    phen.value = "0",
    verbose = NULL
)
```

gl2vcf

Arguments

x	Name of the genlight object containing the SNP data [required].
plink.bin.path	Path of PLINK binary file [default getwd())].
outfile	File name of the output file [default 'gl_vcf'].
outpath	Path where to save the output file [default global working directory or if not specified, tempdir()].
snp.pos	Field name from the slot loc.metrics where the SNP position is stored [default '0'].
snp.chr	Field name from the slot loc.metrics where the chromosome of each is stored [default '0'].
chr.format	Whether chromosome information is stored as 'numeric' or as 'character', see details [default 'character'].
pos.cM	A vector, with as many elements as there are loci, containing the SNP position in morgans or centimorgans [default '0'].
ID.dad	A vector, with as many elements as there are individuals, containing the ID of the father, '0' if father isn't in dataset [default '0'].
ID.mum	A vector, with as many elements as there are individuals, containing the ID of the mother, '0' if mother isn't in dataset [default '0'].
sex.code	A vector, with as many elements as there are individuals, containing the sex code ('male', 'female', 'unknown') [default 'unknown'].
phen.value	A vector, with as many elements as there are individuals, containing the pheno- type value. '1' = control, '2' = case, '0' = unknown [default '0'].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Details

This function requires to download the binary file of PLINK 1.9 and provide its path (plink.bin.path). The binary file can be downloaded from: https://www.cog-genomics.org/plink/ The chromosome information for unmapped SNPS is coded as 0. Family ID is taken from x\$pop Within-family ID (cannot be '0') is taken from indNames(x) Variant identifier is taken from locNames(x)

Value

```
returns no value (i.e. NULL)
```

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

References

Danecek, P., Auton, A., Abecasis, G., Albers, C. A., Banks, E., DePristo, M. A., ... & 1000 Genomes Project Analysis Group. (2011). The variant call format and VCFtools. Bioinformatics, 27(15), 2156-2158.

See Also

```
Other linker: gl2bayesAss(), gl2bayescan(), gl2bpp(), gl2demerelate(), gl2eigenstrat(),
gl2faststructure(), gl2gds(), gl2genalex(), gl2genepop(), gl2geno(), gl2gi(), gl2hiphop(),
gl2phylip(), gl2plink(), gl2related(), gl2sa(), gl2structure(), gl2treemix()
```

Examples

```
## Not run:
#this example needs plink installed to work
require("dartR.data")
gl2vcf(platypus.gl,snp.pos='ChromPos_Platypus_Chrom_NCBIv1',
    snp.chr = 'Chrom_Platypus_Chrom_NCBIv1')
```

End(Not run)

rbind.dartR adjust rbind for dartR

Description

rbind is a bit lazy and does not take care for the metadata (so data in the other slot is lost). You can get most of the loci metadata back using gl.compliance.check.

Usage

```
## S3 method for class 'dartR'
rbind(...)
```

Arguments

... list of dartR objects

Value

A genlight object

Examples

```
t1 <- platypus.gl
class(t1) <- "dartR"
t2 <- rbind(t1[1:5,],t1[6:10,])</pre>
```

theme_dartR

Description

This is the theme used as default for dartR plots. This function controls all non-data display elements in the plots.

Usage

```
theme_dartR(
   base_size = 11,
   base_family = "",
   base_line_size = base_size/22,
   base_rect_size = base_size/22
)
```

Arguments

base_size	base font size, given in pts.
base_family	base font family
<pre>base_line_size</pre>	base size for line elements
<pre>base_rect_size</pre>	base size for rect elements

Value

a the standard dartR theme to be used in ggplots

See Also

Other environment: gl.check.verbosity(), gl.check.wd(), gl.print.history(), gl.set.wd()

Examples

```
ggplot(data.frame(dummy=rnorm(1000)),aes(dummy)) +
geom_histogram(binwidth=0.1) + theme_dartR()
```

utils.basic.stats Calculates mean observed heterozygosity, mean expected heterozygosity and FIS per locus, per population and various population differentiation measures @family utilities

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

```
utils.basic.stats(x)
```

Arguments

Х

A genlight object containing the SNP genotypes [required].

Details

This is a re-implementation of hierfstat::basics.stats specifically for genlight objects. Formula (and hence results) match exactly the original version of hierfstat::basics.stats but it is much faster.

Value

A list with with the statistics for each population

Author(s)

Luis Mijangos and Carlo Pacioni (post to https://groups.google.com/d/forum/dartr)

Examples

```
require("dartR.data")
out <- utils.basic.stats(platypus.gl)</pre>
```

utils.check.datatype Utility function to check the class of an object passed to a function

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

