Package ‘SeqVarTools’

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Author Stephanie M. Gogarten, Xiuwen Zheng
Maintainer Stephanie M. Gogarten <sdmorris@u.washington.edu>, Xiuwen Zheng <zhengx@u.washington.edu>
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SeqVarTools-package  

Tools for Variant Analysis

Description

This package provides tools for data exploration and analysis of variants, extending the functionality of the package SeqArray.

Details

SeqArray provides an alternative to the Variant Call Format (VCF) for storage of variants called from sequencing data, enabling efficient storage, fast access to subsets of the data, and rapid computation.

SeqVarTools provides an interface to the SeqArray storage format with tools for many common tasks in variant analysis and integration with basic S4 classes in Bioconductor.

Author(s)

Stephanie M. Gogarten, Xiuwen Zheng
Maintainer: Stephanie M. Gogarten <sdmorris@u.washington.edu>

allele-methods  

Extract allele information from a GDS object

Description

Extract reference and alternate alleles and allele counts from a GDS object.

Usage

```r
## S4 method for signature 'SeqVarGDSClass'
refChar(gdsobj)
## S4 method for signature 'SeqVarGDSClass'
altChar(gdsobj, n=0)
## S4 method for signature 'SeqVarGDSClass'
nAlleles(gdsobj)
```
Arguments

gdsobj A SeqVarGDSClass object with VCF data.

n An integer indicating which alternate allele to return. $n=0$ returns a comma-separated string of all alternate alleles.

Details

These methods parse the "allele" field of a GDS object.

Value

refChar returns a character vector of reference alleles.

altChar returns a character vector of alternate alleles. If $n=0$, multiple alternate alleles are represented as a comma-separated string. If $n>0$, only the $n$th alternate allele is returned.

nAlleles returns an integer vector of the number of alleles (reference and alternate) for each variant.

Author(s)

Stephanie Gogarten

See Also

SeqVarGDSClass, applyMethod

Examples

gds <- seqOpen(seqExampleFileName("gds"))
table(refChar(gds))
table(altChar(gds))
table(altChar(gds, n=1))
table(altChar(gds, n=2), useNA="ifany")
table(nAlleles(gds))
seqClose(gds)

alleleFrequency Allele frequency

Description

Calculate allele frequency for each variant

Usage

## S4 method for signature 'SeqVarGDSClass'
alleleFrequency(gdsobj, n=0, use.names=FALSE)
Arguments

- **gdobj**: A `SeqVarGDSClass` object with VCF data.
- **n**: An integer indicating which allele to calculate the frequency of. \( n=0 \) is the reference allele, \( n=1 \) is the first alternate allele, and so on.
- **use.names**: A logical indicating whether to assign variant IDs as names of the output vector.

Details

Frequency can be calculated over any allele, specified by the argument \( n \). Default is the reference allele frequency (\( n=0 \)).

Value

A numeric vector of allele frequencies.

Author(s)

Stephanie Gogarten

See Also

- `SeqVarGDSClass`, `applyMethod`, `heterozygosity`

Examples

```r
gds <- seqOpen(seqExampleFileName("gds"))
head(alleleFrequency(gds))
head(alleleFrequency(gds, n=1))
head(alleleFrequency(gds, n=2))
seqClose(gds)
```

Description

Apply a method to a subset of variants and/or samples in a GDS object

Usage

```r
## S4 method for signature 'SeqVarGDSClass, function, character'
applyMethod(gdobj, FUN, variant, sample=NULL, ...)
## S4 method for signature 'SeqVarGDSClass, function, numeric'
applyMethod(gdobj, FUN, variant, sample=NULL, ...)
## S4 method for signature 'SeqVarGDSClass, function, GRanges'
applyMethod(gdobj, FUN, variant, sample=NULL, ...)
## S4 method for signature 'SeqVarGDSClass, function, missing'
applyMethod(gdobj, FUN, variant, sample=NULL, ...)
```
applyMethod

Arguments

gdsobj   A `SeqVarGDSClass` object with VCF data.

