Package 'EFGLmh'

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Type Package
Title Functions For Working With Microhaps for EFGL
Version 0.1.0
Description Written to work with microhaps and SNPs (which are just short microhaps). More generally, will function with codominant, diploid genotypes. Uses ``Progeny-style" (and FishGen-style) inputs. Performs basic manipulations, data summaries, and exporting data in formats for other packages/programs.
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R topics documented:
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Description

calculate allelic richness of loci

Usage

aRich(x)

Arguments

x an EFGLdata object

Value

a tibble giving the allelic richness of each locus

calcAF 3

calcAF

calculate allele frequencies within populations Combinations of pop, locus, allele that are not observed are (silently) excluded from the output. To add these back in, one can use the 'tidyverse' function 'complete' with the output of this function.

Description

calculate allele frequencies within populations Combinations of pop, locus, allele that are not observed are (silently) excluded from the output. To add these back in, one can use the 'tidyverse' function 'complete' with the output of this function.

Usage

```
calcAF(x)
```

Arguments

Х

an EFGLdata object

Value

a tibble

calcHet

calculate expected and observed heterozygosity within populations

Description

calculate expected and observed heterozygosity within populations

Usage

```
calcHet(x)
```

Arguments

Χ

an EFGLdata object

Value

a tibble

4 combineEFGLdata

cleanGrandma	convenience function to remove loci with all fails or no variation from
	gRandma input

Description

convenience function to remove loci with all fails or no variation from gRandma input

Usage

```
cleanGrandma(baseline, mixture = NULL)
```

Arguments

```
baseline a gRandma baseline input mixture a gRandma mixture input
```

Value

a list with two components, one is the baseline, one is the mixture

combineEFGLdata

combine multiple EFGL objects into one

Description

combine multiple EFGL objects into one

Usage

```
combineEFGLdata(
    ...,
    genoComb = c("intersect", "union"),
    metaComb = c("intersect", "union")
)
```

Arguments

... multiple EFGLdata objects separated by commas

genoComb if the objects have different loci, whether to create a new object with the inter-

section or union of loci. If union, genotypes for missing loci are all NA.

metaComb if the objects have different metadata fields, whether to create a new object with

the intersection or union of the fields. If union, missing fields are all NA.

construct_EFGLdata 5

construct_EFGLdata

some basic checks on EFGLdata objects

Description

some basic checks on EFGLdata objects

Usage

```
construct_EFGLdata(x)
```

Arguments

Х

an EFGLdata object

createF1Hybrids

Create F1 hybrids from two populations

Description

Creates hybrids by sampling one allele from one pop and the other allele from the other pop based on observed allele frequencies. Missing genotypes are sampled based on the combined rate of missing genotypes or data can be simulated with no missing genotypes. Alleles at different loci are sampled independently. Note that if a sex marker exists in the data set, it is treated like any other marker, resulting in XX, XY, and YY genotypes in the hybrids. You may want to remove it prior to simulation using the 'removeLoci' function. Loci with all missing genotypes in one population are simulated to have all missing genotypes in the hybrids REGARDLESS of the input value for 'missingGenos'.

Usage

```
createF1Hybrids(x, pop1, pop2, newName, n = 50, missingGenos = TRUE)
```

Arguments

x an EFGLdata object

pop1 one parent population (must be in 'x')
pop2 the other parent population (must be in 'x')

newName a string giving the name of the population to put hybrids into. This MUST be a

new pop.

n the number of hybrids to simulate

missingGenos TRUE to simulate missing genotypes, FALSE to not

Value

an EFGL data object with all the populations in 'x', plus the new simulated hybrid population 6 exampleData

dumpTable

wrapper for write table with commonly used options - carried over from IDFGEN

Description

wrapper for write table with commonly used options - carried over from IDFGEN

Usage

```
dumpTable(x, filename, row.names = FALSE, sep = "\t")
```

Arguments

x object to write out

filename to write out as row.names passed to write.table sep passed to write.table

Value

nothing, just writes a file

exampleData

An example input dataset used in the vignette

Description

Adds a population name column (required by EFGLmh).