```
utils.check.datatype(
    x,
    accept = c("genlight", "SNP", "SilicoDArT", "dartR"),
    verbose = NULL
)
```

Arguments

x	Name of the genlight object, dist matrix, data matrix, glPCA, or fixed difference list (fd) [required].
accept	Vector containing the classes of objects that are to be accepted [default c('genlight','SNP','SilicoDArT'].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity].

Details

Most functions require access to a genlight object, dist matrix, data matrix or fixed difference list (fd), and this function checks that a genlight object or one of the above has been passed, whether the genlight object is a SNP dataset or a SilicoDArT object, and reports back if verbosity is >=2.

This function checks the class of passed object and sets the datatype to 'SNP', 'SilicoDArT', 'dist', 'mat', or class[1](x) as appropriate. Note also that this function checks to see if there are individuals or loci scored as all missing (NA) and if so, issues the user with a warning. Note: One and only one of gl.check, fd.check, dist.check or mat.check can be TRUE.

Value

datatype, 'SNP' for SNP data, 'SilicoDArT' for P/A data, 'dist' for a distance matrix, 'mat' for a data matrix, 'glPCA' for an ordination file, or class(x)[1].

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

```
Other utilities: gl.alf(), utils.collapse.matrix(), utils.dart2genlight(), utils.dist.binary(),
utils.flag.start(), utils.hamming(), utils.het.pop(), utils.impute, utils.is.fixed(),
utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(),
utils.read.ped(), utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(),
utils.recalc.freqhomref(), utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.reset.flags(),
utils.transpose(), utils.vcfr2genlight.polyploid()
```

Examples

```
datatype <- utils.check.datatype(testset.gl)
datatype <- utils.check.datatype(as.matrix(testset.gl),accept='matrix')
fd <- gl.fixed.diff(testset.gl)
datatype <- utils.check.datatype(fd,accept='fd')</pre>
```

utils.collapse.matrix Collapses a distance matrix calculated for individuals to a distance matrix for populations defined in a dartR genlight object

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

```
utils.collapse.matrix(D, x, verbose = NULL)
```

Arguments

D	Name of the matrix containing the distances between individuals [required].
х	Name of the genlight object containing the genotypes [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2].
	@details This script takes a matrix of distances calculated between individuals and collapses it by averaging to a matrix of distances between populations. The script gl.dist.ind has a lot of options for distances for presence absence data, but gl.dist.pop does not. This script allows efficient and consistent transfer of this capability to gl.dist.pop.

Value

An object of class 'dist' or 'matrix' giving distances between individuals

Author(s)

Author: Arthur Georges. Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

```
Other utilities: gl.alf(), utils.check.datatype(), utils.dart2genlight(), utils.dist.binary(),
utils.flag.start(), utils.hamming(), utils.het.pop(), utils.impute, utils.is.fixed(),
utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(),
utils.read.ped(), utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(),
utils.recalc.freqhomref(), utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.reset.flags(),
utils.transpose(), utils.vcfr2genlight.polyploid()
```

utils.dart2genlight An internal function to converts DarT to genlight.

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

```
utils.dart2genlight(
   dart,
   ind.metafile = NULL,
   covfilename = NULL,
   probar = TRUE,
   verbose = NULL
)
```

Arguments

dart	A dart object created via read.dart [required].
ind.metafile	Optional file in csv format with metadata for each individual (see details for explanation) [default NULL].
covfilename	Depreciated, use parameter ind.metafile.
probar	Show progress bar [default TRUE].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL].

Details

Converts a DArT file (read via read.dart) into an genlight object from package adegenet. #' Internal function called by gl.read.dart().

The ind.metadata file needs to have very specific headings. First a heading called id. Here the ids have to match the ids in the dart object colnames(dart[[4]]). The following column headings are optional. pop: specifies the population membership of each individual. lat and lon specify spatial coordinates (in decimal degrees WGS1984 format). Additional columns with individual metadata can be imported (e.g. age, gender).

Value

A genlight object. Including all available slots are filled. loc.names, ind.names, pop, lat, lon (if provided via the ind.metadata file)

Author(s)

Maintainer: Bernd Gruber (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dist.binary(),
utils.flag.start(), utils.hamming(), utils.het.pop(), utils.impute, utils.is.fixed(),
utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(),
utils.read.ped(), utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(),
utils.recalc.freqhomref(), utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.reset.flags(),
utils.transpose(), utils.vcfr2genlight.polyploid()
```

utils.dist.binary Calculates a distance matrix for individuals defined in a dartR genlight object using binary P/A data (SilicoDArT)

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

```
utils.dist.binary(
    x,
    method = "simple",
    scale = FALSE,
    swap = FALSE,
    type = "dist",
    verbose = NULL
)
```
Arguments

x	Name of the genlight containing the genotypes [required].
method	Specify distance measure [default simple].
scale	If TRUE and method='euclidean', the distance will be scaled to fall in the range [0,1] [default FALSE].
swap	If TRUE and working with presence-absence data, then presence (no disrupt- ing mutation) is scored as 0 and absence (presence of a disrupting mutation) is scored as 1 [default FALSE].
type	Specify the format and class of the object to be returned, dist for N11 object of class dist, matrix for an object of class matrix [default "dist"].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2].
	@details This script calculates various distances between individuals based on sequence tag Presence/Absence data.
	The distance measure can be one of:
	• Euclidean – Euclidean Distance applied to cartesian coordinates defined by the loci, scored as 0 or 1. Presence and absence equally weighted.
	• simple – simple matching, both 1 or both 0 = 0; one 1 and the other 0 = 1. Presence and absence equally weighted.
	• Jaccard – ignores matching 0, both 1 = 0; one 1 and the other 0 = 1. Absences could be for different reasons.
	• Bray-Curtis – both 0 = 0; both 1 = 2; one 1 and the other 0 = 1. Absences could be for different reasons. Sometimes called the Dice or Sorensen distance.
	One might choose to disregard or downweight absences in comparison with presences because the homology of absences is less clear (mutation at one or the other, or both restriction sites). Your call.