FUN       A method or function to be applied to gdsobj.

variant A vector of variant.id values or a GRanges object defining the variants to be included in the call to FUN.

sample  A vector of sample.id values defining the samples to be included in the call to FUN.

... Additional arguments, passed to FUN.

Details

applyMethod applies a method or function FUN to the subset of variants defined by variant and samples defined by sample in a GDS object.

If a filter was previously set with `seqSetFilter`, it will be saved and reset after the call to applyMethod.

Value

The result of the call to FUN.

Author(s)

Stephanie Gogarten

See Also

`SeqVarGDSClass`

Examples

gds <- seqOpen(seqExampleFileName("gds"))
variant.id <- seqGetData(gds, "variant.id")
sample.id <- seqGetData(gds, "sample.id")
applyMethod(gds, getGenotype, variant.id[1:5], sample.id[1:10])

library(GenomicRanges)
chrom <- seqGetData(gds, "chromosome")
pos22 <- seqGetData(gds, "position")[chrom == 22]
ranges <- GRanges(seqnames="22", IRanges(min(pos22), max(pos22)))
applyMethod(gds, heterozygosity, ranges, margin="by.sample")
applyMethod(gds, heterozygosity, ranges, margin="by.variant")

seqClose(gds)
duplicateDiscordance  

Duplicate discordance

**Description**

Find discordance rate for duplicate sample pairs

**Usage**

```r
## S4 method for signature 'SeqVarGDSClass'
duplicateDiscordance(gdsobj, samples, check.phase=FALSE, verbose=TRUE)
```

**Arguments**

- `gdsobj`  
  A `SeqVarGDSClass` object with VCF data.
- `samples`  
  A data.frame with columns (sample.id, subject.id). "sample.id" values should correspond to "sample.id" in gdsobj. "subject.id" should match for duplicate samples.
- `check.phase`  
  A logical indicating whether phase should be considered when calculating discordance.
- `verbose`  
  A logical indicating whether to print a progress message for each sample.

**Details**

Duplicate discordance is calculated by sample pair and by variant. If there are more than two samples per subject in `samples`, only the first two samples are used and a warning message is printed.

If `check.phase=TRUE`, variants with mismatched phase are considered discordant. If `check.phase=FALSE`, phase is ignored.

**Value**

A list with the following elements:

- `by.variant`  
  A data.frame with the number of discordances for each variant, the number of sample pairs with non-missing data, and the discordance rate (num.discard / num.pair). Row names are variant ids.
- `by.subject`  
  A data.frame with the sample ids for each pair, the number of discordances, the number of non-missing variants, and the discordance rate (num.discard / num.var). Row names are subject.id (as given in `samples`).

**Author(s)**

Stephanie Gogarten
getGenotype

See Also
SeqVarGDSClass, applyMethod

Examples

```r
gds <- seqOpen(seqExampleFileName("gds"))
  # the example file has one sample per subject, but we
  # will match the first four samples into pairs as an example
  sample.id <- seqGetData(gds, "sample.id")
  samples <- data.frame(subject.id=rep(c("subj1", "subj2"), each=2),
                         sample.id=sample.id[1:4],
                         stringsAsFactors=FALSE)
  disc <- duplicateDiscordance(gds, samples)
  head(disc$by.variant)
  disc$by.subject
  seqClose(gds)
```

Description

Get matrix of genotype values from a GDS object as VCF-style character strings

Usage

```r
## S4 method for signature 'SeqVarGDSClass'
getGenotype(gdsobj, use.names=TRUE)
## S4 method for signature 'SeqVarGDSClass'
getGenotypeAlleles(gdsobj, use.names=TRUE, sort=FALSE)
## S4 method for signature 'SeqVarGDSClass'
refDosage(gdsobj, use.names=TRUE)
```

Arguments

- `gdsobj`: A `SeqVarGDSClass` object with VCF data.
- `use.names`: A logical indicating whether to assign sample and variant IDs as dimnames of the resulting matrix.
- `sort`: Logical for whether to sort alleles lexicographically ("G/T" instead of "T/G").

