

# Package ‘SeqArray’

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**Type** Package

**Title** Data management of large-scale whole-genome sequence variant calls using GDS files

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**LinkingTo** gdsfmt

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**Description** Data management of large-scale whole-genome sequencing variant calls with thousands of individuals: genotypic data (e.g., SNVs, indels and structural variation calls) and annotations in SeqArray GDS files are stored in an array-oriented and compressed manner, with efficient data access using the R programming language.

**License** GPL-3

**VignetteBuilder** knitr

**ByteCompile** TRUE

**LazyData** true

**URL** <https://github.com/zhengxwen/SeqArray>

**BugReports** <https://github.com/zhengxwen/SeqArray/issues>

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SeqArray-package	<i>Data Management of Large-scale Whole-Genome Sequence Variant Calls</i>
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## Description

Data management of large-scale whole-genome sequencing variants.

## Details

As the cost of DNA sequencing rapidly decreases, whole-genome sequencing (WGS) is generating data at an unprecedented rate. Scientists are being challenged to manage data sets that are terabyte-sized, contain diverse types of data and complex data relationships. Data analyses of WGS requires a general file format for storing genetic variants including single nucleotide variations (SNVs), insertions and deletions (indels) and structural variants. The variant call format (VCF) is a generic and flexible format for storing DNA polymorphisms developed for the 1000 Genomes Project that is the standard WGS format in use today. VCF is a textual format usually stored in compressed files that supports rich annotations and relatively efficient data retrieval. However, VCF files are large and the computational burden associated with all data retrieval from text files can be significant for a large WGS study with thousands of samples.

To provide an efficient alternative to VCF for WGS data, we developed a new data format and accompanying Bioconductor package, “SeqArray”. Key features of SeqArray are efficient storage including multiple high compression options, data retrieval by variant or sample subsets, support for parallel access and computing, and C++ integration in the R programming environment. The SeqArray package provides R functions for efficient block-wise computations, and enables scientists to develop custom R scripts for exploratory data analysis.

Webpage: <https://github.com/zhengxwen/SeqArray>, <http://bioconductor.org/packages/SeqArray/>

## Author(s)

Xiuwen Zheng <zhengx@u.washington.edu>

## Examples

```
# the file of VCF
vcf.fn <- seqExampleFileName("vcf")
vcf.fn
# or vcf.fn <- "C:/YourFolder/Your_VCF_File.vcf"

# parse the header
seqVCF_Header(vcf.fn)

# get sample id
seqVCF_SampID(vcf.fn)

# convert
```

```

seqVCF2GDS(vcf.fn, "tmp.gds", storage.option="ZIP_RA")
seqSummary("tmp.gds")

# list the structure of GDS variables
f <- seqOpen("tmp.gds")
f

seqClose(f)
unlink("tmp.gds")

#####

# the GDS file
(gds.fn <- seqExampleFileName("gds"))

# display
(f <- seqOpen(gds.fn))

# get 'sample.id'
(samp.id <- seqGetData(f, "sample.id"))
# "NA06984" "NA06985" "NA06986" ...

# get 'variant.id'
head(variant.id <- seqGetData(f, "variant.id"))

# get 'chromosome'
table(seqGetData(f, "chromosome"))

# get 'allele'
head(seqGetData(f, "allele"))
# "T,C" "G,A" "G,A" ...

# set sample and variant filters
seqSetFilter(f, sample.id=samp.id[c(2,4,6,8,10)])
set.seed(100)
seqSetFilter(f, variant.id=sample(variant.id, 10))

# get genotypic data
seqGetData(f, "genotype")

# get annotation/info/DP
seqGetData(f, "annotation/info/DP")

# get annotation/info/AA, a variable-length dataset
seqGetData(f, "annotation/info/AA")
# $length          <- indicating the length of each variable-length data
# [1] 1 1 1 1 1 1 ...
# $data            <- the data according to $length
# [1] "T" "C" "T" "C" "G" "C" ...

# get annotation/format/DP, a variable-length dataset
seqGetData(f, "annotation/format/DP")
# $length          <- indicating the length of each variable-length data
# [1] 1 1 1 1 1 1 ...
# $data            <- the data according to $length