Value

An object of class 'dist' or 'matrix' giving distances between individuals

Author(s)

Author: Arthur Georges. Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.flag.start(), utils.hamming(), utils.het.pop(), utils.impute, utils.is.fixed(),
utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(),
utils.read.ped(), utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(),
utils.recalc.freqhomref(), utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.reset.flags(),
utils.transpose(), utils.vcfr2genlight.polyploid()
```

utils.dist.ind.snp

Calculates a distance matrix for individuals defined in a genlight object using SNP data (DArTseq)

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

```
utils.dist.ind.snp(
    x,
    method = "Euclidean",
    scale = FALSE,
    type = "dist",
    verbose = NULL
)
```

Arguments

x	Name of the genlight containing the genotypes [required].
method	Specify distance measure [default Euclidean].
scale	If TRUE and method='Euclidean', the distance will be scaled to fall in the range [0,1] [default FALSE].
type	Specify the format and class of the object to be returned, dist for a object of class dist, matrix for an object of class matrix [default "dist"].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2].

Details

This script calculates various distances between individuals based on SNP genotypes. The distance measure can be one of:

- Euclidean Euclidean Distance applied to Cartesian coordinates defined by the loci, scored as 0, 1 or 2.
- Simple simple mismatch, 0 where no alleles are shared, 1 where one allele is shared, 2 where both alleles are shared.
- Absolute absolute mismatch, 0 where no alleles are shared, 1 where one or both alleles are shared.
- Czekanowski (or Manhattan) calculates the city block metric distance by summing the scores on each axis (locus).

utils.flag.start

Value

An object of class 'dist' or 'matrix' giving distances between individuals

Author(s)

Author(s): Arthur Georges. Custodian: Arthur Georges – Post to https://groups.google.com/ d/forum/dartr

See Also

Other distance: gl.dist.ind(), gl.dist.pop(), gl.fdsim()

utils.flag.start *A utility script to flag the start of a script*

Description

A utility script to flag the start of a script

Usage

utils.flag.start(func = NULL, build = NULL, verbose = NULL)

Arguments

func	Name of the function that is starting [required].
build	Name of the build [default NULL].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity]

Value

calling function name

Author(s)

Custodian: Arthur Georges – Post to https://groups.google.com/d/forum/dartr

See Also

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.dist.binary(), utils.hamming(), utils.het.pop(), utils.impute, utils.is.fixed(),
utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(),
utils.read.ped(), utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(),
utils.recalc.freqhomref(), utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.reset.flags(),
utils.transpose(), utils.vcfr2genlight.polyploid()
```

utils.hamming

Calculates the Hamming distance between two DArT trimmed DNA sequences

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES. The algorithm is that of Johann de Jong https://johanndejong. wordpress.com/2015/10/02/faster-hamming-distance-in-r-2/

Usage

utils.hamming(str1, str2, r = 4)

Arguments

str1	String containing the first sequence [required].
str2	String containing the second sequence [required].
r	Number of bases in the restriction enzyme recognition sequence [default 4].

Details

Hamming distance is calculated as the number of base differences between two sequences which can be expressed as a count or a proportion. Typically, it is calculated between two sequences of equal length. In the context of DArT trimmed sequences, which differ in length but which are anchored to the left by the restriction enzyme recognition sequence, it is sensible to compare the two trimmed sequences starting from immediately after the common recognition sequence and terminating at the last base of the shorter sequence. The Hamming distance between the rows of a matrix can be computed quickly by exploiting the fact that the dot product of two binary vectors x and (1-y) counts the corresponding elements that are different between x and y. This matrix multiplication can also be used for matrices with more than two possible values, and different types of elements, such as DNA sequences. The function calculates the Hamming distance between all columns of a matrix X, or two matrices X and Y. Again matrix multiplication is used, this time for counting, between two columns x and y, the number of cases in which corresponding elements have the same value (e.g. A, C, G or T). This counting is done for each of the possible values individually, while iteratively adding the results. The end result of the iterative adding is the sum of all corresponding elements that are the same, i.e. the inverse of the Hamming distance. Therefore, the last step is to subtract this end result H from the maximum possible distance, which is the number of rows of matrix X. If the two DNA sequences are of differing length, the longer is truncated. The initial common restriction enzyme recognition sequence is ignored.