Details

In `getGenotype`, genotypes are coded as in the VCF file, where "0/0" is homozygous reference, "0/1" is heterozygous for the first alternate allele, "0/2" is heterozygous for the second alternate allele, etc.

Separators are "/" for unphased and "|" for phased. If `sort=TRUE`, all returned genotypes will be unphased. Missing genotypes are coded as NA.

Only diploid genotypes (the first two alleles at a given site) are returned.
getVariableLengthData

Value

getGenotype and getGenotypeAlleles return a character matrix with dimensions [sample,variant] containing diploid genotypes.
getGenotype returns alleles as "0", "1", "2", etc. indicating reference and alternate alleles.
getGenotypeAlleles returns alleles as "A", "C", "G", "T". sort=TRUE sorts lexicographically, which may be useful for comparing genotypes with data generated using a different reference sequence.
refDosage returns an integer matrix with the dosage of the reference allele: 2 for two copies of the reference allele ("0/0"), 1 for one copy of the reference allele, and 0 for two alternate alleles.

Author(s)
Stephanie Gogarten

See Also
SeqVarGDSClass, applyMethod, seqGetData

Examples

gds <- seqOpen(seqExampleFileName("gds"))
variant.id <- seqGetData(gds, "variant.id")
sample.id <- seqGetData(gds, "sample.id")
seqSetFilter(gds, variant.id=variant.id[1:5],
              sample.id=sample.id[1:10])
getGenotype(gds)
getGenotypeAlleles(gds)
refDosage(gds)
seqClose(gds)

---

getVariableLengthData  Get variable-length data

Description

Get data with multiple values per sample from a GDS object and return as an array

Usage

## S4 method for signature 'SeqVarGDSClass,character'
getVariableLengthData(gdsobj, var.name, use.names=TRUE)

Arguments

gdsobj  A SeqVarGDSClass object with VCF data.
var.name  Character string with name of the variable, most likely "annotation/format/VARIABLE_NAME".
use.names  A logical indicating whether to assign sample and variant IDs as dimnames of the resulting matrix.
getVariableLengthData

Details

Data which are indicated as having variable length (possibly different numbers of values for each variant) in the VCF header are stored as variable-length data in the GDS file. Each such data object has two components, "length" and "data." "length" indicates how many values there are for each variant, while "data" is a matrix with one row per sample and columns defined as all values for variant 1, followed by all values for variant 2, etc.

getVariableLengthData converts this format to a 3-dimensional array, where the length of the first dimension is the maximum number of values in "length," and the remaining dimensions are sample and variant. Missing values are given as NA. If the first dimension of this array would have length 1, the result is converted to a matrix.

Value

An array with dimensions [n, sample, variant] where n is the maximum number of values possible for a given sample/variant cell. If n=1, a matrix with dimensions [sample,variant].

Author(s)

Stephanie Gogarten

See Also

SeqVarGDSClass, applyMethod, seqGetData

Examples

```r
file <- system.file("extdata", "gl_chr1.gds", package="SeqVarTools")
gds <- seqOpen(file)
## genotype likelihood
gl <- seqGetData(gds, "annotation/format/GL")
names(gl)
gl[length]
## 3 values per variant - likelihood of RR,RA,AA genotypes
dim(gl$data)
## 85 samples (rows) and 9 variants with 3 values each - 27 columns

gl.array <- getVariableLengthData(gds, "annotation/format/GL")
dim(gl.array)
## 3 genotypes x 85 samples x 9 variants
head(gl.array[1,,])
head(gl.array[2,,])
head(gl.array[3,,])

## genotype dosage
ds <- seqGetData(gds, "annotation/format/DS")
names(ds)
ds[length]
## 1 value per variant
dim(ds$data)
## 85 samples (rows) and 9 variants (columns)
```
ds.array <- getVariableLengthData(gds, "annotation/format/DS")
dim(ds.array)
## 85 samples x 9 variants
head(ds.array)

seqClose(gds)

---

heterozygosity  Heterozygosity and Homozygosity

Description

Calculate heterozygosity and homozygosity by variant or by sample.