```

```

#      variant
# sample [,1] [,2] [,3] [,4] [,5] [,6] ...
# [1,]    25    25    22     3     4    17 ...

# read multiple variables variant by variant
seqApply(f, c(geno="genotype", phase="phase", qual="annotation/id"),
         FUN=function(x) print(x), as.is="none")

# get the numbers of alleles per variant
head(seqApply(f, "allele",
             FUN=function(x) length(unlist(strsplit(x,""))), as.is="integer"))
# or
head(seqGetData(f, "$num_allele"))

#####

# remove the sample and variant filters
seqResetFilter(f)

# calculate the frequency of reference allele,
# a faster version could be obtained by C coding
af <- seqApply(f, "genotype", FUN=function(x) mean(x=="L", na.rm=TRUE),
              as.is="double")
length(af)
summary(af)

# close the GDS file
seqClose(f)

```

---

KG\_P1\_SampData

*Simulated sample data for 1000 Genomes Phase 1*


---

## Description

An AnnotatedDataFrame with columns sample.id, sex, age, and phenotype, where the identifiers in sample.id match those in the SeqArray file.

## Usage

```
KG_P1_SampData
```

## Value

An AnnotatedDataFrame

---

seqAddValue                      *Add values to a GDS File*

---

### Description

Add or modify the values in a GDS file with hash code

### Usage

```
seqAddValue(gdsfile, varnm, val, desp=character(), replace=FALSE, compress="LZMA_RA",
  packed=TRUE, packed.idx=TRUE, use_float32=TRUE, verbose=TRUE, verbose.attr=TRUE)
```

### Arguments

gdsfile	character for file name, or a <a href="#">SeqVarGDSClass</a> object
varnm	the variable name, e.g., "sample.id", "variant.id", "chromosome", "annotation/info/NEW_VARIABLE"
val	the R value can be integers, real numbers, characters, factor, logical, raw variable, data.frame or a list; a list of vectors is used for variable-length annotation data; or NULL for adding a new folder
desp	variable description
replace	if TRUE, replace the existing variable silently if possible
compress	the compression method can be "" (no compression), see <a href="#">add.gdsn</a>
packed	TRUE, pack data if there is any missing value
packed.idx	TRUE, store the index variable using integers with the fewest bits if possible
use_float32	if TRUE, use 32 bits instead of 64 bits to store floating numeric values (i.e., double)
verbose	if TRUE, show information
verbose.attr	if TRUE, show attribute information in a GDS node

### Value

Return none.

### Author(s)

Xiuwen Zheng

### See Also

[seqVCF2GDS](#), [seqNewVarData](#), [add.gdsn](#)

### Examples

```
# the file of GDS
gds.fn <- seqExampleFileName("gds")
file.copy(gds.fn, "tmp.gds", overwrite=TRUE)

# display
(f <- seqOpen("tmp.gds", readonly=FALSE))
```

```

show(index.gdsn(f, "sample.id"))
seqAddValue(f, "sample.id", 1:90, replace=TRUE)
show(index.gdsn(f, "sample.id"))

show(index.gdsn(f, "chromosome"))
v <- seqGetData(f, "chromosome")
seqAddValue(f, "chromosome", paste0("chr", v), replace=TRUE)
show(index.gdsn(f, "chromosome"))
table(seqGetData(f, "chromosome"))

# annotation info
seqAddValue(f, "annotation/info/folder", NULL) # add a new folder
seqAddValue(f, "annotation/info/folder/val", 1:1348, "random number")
seqAddValue(f, "annotation/info/folder/logical", rep(c(TRUE, FALSE), length.out=1348))
seqAddValue(f, "annotation/info/folder/packed", c(rep(2L, 1000), rep(NA, 348)))
seqAddValue(f, "annotation/info/newff",
  data.frame(x=1:1348, y=rep("s", 1348), stringsAsFactors=FALSE),
  desp=c("integer numbers", "character"))

# variable-length annotation info data
v <- lapply(1:1348, function(x) as.character(x))
v[[1]] <- 1:10
seqAddValue(f, "annotation/info/folder/val1", v)
head(seqGetData(f, "annotation/info/folder/val1", .tolist=TRUE))

# sample annotation
seqAddValue(f, "sample.annotation", data.frame(ii=1:90, y=rep("A", 90)),
  replace=TRUE)
seqAddValue(f, "sample.annotation/float", (1:90)/90)

# close the GDS file
seqClose(f)

# remove the temporary file
unlink("tmp.gds", force=TRUE)

```

---

seqAlleleFreq

*Get Allele Frequencies or Counts*


---

## Description

Calculates the allele frequencies or counts for reference or minor alleles.