Value

Hamming distance between the two strings

utils.het.pop

Author(s)

Custodian: Arthur Georges (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.dist.binary(), utils.flag.start(), utils.het.pop(), utils.impute, utils.is.fixed(),
utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(),
utils.read.ped(), utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(),
utils.recalc.freqhomref(), utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.reset.flags(),
utils.transpose(), utils.vcfr2genlight.polyploid()
```

utils.het.pop	An internal function that calculates expected mean heterozygosity per
	population

Description

An internal function that calculates expected mean heterozygosity per population

Usage

ind.count(x)

Arguments

х

A genlight object containing the SNP genotypes [required].

Value

A vector with the mean expected heterozygosity for each population

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.dist.binary(), utils.flag.start(), utils.hamming(), utils.impute, utils.is.fixed(),
utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(),
utils.read.ped(), utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(),
utils.recalc.freqhomref(), utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.reset.flags(),
utils.transpose(), utils.vcfr2genlight.polyploid()
```

utils.impute

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

matrix2gen(snp_matrix, parallel = FALSE)

Arguments

<pre>snp_matrix</pre>	[Custodian to provide parameter description]
parallel	[Custodian to provide parameter description]

Details

#[Custodian to provide details for future you]

Value

The resultant genlight object

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.dist.binary(), utils.flag.start(), utils.hamming(), utils.het.pop(), utils.is.fixed(),
utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(),
utils.read.ped(), utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(),
utils.recalc.freqhomref(), utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.reset.flags(),
utils.transpose(), utils.vcfr2genlight.polyploid()
```

utils.is.fixed

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

```
utils.is.fixed(s1, s2, tloc = 0)
```

Arguments

s1	Percentage SNP allele or sequence tag frequency for the first population [re- quired].
s2	Percentage SNP allele or sequence tag frequency for the second population [re- quired].
tloc	Threshold value for tolerance in when a difference is regarded as fixed [default 0].

Details

This script compares two percent allele frequencies and reports TRUE if they represent a fixed difference, FALSE otherwise.

A fixed difference at a locus occurs when two populations share no alleles, noting that SNPs are biallelic (ploidy=2). Tolerance in the definition of a fixed difference is provided by the t parameter. For example, t=0.05 means that SNP allele frequencies of 95,5 and 5,95 percent will be reported as fixed (TRUE).

Value

TRUE (fixed difference) or FALSE (alleles shared) or NA (one or both s1 or s2 missing)

Author(s)

Maintainer: Arthur Georges (Post to https://groups.google.com/d/forum/dartr)

See Also

gl.fixed.diff

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.dist.binary(), utils.flag.start(), utils.hamming(), utils.het.pop(), utils.impute,
utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(),
utils.read.ped(), utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(),
```

utils.recalc.freqhomref(),utils.recalc.freqhomsnp(),utils.recalc.maf(),utils.reset.flags(), utils.transpose(),utils.vcfr2genlight.polyploid()

utils.jackknife An internal function to conducts jackknife resampling using a genlight object

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

```
utils.jackknife(
    x,
    FUN,
    unit = "loc",
    recalc = FALSE,
    mono.rm = FALSE,
    n.cores = 1,
    verbose = NULL,
    ...
)
```

Arguments

x	Name of the genlight object [required].
FUN	the name of the function to be used to calculate the statistic
unit	The unit to use for resampling. One of c("loc", "ind", "pop"): loci, individuals or populations
recalc	If TRUE, recalculate the locus metadata statistics [default FALSE].
mono.rm	If TRUE, remove monomorphic and all NA loci [default FALSE].
n.cores	The number of cores to use. If "auto", it will use all but one available cores.
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress but not results; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].
	any additional arguments to be passed to FUN

Details

Jackknife resampling is a statistical procedure where for a dataset of sample size n, subsamples of size n-1 are used to compute a statistic. The collection of the values obtained can be used to evaluate the variability around the point estimate. This function can take the loci, the individuals or the populations as units over which to conduct resampling. Note: when n is very small, jackknife resampling is not recommended. Parallel computation is implemented. The argument n.cores indicates the number of core to use. If "auto", it will use all but one available cores. If the number of units is small (e.g. a few populations), there is not real advantage in using parallel computation. On the other hand, if the number of units is large (e.g. thousands of loci), even with parallel computation, this function can be very slow.

Value

A list of length n where each element is the output of FUN

Author(s)

Custodian: Carlo Pacioni – Post to https://groups.google.com/d/forum/dartr

See Also

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.dist.binary(), utils.flag.start(), utils.hamming(), utils.het.pop(), utils.impute,
utils.is.fixed(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(), utils.read.ped(),
utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(), utils.recalc.freqhomref(),
utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.reset.flags(), utils.transpose(),
utils.vcfr2genlight.polyploid()
```

Examples