Usage

```
## S4 method for signature 'SeqVarGDClass'
heterozygosity(gdsobj, margin=c("by.variant", "by.sample"), use.names=FALSE)
## S4 method for signature 'SeqVarGDClass'
homozygosity(gdsobj, allele=c("any", "ref", "alt"), margin=c("by.variant", "by.sample"), use.names=FALSE)
```

Arguments

- **gdsobj**: A `SeqVarGDClass` object with VCF data.
- **margin**: Possible values are "by.variant" or "by.sample," indicating whether the calculation should be done over all samples for each variant, or over all variants for each sample.
- **use.names**: A logical indicating whether to assign variant or samples IDs as names of the output vector.
- **allele**: Possible values are "any", "ref," or "alt," indicating which alleles to consider when calculating homozygosity.

Details

- `heterozygosity` calculates the fraction of heterozygous genotypes in a GDS object, either by variant or by sample.
- `homozygosity` calculates the rate of homozygous genotypes in a GDS object, either by sample or by variant. If `allele="any"`, all homozygous genotypes are considered (reference or any alternate allele). If `allele="ref"`, only reference homozygotes are considered. If `allele="alt"`, any alternate allele homozygote is considered. For example, "ref" will count "0/0" genotypes only, "alt" will count "1/1", "2/2", etc. (but not "0/0"), and "any" will count all of the above.

Value

A numeric vector of heterozygosity or homozygosity rates. If `margin="by.variant"`, the vector will have length equal to the number of variants in the GDS object. If `margin="by.sample"`, the vector will have length equal to the number of samples.
**hwe**

*Exact test for Hardy-Weinberg equilibrium*

**Description**

Performs an exact test for Hardy-Weinberg equilibrium on Single-Nucleotide Variants

**Usage**

```r
## S4 method for signature 'SeqVarGDSClass'
hwe(gdsobj, use.names=FALSE)
```

**Arguments**

- **gdsobj**
  A `SeqVarGDSClass` object with VCF data.
- **use.names**
  A logical indicating whether to assign variant IDs as names of the output vector.

**Details**

HWE calculations are performed with the `HWExact` function in the `GWASExactHW` package. P values are set to `NA` for all non-single-nucleotide variants and monomorphic variants.

**Value**

A vector of p values for the exact test.

---

**Author(s)**

Stephanie Gogarten

**See Also**

`SeqVarGDSClass`, `applyMethod`, `alleleFrequency`


**Author(s)**

Stephanie Gogarten

**See Also**

SeqVarGDSClass, applyMethod

**Examples**

```r
library(SeqVarGDSClass)

gds <- seqOpen(seqExampleFileName("gds"))
## autosomal variants only
auto <- seqGetData(gds, "chromosome") %in% 1:22
var.auto <- seqGetData(gds, "variant.id"))[auto]
pv <- applyMethod(gds, hwe, variant=var.auto)
head(pv)
sum(is.na(pv))
range(pv, na.rm=TRUE)
seqClose(gds)
```

### inbreedCoeff

**Inbreeding coefficient**

**Description**

Calculates the inbreeding coefficient by variant or by sample

**Usage**

```r
## S4 method for signature 'SeqVarGDSClass'
inbreedCoeff(gdsobj, margin=c("by.variant", "by.sample"), use.names=FALSE)
```

**Arguments**

- `gdsobj`: A SeqVarGDSClass object with VCF data.
- `margin`: Possible values are "by.variant" or "by.sample," indicating whether the calculation should be done over all samples for each variant, or over all variants for each sample.
- `use.names`: A logical indicating whether to assign variant or sample IDs as names of the output vector.