Usage

exampleData

Format

a tibble

exportCKMRsimAF 7

export	CKMD	cimA	_
export	U.KIYIK	STIIIA	г

return a CKMRsim allele frequency input tibble.

Description

No chromosome and position information is used. All populations specified are combined into one allele frequency output. Any loci with only missing genotypes are omitted from the output. The output should then be run through the CKMRsim function reindex_markers().

Usage

```
exportCKMRsimAF(x, pops = NULL, loci = NULL)
```

Arguments

x an EFGLdata object

pops a vector of pops to include. If not specified, all pops are used.

loci a vector of loci to include. If not specified, all loci are used.

Value

a tibble

exportCKMRsimLG

Return a long genotype tibble for input to CKMRsim for relationship inference.

Description

Return a long genotype tibble for input to CKMRsim for relationship inference.

Usage

```
exportCKMRsimLG(x, pops = NULL, loci = NULL)
```

Arguments

x an EFGLdata object

pops a vector of pops to include. If not specified, all pops are used.

loci a vector of loci to include. If not specified, all loci are used.

Value

a tibble

8 exportColony

exportColony

write a Colony input file.

Description

write a Colony input file.

Usage

```
exportColony(
    x,
    filename = "colony2.dat",
    pops = NULL,
    loci = NULL,
    candMales = NULL,
    candFemales = NULL,
    candFemaleProb = 0.01,
    otherGenotypingErrorRate = 0.01
)
```

Arguments

an EFGLdata object filename the name of the file to write. If NULL, the colony input is returned as a character vector with each line a separate entry in the vector. pops a vector of pops to include. If not specified, all pops are used. loci a vector of loci to include. If not specified, all loci are used. candMales candidate male names candFemales candidate female names candMaleProb probability father is included in candidate males. if NULL, either 0 or 0.5 depending on whether candMales is NULL. candFemaleProb probability father is included in candidate Females. if NULL, either 0 or 0.5 depending on whether candFemales is NULL. projName project name in colony input colOutFileName output file name to direct colony to use alleleDropoutRate allelic dropout rate to use for all loci $other {\tt Genotyping Error Rate}$ other genotyping error rate to use for all loci

Value

nothing, just writes a file

exportGenAlEx 9

 ${\tt exportGenAlEx}$

write a GenAlEx input file

Description

write a GenAlEx input file

Usage

```
exportGenAlEx(
   x,
   filename,
   pops = NULL,
   loci = NULL,
   title = "",
   useNames = TRUE
)
```

Arguments

x an EFGLdata object

filename the name of the file to write

pops a vector of pops to include. If not specified, all pops are used.

loci a vector of loci to include. If not specified, all loci are used.

title a string to use as the "title" row

useNames TRUE to use samples names, FALSE to replace with unique numerical identi-

fiers

Value

nothing, just writes a file

exportGenePop

write a genepop input file

Description

write a genepop input file

Usage

```
exportGenePop(
    x,
    filename,
    header = "genePop file",
    pops = NULL,
    loci = NULL,
    useIndNames = FALSE
)
```

10 exportGrandma

Arguments

x an EFGLdata object

filename the name of the file to write

header a string to use as the header line of the genepop file

pops a vector of pops to include. If not specified, all pops are used.

loci a vector of loci to include. If not specified, all loci are used.

useIndNames TRUE to use individual names as sample identifiers. Otherwise, population

names are used

Value

nothing, just writes a file

exportGrandma export a gRandma baseline or mixture

Description

export a gRandma baseline or mixture

Usage

```
exportGrandma(x, pops = NULL, loci = NULL, baseline = TRUE)
```

Arguments

x an EFGLdata object

pops a vector of pops to include in the baseline. If not specified, all pops are used.

loci a vector of loci to use. If not specified, all loci are used.

baseline TRUE to make a baseline input, FALSE to make a mixture input.

Value

a tibble

exportHierFstat 11

exportHierFstat

export a hierfstat input dataframe

Description

export a hierfstat input dataframe

Usage

```
exportHierFstat(x, pops = NULL, loci = NULL)
```

Arguments

x an EFGLdata object

pops a vector of pops to include. If not specified, all pops are used.

loci a vector of loci to use. If not specified, all loci are used.