```
require("dartR.data")
platMod.gl <- gl.filter.allna(platypus.gl)
chk.pop <- utils.jackknife(x=platMod.gl, FUN="gl.alf", unit="pop",
recalc = FALSE, mono.rm = FALSE, n.cores = 1, verbose=0)</pre>
```

utils.n.var.invariant An internal utility function to calculate the number of variant and invariant sites by locus

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

utils.n.var.invariant(x, verbose = NULL)

Arguments

x	Name of the genlight object containing the SNP data [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL].
	@details Calculate the number of variant and invariant sites by locus and add them as columns in loc.metrics. This can be useful to conduct further fil- tering, for example where only loci with secondaries are wanted for phyloge- netic analyses. Invariant sites are the sites (nucleotide) that are not polymorphic. When the locus metadata supplied by DArT includes the sequence of the allele (TrimmedSequence), it is used by this function to estimate the number of sites that were sequenced in each tag (read). This script then subtracts the number of polymorphic sites. The length of the trimmed sequence (lenTrimSeq), the number of variant (n.variant) and invariant (n.invariant) sites are the added to the table in gl@others\$loc.metrics. NOTE: It is important to realise that this function correctly estimates the number of variant and invariant sites only when it is executed on genlight objects before secondaries are removed.

Value

The modified genlight object.

Author(s)

Custodian: Carlo Pacioni (Post to https://groups.google.com/d/forum/dartr)

See Also

gl.filter.secondaries,gl.report.heterozygosity

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.dist.binary(), utils.flag.start(), utils.hamming(), utils.het.pop(), utils.impute,
utils.is.fixed(), utils.jackknife(), utils.plot.save(), utils.read.fasta(), utils.read.ped(),
utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(), utils.recalc.freqhomref(),
utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.reset.flags(), utils.transpose(),
utils.vcfr2genlight.polyploid()
```

utils.plink.run Runs PLINK from within R

Description

Runs PLINK from within R.

utils.plink.run

Usage

```
utils.plink.run(
  dir.in,
  plink.cmd = "plink",
  plink.path = "path",
  out = "hapmap1",
  syntax,
  verbose = NULL
)
```

Arguments

The path where the data files are
The 'name' to call plink. This will depend on the file name (without the extension '.exe' if on windows) or the name of the PATH variable
The path where the executable is. If plink is listed in the PATH then there is no need for this. This is what the option "path" means
The root of the output file name
the flags to pass to plink call
Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL].

Details

PLINK needs to be installed on the machine and syntax used need to be appropriate for the version installed.

Value

A character vector with the command used for PLINK.

Author(s)

Custodian: Carlo Pacioni and Luis Mijangos (Post to https://groups.google.com/d/forum/ dartr)

References

Purcell, Shaun, et al. 'PLINK: a tool set for whole-genome association and population-based linkage analyses.' The American journal of human genetics 81.3 (2007): 559-575. utils.plot.save

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

```
utils.plot.save(x, dir = NULL, file = NULL, verbose = NULL, ...)
```

Arguments

х	Name of the ggplot object.
dir	Name of the directory to save the file.
file	Name of the file to save the plot to (omit file extension)
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL, unless specified using gl.set.verbosity]
	Parameters passed to function ggsave, such as width and height, when the ggplot is to be saved.

Details

An internal function to save a ggplot object to disk in RDS binary format. Uses saveRDS() to save the file with an .RDS extension; can be reloaded with gl.load().

Value

returns NULL

Author(s)

Custodian: Arthur Georges (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.dist.binary(), utils.flag.start(), utils.hamming(), utils.het.pop(), utils.impute,
utils.is.fixed(), utils.jackknife(), utils.n.var.invariant(), utils.read.fasta(), utils.read.ped(),
utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(), utils.recalc.freqhomref(),
utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.reset.flags(), utils.transpose(),
utils.vcfr2genlight.polyploid()
```

utils.read.dart

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

```
utils.read.dart(
  filename,
  nas = "-",
  topskip = NULL,
  lastmetric = "RepAvg",
  service.row = 1,
  plate.row = 3,
  verbose = NULL
)
```

Arguments

filename	Path to file (csv file only currently) [required].
nas	A character specifying NAs [default '-'].
topskip	A number specifying the number of rows to be skipped. If not provided the number of rows to be skipped are 'guessed' by the number of rows with '*' at the beginning [default NULL].
lastmetric	Specifies the last non genetic column [default 'RepAvg']. Be sure to check if that is true, otherwise the number of individuals will not match. You can also specify the last column by a number.
service.row	The row number in which the information of the DArT service is contained [default 1].
plate.row	The row number in which the information of the plate location is contained [default 3].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default NULL].

Details

Internal function called by gl.read.dart()

Value

A list of length 5. #dart format (one or two rows) #individuals, #snps, #non genetic metrics, #genetic data (still two line format, rows=snps, columns=individuals)

Author(s)

Custodian: Bernd Gruber (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Otherio:gl.load(),gl.read.csv(),gl.read.dart(),gl.read.fasta(),gl.read.silicodart(),
gl.save(),gl.write.csv()
```

utils.read.fasta An internal script to read a fastA file into a genlight object

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

```
utils.read.fasta(file, parallel = parallel, n.cores = NULL, verbose = verbose)
```

Arguments

file	Name of the fastA file [required]
parallel	Switch to deactivate parallel version. It might not be worth to run it parallel most of the times [default FALSE]
n.cores	Number of cores to use in parallel [default 4]
verbose	Verbosity: 0, silent, fatal errors only; 1, flag function begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity].