**Details**

For inbreeding coefficients by variant, calculates $1 - \text{observed heterozygosity} / \text{expected heterozygosity}$.

For individual inbreeding coefficients (`margin=\"by.sample\"`), calculates Visscher's estimator described in Yang et al. (2010).
isSNV

Value

Values for the inbreeding coefficient.

Author(s)

Xiuwen Zheng, Stephanie Gogarten

References


See Also

SeqVarGDSClass, applyMethod

Examples

gds <- seqOpen(seqExampleFileName("gds"))
f <- inbreedCoeff(gds, margin="by.variant")
range(f, na.rm=TRUE)

ic <- inbreedCoeff(gds, margin="by.sample")
range(ic)
seqClose(gds)

isSNV

Flag single nucleotide variants

Description

Flag single nucleotide variants

Usage

## S4 method for signature 'SeqVarGDSClass'
isSNV(x, biallelic=TRUE)

Arguments

x A SeqVarGDSClass object with VCF data.
biallelic A logical indicating whether only biallelic SNVs are considered.
Details

If biallelic=TRUE, a variant is considered a single nucleotide variant (SNV) if there is one reference allele and one alternate allele, each one base in length. If biallelic=FALSE, there may be multiple alternate alleles, each one base in length.

Setting biallelic=TRUE is considerably faster for large data sets.

Value

A logical vector indicating which variants are SNVs.

Author(s)

Stephanie Gogarten

See Also

SeqVarGDSClass, allele-methods, applyMethod

Examples

gds <- seqOpen(seqExampleFileName("gds"))
table(isSNV(gds))
seqClose(gds)

isVariant

Locate variant samples across sites

Description

Locate which samples are variant for each site in a GDS object

Usage

## S4 method for signature 'SeqVarGDSClass'
isVariant(gdsobj, use.names=FALSE)

Arguments

<table>
<thead>
<tr>
<th>gdsobj</th>
<th>A SeqVarGDSClass object with VCF data.</th>
</tr>
</thead>
<tbody>
<tr>
<td>use.names</td>
<td>A logical indicating whether to assign sample and variant IDs as dimnames of the resulting matrix.</td>
</tr>
</tbody>
</table>

Details

Each sample/site cell of the resulting matrix is TRUE if the genotype at that location for that sample contains an alternate allele. A genotype of "0/0" is not variant, while genotypes "0/1", "1/0", "0/2", etc. are variant.
meanBySample

Value
A logical matrix with dimensions [sample, site] which is TRUE for cells where the genotype contains an alternate allele.

Author(s)
Stephanie Gogarten

See Also
SeqVarGDSClass, applyMethod, getGenotype

Examples
```r
gds <- seqOpen(seqExampleFileName("gds"))
variant.id <- seqGetData(gds, "variant.id")
sample.id <- seqGetData(gds, "sample.id")
applyMethod(gds, isVariant, variant.id[1:5], sample.id[1:10])
applyMethod(gds, isVariant, variant.id[1:5], sample.id[1:10], use.names=TRUE)
seqClose(gds)
```

---

meanBySample  
Mean value by sample

Description
Calculate the mean value of a variable by sample over all variants

Usage
```r
## S4 method for signature 'SeqVarGDSClass'
meanBySample(gdsobj, var.name, use.names=FALSE)
```

Arguments
- **gdsobj**: A SeqVarGDSClass object with VCF data.
- **var.name**: Character string with name of the variable, most likely "annotation/format/VARIABLE_NAME".
- **use.names**: A logical indicating whether to assign sample IDs as names of the output vector.

Details
Mean values by variant can be calculated using `seqApply(gdsobj, var.name, mean, na.rm=TRUE)`. Currently `seqApply` can only be used with the option `margin="by.variant"`. This method provides a way to calculate mean values by sample.