Value

a dataframe coded to be used as input for hierfstat

exportPlink

export a Plink PED file and optionally a (not very useful except as a template) MAP file

Description

Only biallelic markers are used, exports as two column per call. This format can be used as the input for "Admixture".

Usage

```
exportPlink(
    x,
    filename,
    pops = NULL,
    loci = NULL,
    FID = "Ind",
    IID = "Ind",
    pa = NULL,
    ma = NULL,
    sex = NULL,
    pheno = NULL,
    map = NULL)
```

12 exportProgenyStyle

Arguments

X	an EFGLdata object
filename	the name of the file to write
pops	a vector of pops to include. If not specified, all pops are used.
loci	a vector of loci to use. If not specified, all loci are used.
FID	Metadata column name to use as the family ID
IID	Metadata column name to use as the within-family ID
pa	Metadata column name to use as the within-family ID of the father. If NULL, all are listed as unknown (0) .
ma	Metadata column name to use as the within-family ID of the mother. If NULL, all are listed as unknown (0) .
sex	Metadata column to use as the sex. This should be coded as "M" for male, "F" for female, and any other values are treated as Unknown. If NULL, all are listed as unknown (0).
pheno	Metadata column to use as the phenotype. This should be coded as "1" for control, "2" for case, and "-9" or "0" for Unknown. If NULL, all are listed as unknown (0) .
map	filename to write a MAP file. If NULL, no MAP file is written. This just writes a dummy MAP file with the loci names in order and all given the same chromosome code and positions of "0". If you need a valid MAP file, you will need to edit this.

Value

nothing, just writes a file

 $export Progeny Style \qquad \textit{export a "Progeny-style" export file for later reading into EFGLmh}$

Description

Columns in order are Pop, Ind, metadata, genotypes (2-column per call) Missing genotypes are "0" for SNPs (biallelic or nonvariable with alleles represented by 1 character) and "000" for others. If a locus is all missing, it is treated as a SNP. Pop column is called "Pedigree" and Ind column is called "Individual.Name".

Usage

```
exportProgenyStyle(x, filename, pops = NULL, loci = NULL, metadata = NULL)
```

Arguments

x an EFGLdata object filename the name of the file to write

pops a vector of pops to include. If not specified, all pops are used.

loci a vector of loci to use. If not specified, all loci are used.

metadata a vector of metadata fields to include. If not specified, all fields are used.

exportRubias_baseline 13

Value

nothing, just writes a file

exportRubias_baseline export a rubias baseline

Description

export a rubias baseline

Usage

```
exportRubias_baseline(
   x,
   pops = NULL,
   repunit = NULL,
   collection = NULL,
   loci = NULL
)
```

Arguments

x an EFGLdata object

pops a vector of pops to include in the baseline. If not specified, all pops are used.

repunit the column name of the metadata variable designating repunit. This can be Pop

to use the population name. If not specified, NA is used for all samples.

collection the column name of the metadata variable designating collection. This can be

Pop to use the population name. If not specified, NA is used for all samples.

loci a vector of loci to use. If not specified, all loci are used.

Value

a tibble

```
exportRubias_mixture export a rubias mixture
```

Description

export a rubias mixture

Usage

```
exportRubias_mixture(x, pops = NULL, collection = NULL, loci = NULL)
```

14 exportSNPPIT

Arguments

x an EFGLdata object

pops a vector of pops to include in the baseline. If not specified, all pops are used.

collection the column name of the metadata variable designating collection. This can be

Pop to use the population name. If not specified, NA is used for all samples. For mixtures, this variable indicates what samples come from the same "stratum" -

to be analyzed together.

loci a vector of loci to use. If not specified, all loci are used.

Value

a tibble

exportSNPPIT write a SNPPIT input file. Will warn about skipping loci with > 2

alleles.

Description

write a SNPPIT input file. Will warn about skipping loci with > 2 alleles.