Value

The resultant genlight object

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.dist.binary(), utils.flag.start(), utils.hamming(), utils.het.pop(), utils.impute,
utils.is.fixed(), utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.ped(),
utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(), utils.recalc.freqhomref(),
utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.reset.flags(), utils.transpose(),
utils.vcfr2genlight.polyploid()
```

utils.read.ped

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

```
utils.read.ped(
  file,
  snps,
  which,
  split = "\t| +",
  sep = ".",
  na.strings = "0",
  lex.order = FALSE,
  show_warnings = TRUE
)
```

Arguments

file	Custodian to provide
snps	Custodian to provide
which	Custodian to provide
split	Custodian to provide
sep	Custodian to provide
na.strings	Custodian to provide
lex.order	Custodian to provide
show_warnings	Custodian to provide
	#[Custodian to provide other parameter descriptions]

Details

#[Custodian to provide details]

Value

The resultant genlight object

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

See Also

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.dist.binary(), utils.flag.start(), utils.hamming(), utils.het.pop(), utils.impute,
utils.is.fixed(), utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(),
utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(), utils.recalc.freqhomref(),
utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.reset.flags(), utils.transpose(),
utils.vcfr2genlight.polyploid()
```

utils.recalc.avgpic A utility function to recalculate intermediate locus metrics

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

utils.recalc.avgpic(x, verbose = NULL)

Arguments

х	Name of the genlight [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress
	and results summary; 5, full report [default 2 or as specified using gl.set.verbosity]

Details

Recalculates OneRatioRef, OneRatioSnp, PICRef, PICSnp, and AvgPIC by locus after some individuals or populations have been deleted.

The locus metadata supplied by DArT has OneRatioRef, OneRatioSnp, PICRef, PICSnp, and Avg-PIC included, but the allelic composition will change when some individuals,or populations, are removed from the dataset and so the initial statistics will no longer apply. This script recalculates these statistics and places the recalculated values in the appropriate place in the genlight object. If the locus metadata OneRatioReflSnp, PICReflSnp and/or AvgPIC do not exist, the script creates and populates them.

Value

The modified genlight object.

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

utils.recalc.callrate

See Also

utils.recalc.metrics for recalculating all metrics, utils.recalc.callrate for recalculating CallRate, utils.recalc.freqhomref for recalculating frequency of homozygous reference, utils.recalc.freqhomsnp for recalculating frequency of homozygous alternate, utils.recalc.freqhet for recalculating frequency of heterozygotes, gl.recalc.maf for recalculating minor allele frequency, gl.recalc.rdepth for recalculating average read depth

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.dist.binary(), utils.flag.start(), utils.hamming(), utils.het.pop(), utils.impute,
utils.is.fixed(), utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(),
utils.read.ped(), utils.recalc.callrate(), utils.recalc.freqhets(), utils.recalc.freqhomref(),
utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.reset.flags(), utils.transpose(),
utils.vcfr2genlight.polyploid()
```

utils.recalc.callrate A utility script to recalculate the callrate by locus after some populations have been deleted

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

utils.recalc.callrate(x, verbose = NULL)

Arguments

х	Name of the genlight object containing the SNP data [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity]

Details

SNP datasets generated by DArT have missing values primarily arising from failure to call a SNP because of a mutation at one or both of the restriction enzyme recognition sites. The locus metadata supplied by DArT has callrate included, but the call rate will change when some individuals are removed from the dataset. This script recalculates the callrate and places these recalculated values in the appropriate place in the genlight object. It sets the Call Rate flag to TRUE.

Value

The modified genlight object

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

See Also

utils.recalc.metrics for recalculating all metrics, utils.recalc.avgpic for recalculating avg-PIC, utils.recalc.freqhomref for recalculating frequency of homozygous reference, utils.recalc.freqhomsnp for recalculating frequency of homozygous alternate, utils.recalc.freqhet for recalculating frequency of heterozygotes, gl.recalc.maf for recalculating minor allele frequency, gl.recalc.rdepth for recalculating average read depth

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.dist.binary(), utils.flag.start(), utils.hamming(), utils.het.pop(), utils.impute,
utils.is.fixed(), utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(),
utils.read.ped(), utils.recalc.avgpic(), utils.recalc.freqhets(), utils.recalc.freqhomref(),
utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.reset.flags(), utils.transpose(),
utils.vcfr2genlight.polyploid()
```

utils.recalc.freqhets A utility script to recalculate the frequency of the heterozygous SNPs by locus after some populations have been deleted

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

```
utils.recalc.freqhets(x, verbose = NULL)
```

Arguments

х	Name of the genlight object containing the SNP data [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity]

Details

The locus metadata supplied by DArT has FreqHets included, but the frequency of the heterozygotes will change when some individuals are removed from the dataset. This script recalculates the FreqHets and places these recalculated values in the appropriate place in the genlight object. Note that the frequency of the homozygote reference SNPS is calculated from the individuals that could be scored.

Value

The modified genlight object.