Value
A numeric vector of mean values.
**Author(s)**
Stephanie Gogarten

**See Also**
`SeqVarGDSClass`, `applyMethod`, `seqApply`

**Examples**
```
gds <- seqOpen(seqExampleFileName("gds"))
head(meanBySample(gds, "annotation/format/DP", use.names=TRUE))
seqClose(gds)
```

---

**mendelErr**  
*Mendelian errors*

**Description**
Detect Mendelian errors

**Usage**
```
## S4 method for signature 'SeqVarGDSClass'
mendelErr(gdsobj, pedigree, use.names=FALSE, 
          autosomes=1:22, xchrom="X", ychrom="Y", verbose=TRUE)
```

**Arguments**
- `gdsobj`  
  A `SeqVarGDSClass` object with VCF data.
- `pedigree`  
  A data.frame with columns (family, individ, father, mother, sex, sample.id).  
  "sex" column should have values "M"/"F".  "sample.id" values should correspond to "sample.id" in gdsobj.
- `use.names`  
  A logical indicating whether to assign variant IDs as names of the output vector.
- `autosomes`  
  A vector with chromosome values in gdsobj corresponding to autosomes.
- `xchrom`  
  The chromosome value in gdsobj corresponding to the X chromosome.
- `ychrom`  
  The chromosome value in gdsobj corresponding to the Y chromosome.
- `verbose`  
  A logical indicating whether to print the number of samples selected for each trio.

**Details**
Mendelian errors are detected for each trio in pedigree. Duos (mother or father missing) are included. The pedigree must have only one sample per individual.
Value

A list with the following elements:

by.variant An integer vector with the number of mendelian errors detected for each variant. If use.names=TRUE, the vector will be named with variant IDs.

by.trio An integer vector with the number of mendelian errors detected for each trio. The vector will be named with the sample ID of the child in each trio.

Author(s)

Stephanie Gogarten

See Also

SeqVarGDSClass, applyMethod

Examples

gds <- seqOpen(seqExampleFileName("gds"))
data(pedigree)
err <- mendelErr(gds, pedigree)
table(err$by.variant)
err$by.trio
seqClose(gds)

missingGenotypeRate  Missing genotype rate

Description

Calculate missing genotype rate by variant or by sample

Usage

## S4 method for signature 'SeqVarGDSClass'
missingGenotypeRate(gdsobj, margin=c("by.variant", "by.sample"), use.names=FALSE)

Arguments

gdsobj A SeqVarGDSClass object with VCF data.
margin Possible values are "by.variant" or "by.sample," indicating whether the calculation should be done over all samples for each variant, or over all variants for each sample.
use.names A logical indicating whether to assign variant IDs as names of the output vector.

Details

Calculates the fraction of missing genotypes in a GDS object, either by variant or by sample.
Value

A numeric vector of missing genotype rates. If margin="by.variant", the vector will have length equal to the number of variants in the GDS object. If margin="by.sample", the vector will have length equal to the number of samples.

Author(s)

Stephanie Gogarten

See Also

SeqVarGDSClass, applyMethod, getGenotype

Examples

gds <- seqOpen(seqExampleFileName("gds"))
head(missingGenotypeRate(gds, margin="by.variant"))
head(missingGenotypeRate(gds, margin="by.sample"))
seqClose(gds)

pca  Principal Component Analysis

Description

Calculates the eigenvalues and eigenvectors of a SeqVarGDSClass object with Principal Component Analysis

Usage

## S4 method for signature 'SeqVarGDSClass'
pca(gdsobj, eigen.cnt=32)

Arguments

gdsobj  A SeqVarGDSClass object with VCF data.
eigen.cnt  An integer indicating how many eigenvalues and eigenvectors to return.

Details

Calculates the genetic covariance matrix and finds the eigen decomposition.