Usage

```
exportSNPPIT(
    x,
    filename,
    baseline,
    mixture,
    loci = NULL,
    errorRate = 0.005,
    POPCOLUMN_SEX = NULL,
    POPCOLUMN_REPRO_YEARS = NULL,
    POPCOLUMN_SPAWN_GROUP = NULL,
    OFFSPRINGCOLUMN_BORN_YEAR = NULL,
    OFFSRPINGCOLUMN_SAMPLE_YEAR = NULL,
    OFFSPRINGCOLUMN_AGE_AT_SAMPLING = NULL
)
```

Arguments

x an EFGLdata object

filename the name of the file to write

baseline a vector of pops to use as the baseline (potential parents).

mixture a vector of pops to use as the mixture (potential offspring).

loci a vector of loci to include. If not specified, all loci are used.

errorRate per allele error rate for all loci

POPCOLUMN_SEX metadata column with sex info (coded as M, F, and ?)

exportStructure 15

POPCOLUMN_REPRO_YEARS

metadata column with repro years

POPCOLUMN_SPAWN_GROUP

metadata column with spawn group

OFFSPRINGCOLUMN_BORN_YEAR

metadata column with birth year

OFFSRPINGCOLUMN_SAMPLE_YEAR

metadata column with sample year

OFFSPRINGCOLUMN_AGE_AT_SAMPLING

metadata column with age at sampling

Value

nothing, just writes a file

exportStructure

write a Structure input file.

Description

Columns are Ind, Pop (converted to integers), then genotypes. Missing alleles are coded as -9

Usage

```
exportStructure(x, filename, pops = NULL, loci = NULL)
```

Arguments

x an EFGLdata object

filename the name of the file to write

pops a vector of pops to include. If not specified, all pops are used.

loci a vector of loci to include. If not specified, all loci are used.

Value

nothing, just writes a file

16 getInds

genoSuccess	calculate genotyping success of individuals (uses only allele 1 for each
	genotype - assumes if allele 1 is (is not) NA, so is (is not) allele 2)

Description

calculate genotyping success of individuals (uses only allele 1 for each genotype - assumes if allele 1 is (is not) NA, so is (is not) allele 2)

Usage

```
genoSuccess(x, loci = NULL)
```

Arguments

x an EFGLdata object

loci a vector of loci to include. If not specified, all loci are used.

Value

a tibble giving the genotyping success of each individual as a proportion and number of missing genotypes

getInds

get a vector of individuals present

Description

get a vector of individuals present

Usage

```
getInds(x, pops = NULL)
```

Arguments

x an EFGLdata object

pops a vector of pops that you want individual names for. If not specified, names for

all pops are returned

Value

a vector of the Individual names present

getLoci 17

getLoci

get a vector of loci names present

Description

get a vector of loci names present

Usage

getLoci(x)

Arguments

Х

an EFGLdata object

getMeta

get a vector of metadata column names present

Description

get a vector of metadata column names present

Usage

getMeta(x)

Arguments

Х

an EFGLdata object

getPops

get a vector of populations (pedigrees) present

Description

get a vector of populations (pedigrees) present

Usage

getPops(x)

Arguments

Х

an EFGLdata object

Value

a vector of the unique population names present

18 moveInds

lociSuccess	calculate genotyping success of loci (uses only allele 1 for each geno-
	type - assumes if allele 1 is (is not) NA, so is (is not) allele 2)

Description

calculate genotyping success of loci (uses only allele 1 for each genotype - assumes if allele 1 is (is not) NA, so is (is not) allele 2)

Usage

lociSuccess(x)

Arguments

x an EFGLdata object

Value

a tibble giving the genotyping success of each locus as a proportion

moveInds combine individuals into one population AND REMOVE the previous entry for those individuals

Description

combine individuals into one population AND REMOVE the previous entry for those individuals

Usage

```
moveInds(x, inds, newName)
```

Arguments

x an EFGLdata object

inds a vector of individuals to put in the new pop

newName a string giving the name of population to add the individuals too. This can be a

new pop or an existing pop (a warning is issued if existing).

Value

an EFGLdata object

movePops 19

movePops	combine populations into one AND REMOVE the old populations

Description

combine populations into one AND REMOVE the old populations

Usage

```
movePops(x, pops, newName)
```

Arguments

x an EFGLdata object

pops a vector of populations to combine

newName a string giving the name of the population to combine pops into. This can be a

new pop or an existing pop (a warning is issued if existing).