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

See Also

utils.recalc.metrics for recalculating all metrics, utils.recalc.callrate for recalculating CallRate, utils.recalc.freqhomref for recalculating frequency of homozygous reference, utils.recalc.freqhomsnp for recalculating frequency of homozygous alternate, utils.recalc.AvgPIC for recalculating RepAvg, gl.recalc.maf for recalculating minor allele frequency, gl.recalc.rdepth for recalculating average read depth

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.dist.binary(), utils.flag.start(), utils.hamming(), utils.het.pop(), utils.impute,
utils.is.fixed(), utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(),
utils.read.ped(), utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhomref(),
utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.reset.flags(), utils.transpose(),
utils.vcfr2genlight.polyploid()
```

utils.recalc.freqhomref

#' An internal utility function to recalculate the frequency of the homozygous reference SNP by locus after some populations have been deleted

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

utils.recalc.freqhomref(x, verbose = NULL)

Arguments

х	Name of the genlight [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress
	and results summary; 5, full report [default 2 or as specified using gl.set.verbosity]

Details

The locus metadata supplied by DArT has FreqHomRef included, but the frequency of the homozygous reference will change when some individuals are removed from the dataset. This script recalculates the FreqHomRef and places these recalculated values in the appropriate place in the genlight object. Note that the frequency of the homozygote reference SNPS is calculated from the individuals that could be scored.

Value

The modified genlight object

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

See Also

utils.recalc.metrics for recalculating all metrics, utils.recalc.callrate for recalculating CallRate, utils.recalc.avgpic for recalculating AvgPIC, utils.recalc.freqhomsnp for recalculating frequency of homozygous alternate, utils.recalc.freqhet for recalculating frequency of heterozygotes, gl.recalc.maf for recalculating minor allele frequency, gl.recalc.rdepth for recalculating average read depth

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.dist.binary(), utils.flag.start(), utils.hamming(), utils.het.pop(), utils.impute,
utils.is.fixed(), utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(),
utils.read.ped(), utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(),
utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.reset.flags(), utils.transpose(),
utils.vcfr2genlight.polyploid()
```

utils.recalc.freqhomsnp

A utility function to recalculate the frequency of the homozygous alternate SNP by locus after some populations have been deleted

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

utils.recalc.freqhomsnp(x, verbose = NULL)

Arguments

x	Name of the genlight object [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress
	and results summary; 5, full report [default 2 or as specified using gl.set.verbosity]

Details

The locus metadata supplied by DArT has FreqHomSnp included, but the frequency of the homozygous alternate will change when some individuals are removed from the dataset. This function recalculates the FreqHomSnp and places these recalculated values in the appropriate place in the genlight object. Note that the frequency of the homozygote alternate SNPS is calculated from the individuals that could be scored. This function only applies to SNP genotype data not Tag P/A data (SilicoDArT).

Value

The modified genlight object.

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

See Also

utils.recalc.metrics for recalculating all metrics, utils.recalc.callrate for recalculating CallRate, utils.recalc.freqhomref for recalculating frequency of homozygous reference, utils.recalc.avgpic for recalculating AvgPIC, utils.recalc.freqhet for recalculating frequency of heterozygotes, gl.recalc.maf for recalculating minor allele frequency, gl.recalc.rdepth for recalculating average read depth

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.dist.binary(), utils.flag.start(), utils.hamming(), utils.het.pop(), utils.impute,
utils.is.fixed(), utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(),
utils.read.ped(), utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(),
utils.recalc.freqhomref(), utils.recalc.maf(), utils.reset.flags(), utils.transpose(),
utils.vcfr2genlight.polyploid()
```

utils.recalc.maf

A utility function to recalculate the minor allele frequency by locus, typically after some populations have been deleted

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

utils.recalc.maf(x, verbose = NULL)

Arguments

х	Name of the genlight object [required].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress
	and results summary; 5, full report [default 2 or as specified using gl.set.verbosity]

Details

The locus metadata supplied by DArT does not have MAF included, so it is calculated and added to the locus.metadata by this script. The minimum allele frequency will change when some individuals are removed from the dataset. This script recalculates the MAF and places these recalculated values in the appropriate place in the genlight object. This function only applies to SNP genotype data.

Value

The modified genlight dataset.

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

See Also

utils.recalc.metrics for recalculating all metrics, utils.recalc.callrate for recalculating CallRate, utils.recalc.freqhomref for recalculating frequency of homozygous reference, utils.recalc.freqhomsnp for recalculating frequency of homozygous alternate, utils.recalc.freqhet for recalculating frequency of heterozygotes, gl.recalc.avgpic for recalculating AvgPIC, gl.recalc.rdepth for recalculating average read depth

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.dist.binary(), utils.flag.start(), utils.hamming(), utils.het.pop(), utils.impute,
utils.is.fixed(), utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(),
utils.read.ped(), utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(),
utils.recalc.freqhomref(), utils.recalc.freqhomsnp(), utils.reset.flags(), utils.transpose(),
utils.vcfr2genlight.polyploid()
```

utils.reset.flags *#'* An internal utility function to reset to FALSE (or TRUE) the locus metric flags after some individuals or populations have been deleted.

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

utils.reset.flags

Usage

```
utils.reset.flags(x, set = FALSE, value = 2, verbose = NULL)
```

Arguments

х	Name of the genlight object [required].
set	Set the flags to TRUE or FALSE [default FALSE].
value	Set the default verbosity for all functions, where verbosity is not specified [de-fault 2].
verbose	Verbosity: 0, silent or fatal errors; 1, begin and end; 2, progress log; 3, progress and results summary; 5, full report [default 2 or as specified using gl.set.verbosity]

Details

The locus metadata supplied by DArT has OneRatioRef, OneRatioSnp, PICRef, PICSnp, and Avg-PIC included, but the allelic composition will change when some individuals are removed from the dataset and so the initial statistics will no longer apply. This applies also to some variable calculated by dartR (e.g. maf). This script resets the locus metrics flags to FALSE to indicate that these statistics in the genlight object are no longer current. The verbosity default is also set, and in the case of SilcoDArT, the flags PIC and OneRatio are also set. If the locus metrics flags do not exist then they are added to the genlight object but not populated. If the locus metrics flags do not exist, then they are added to the genlight object and set to FALSE (or TRUE).