Value

A list with two elements:
eigenva  A vector of length eigen.cnt with eigenvalues
eigenvect  A matrix of dimension ("selected samples", eigen.cnt).
pedigree

Author(s)

Xiuwen Zheng, Stephanie Gogarten

References


See Also

SeqVarGDSClass, applyMethod

Examples

```r
gds <- seqOpen(seqExampleFileName("gds"))
pca <- pca(gds)
pca$eigenval
head(pca$eigenvect)
seqClose(gds)
```

---

pedigree  Pedigree for example data

Description

Pedigree for example data files in SeqArray.

Usage

pedigree

Format

A data.frame with the following columns.

- family: Family ID
- individ: Individual ID
- father: Father ID
- mother: Mother ID
- sex: Sex
- sample.id: sample.id in VCF/GDS files

Details

There is one trio in the pedigree.
setVariantID

Source
HapMap

Examples

data(pedigree)
head(pedigree)
gds <- seqOpen(seqExampleFileName("gds"))
setdiff(seqGetData(gds, "sample.id"), pedigree$sample.id)
seqClose(gds)

---

setVariantID (Change the variant ID of a GDS file)

Description
Replace the variable "variant.id" in a GDS file with a user-supplied unique vector of the same length.

Usage
setVariantID(gdsfile, variant.id)

Arguments

- gdsfile: A character string with the file path of a GDS file.
- variant.id: A vector with the new variant IDs.

Details
A VCF file created by seqVCF2GDS creates a variable "variant.id" containing sequential integers to identify each variant. setVariantID allows the user to replace these values with something more meaningful. The replacement values in variant.id must be unique and have the same length as the original "variant.id" vector.

Using character values for variant.id may affect performance for large datasets.

Author(s)
Stephanie Gogarten

See Also
SeqVarGDSClass, seqVCF2GDS
Examples

```r
oldfile <- system.file("extdata", "gl_chr1.gds", package="SeqVarTools")
newfile <- tempfile()
file.copy(oldfile, newfile)

gds <- seqOpen(newfile)
rsID <- seqGetData(gds, "annotation/id")
seqClose(gds)

setVariantID(newfile, rsID)
gds <- seqOpen(newfile)
seqGetData(gds, "variant.id")
head(getGenotype(gds))
seqClose(gds)
unlink(newfile)
```

---

**titv**

*Transition/Transversion Ratio*

**Description**

Calculate transition/transversion ratio overall or by sample

**Usage**

```r
## S4 method for signature 'SeqVarGDSClass'
titv(gdsobj, by.sample=FALSE, use.names=FALSE)
```

**Arguments**

- `gdsobj` A `SeqVarGDSClass` object with VCF data.
- `by.sample` A logical indicating whether TiTv should be calculated by sample or overall for the entire GDS object.
- `use.names` A logical indicating whether to assign sample IDs as names of the output vector (if `by.sample=TRUE`).

**Details**

If `by.sample=FALSE` (the default), `titv` calculates the transition/transversion ratio (TiTv) over all samples.

If `by.sample=TRUE`, `titv` calculates TiTv over all variant genotypes (heterozygous or homozygous non-reference) for each sample.

**Value**

A single value for TiTv if `by.sample=FALSE`. If `by.sample=TRUE`, a numeric vector containing TiTv for each sample.
Author(s)
Stephanie Gogarten

See Also
SeqVarGDSClass, applyMethod, isVariant

Examples

gds <- seqOpen(seqExampleFileName("gds"))
titv(gds)
titv(gds, by.sample=TRUE)

## apply to a subset of variants
library(GenomicRanges)
chrom <- seqGetData(gds, "chromosome")
pos22 <- seqGetData(gds, "position")[chrom == 22]
ranges <- GRanges(seqnames="22", IRanges(min(pos22), max(pos22)))
applyMethod(gds, titv, ranges)

seqClose(gds)
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