Value

an EFGLdata object

mtype2wide	A function for turning the genotype output of mtype2 (microTyper) into a wide format for input into EFGLmh

Description

A function for turning the genotype output of mtype2 (microTyper) into a wide format for input into EFGLmh

Usage

```
mtype2wide(x, popName = "PopulationName")
```

Arguments

x the output of mtype2, either a tibble or a path to a file popName The value to give all samples for population name.

20 print.EFGLdata

numInds

get the number of individuals present in each pop

Description

get the number of individuals present in each pop

Usage

```
numInds(x, pops = NULL)
```

Arguments

x an EFGLdata object

pops a vector of pops that you want individual names for. If not specified, numbers

for all pops are returned

Value

a named vector with the number of individuals in each pop

print.EFGLdata

print method for EFGLdata

Description

print method for EFGLdata

Usage

```
## S3 method for class 'EFGLdata'
print(x, ...)
```

Arguments

x an EFGLdata object

... ignored

readInData 21

readInData	read	in	data	from	a	Progeny-style	output	file	or	ma-
	trix/d	atafr	ame/til	bble adı	ı cre	eate an EFGLdai	ta object			

Description

read in data from a Progeny-style output file or matrix/dataframe/tibble adn create an EFGLdata object

Usage

```
readInData(
  input,
  genotypeStart = NULL,
  pedigreeColumn = 1,
 nameColumn = 2,
  convertNames = TRUE,
  convertMetaDataNames = TRUE,
 missingAlleles = c("0", "00", "000"),
 guess_max = 10000
)
```

Arguments

input

Either a character string to the tab-separated input file with a header row or a matrix/dataframe/tibble. Structure of the input: one row per individual. One column giving pedigree (population) names, one column giving individual names, optional additional metadata columns, genotype columns. Pedigree and individual name columns can be anywhere, if specified. Genotype columns MUST be consecutive and be the right most columns. Genotypes are given as two columns per call (diploidy assumed).

genotypeStart

The column number that genotypes start at. If not specified, the first column with a column name ending in ".A1", ".a1", "-A1", or "-a1" is chosen.

pedigreeColumn The column number that contains pedigree (population) names.

nameColumn

The column number that contains individual names. These MUST be unique.

convertNames

TRUE to convert genotype and pedigree names in the same way that IDF-GEN does (remove special characters from both and remove "." from genotype names).

convertMetaDataNames

TRUE to remove special characters and spaces from metadata column names. This makes accessing them easier.

missingAlleles a vector of values (not NA) to treat as missing alleles. They will be converted to

guess_max

If input is a character string, this is the maximum number of lines to use when guessing input data types. Making this smaller results in quicker loading, making it larger can fix some parsing errors

22 removeLoci

Value

An EFGLdata object, which is just a list with two elements. The first element is a tibble with genotype data, the second is a tibble with metadata

removeInds

remove individuals from an EFGLdata object

Description

remove individuals from an EFGLdata object

Usage

```
removeInds(x, inds)
```

Arguments

x an EFGLdata object

inds a vector of individuals to remove

Value

an EFGLdata object

 $\verb"removeLoci"$

remove loci from an EFGLdata object

Description

remove loci from an EFGLdata object

Usage

```
removeLoci(x, lociRemove)
```

Arguments

x an EFGLdata object

lociRemove a vector of loci names to remove

removePops 23

removePops

remove pops from an EFGLdata object

Description

remove pops from an EFGLdata object

Usage

```
removePops(x, pops)
```

Arguments

x an EFGLdata object

pops a vector of pops to remove

Value

an EFGLdata object

whichLower

Identify which individual out of duplicate pairs has the lower genotyp-

ing success

Description

Identify which individual out of duplicate pairs has the lower genotyping success

Usage

```
whichLower(dupTable, geno_success)
```

Arguments

 ${\tt dupTable} \qquad \qquad {\tt the \ output \ of \ close_matching_samples \ (from \ rubias)}$

geno_success the output of genoSuccess

Value

a vector of unique individual names representing the individuals with lower genotyping success from each pair

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