Value

The modified genlight object

Author(s)

Custodian: Luis Mijangos (Post to https://groups.google.com/d/forum/dartr)

See Also

utils.recalc.metrics for recalculating all metrics, utils.recalc.callrate for recalculating CallRate, utils.recalc.freqhomref for recalculating frequency of homozygous reference, utils.recalc.freqhomsnp for recalculating frequency of homozygous alternate, utils.recalc.freqhet for recalculating frequency of heterozygotes, gl.recalc.maf for recalculating minor allele frequency, gl.recalc.rdepth for recalculating average read depth

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.dist.binary(), utils.flag.start(), utils.hamming(), utils.het.pop(), utils.impute,
utils.is.fixed(), utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(),
utils.read.ped(), utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(),
utils.recalc.freqhomref(), utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.transpose(),
utils.vcfr2genlight.polyploid()
```

Examples

result <- utils.reset.flags(testset.gl)</pre>

utils.transpose

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

utils.transpose(x, parallel = FALSE)

Arguments

х	name of the genlight object
parallel	if TRUE, use parallel processing capability

Details

This is a function to transpose a genlight object, that is, to set loci as entities and individuals as attributes.

Value

a transposed genlight object

See Also

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.dist.binary(), utils.flag.start(), utils.hamming(), utils.het.pop(), utils.impute,
utils.is.fixed(), utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(),
utils.read.ped(), utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(),
utils.recalc.freqhomref(), utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.reset.flags(),
utils.vcfr2genlight.polyploid()
```

utils.vcfr2genlight.polyploid

Utility function to convert polyploid vcfR object as genlight

Description

WARNING: UTILITY SCRIPTS ARE FOR INTERNAL USE ONLY AND SHOULD NOT BE USED BY END USERS AS THEIR USE OUT OF CONTEXT COULD LEAD TO UNPRE-DICTABLE OUTCOMES.

Usage

```
utils.vcfr2genlight.polyploid(x, n.cores = 1, mode2 = mode)
```

Arguments

х	Name of the vcfR object [defined in function gl.read.vcf].
n.cores	Number of cores [default 1]
mode2	genotype: all heterozygous sites will be coded as 1 regardless ploidy level, dosage: sites will be codes as copy number of alternate allele [defined in function gl.read.vcf].

Details

This function uses parameters from gl.read.vcf for conversion Note also that this function checks to see if there are input of mode, missing input of mode will issued the user with a error. "Dosage" mode of this function assign ploidy levels as maximum copy number of alternate alleles. Please carefully check the data if "dosage" mode is used. (codes were modified from 'vcfR2genlight' in vcfR packge to convert polyploid data)

Value

genlight object

Author(s)

Custodian: Ching Ching Lau – Post to https://groups.google.com/d/forum/dartr

References

- Knaus, B. J., & Grunwald, N. J. (2017). vcfr: a package to manipulate and visualize variant call format data in R. Molecular ecology resources, 17(1), 44-53.
- Knaus, B. J., Grunwald, N. J., Anderson, E. C., Winter, D. J., Kamvar, Z. N., & Tabima, J. F. (2023). Package 'vcfR'. vcfR

See Also

```
Other utilities: gl.alf(), utils.check.datatype(), utils.collapse.matrix(), utils.dart2genlight(),
utils.dist.binary(), utils.flag.start(), utils.hamming(), utils.het.pop(), utils.impute,
utils.is.fixed(), utils.jackknife(), utils.n.var.invariant(), utils.plot.save(), utils.read.fasta(),
utils.read.ped(), utils.recalc.avgpic(), utils.recalc.callrate(), utils.recalc.freqhets(),
utils.recalc.freqhomref(), utils.recalc.freqhomsnp(), utils.recalc.maf(), utils.reset.flags(),
utils.transpose()
```

Examples

```
## Not run:
datatype <- utils.vcfr2genlight.polyploid(x=vcfr, mode2="genotype")</pre>
```

End(Not run)

zzz

Description

Setting up the package Setting theme, colors and verbosity

Usage

zzz

Format

An object of class NULL of length 0.

[,dartR,ANY,ANY,ANY-method

indexing dartR objects correctly...

Description

indexing dartR objects correctly...

Usage

S4 method for signature 'dartR,ANY,ANY,ANY'
x[i, j, ..., pop = NULL, treatOther = TRUE, quiet = TRUE, drop = FALSE]

Arguments

х	dartR object
i	index for individuals
j	index for loci
	other parameters
рор	list of populations to be kept
treat0ther	elements in other (and ind.metrics & loci.metrics) as indexed as well. default: TRUE
quiet	warnings are suppressed. default: TRUE
drop	reduced to a vector if a single individual/loci is selected. default: FALSE [should never set to TRUE]

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