## Usage

```

seqAlleleFreq(gdsfile, ref.allele=0L, minor=FALSE, parallel=seqGetParallel(),
  balancing=NA, verbose=FALSE)
seqAlleleCount(gdsfile, ref.allele=0L, minor=FALSE, parallel=seqGetParallel(),
  balancing=NA, verbose=FALSE)
seqGetAF_AC_Missing(gdsfile, minor=FALSE, alt=FALSE, ns=FALSE,
  parallel=seqGetParallel(), balancing=NA, verbose=FALSE)

```

**Arguments**

<code>gdsfile</code>	a <a href="#">SeqVarGDSCClass</a> object
<code>ref.allele</code>	NULL, a single numeric value, a numeric vector or a character vector; see <a href="#">Value</a>
<code>minor</code>	if TRUE, return minor allele frequency/count
<code>alt</code>	if TRUE and <code>minor=FALSE</code> , return the frequencies and counts for the alternative alleles in <code>seqGetAF_AC_Missing</code> ; otherwise, return the frequencies and counts for the reference alleles; if there are more than two alleles at a site, the alternative alleles will be collapsed
<code>ns</code>	if TRUE, return the numbers of samples without missing genotypes in the columns, used in <code>seqGetAF_AC_Missing</code>
<code>parallel</code>	FALSE (serial processing), TRUE (multicore processing), numeric value or other value; <code>parallel</code> is passed to the argument <code>cl</code> in <a href="#">seqParallel</a> , see <a href="#">seqParallel</a> for more details.
<code>balancing</code>	whether to perform workload balancing or not, only applicable when multiple cores are used; if NA, use TRUE as a default until <code>getOption("seqarray.balancing")</code> is set and not TRUE
<code>verbose</code>	if TRUE, show progress information

**Details**

If the `gds` node 'genotype/data' (integer genotypes) is not available, the node 'annotation/format/DS' (numeric genotype dosages for alternative alleles) will be used to calculate allele frequencies. At a site, it assumes 'annotation/format/DS' stores the dosage of the 1st alternative allele in the 1st column, 2nd alt. allele in the 2nd column if it is multi-allelic, and so on.

**Value**

If `ref.allele=NULL`, the function returns a list of allele frequencies/counts according to all allele per site. If `ref.allele` is a single numeric value (like 0L), it returns a numeric/integer vector for the specified allele (0L for the reference allele, 1L for the first alternative allele, etc). If `ref.allele` is a numeric vector, `ref.allele` specifies each allele per site. If `ref.allele` is a character vector, `ref.allele` specifies the desired allele for each site (e.g. ancestral allele for the derived allele frequency/count).

`seqGetAF_AC_Missing()` returns `data.frame(af, ac, miss)` for allele frequencies, allele counts and missing rates. It is faster than calling `seqAlleleFreq()`, `seqAlleleCount()` and `seqMissing` sequentially.

**Author(s)**

Xiuwen Zheng

**See Also**

[seqMissing](#), [seqNumAllele](#), [seqParallel](#), [seqGetParallel](#)

**Examples**

```
# the GDS file
(gds.fn <- seqExampleFileName("gds"))

# display
```

```

f <- seqOpen(gds.fn)

# return a list
head(seqAlleleFreq(f, NULL, verbose=TRUE))

# return a numeric vector
summary(seqAlleleFreq(f, 0L, verbose=TRUE))

# return a numeric vector
summary(seqAlleleFreq(f, 0L, minor=TRUE, verbose=TRUE))

# return a numeric vector, AA is ancestral allele
AA <- seqGetData(f, "annotation/info/AA", .padNA=TRUE)
summary(seqAlleleFreq(f, AA))
summary(seqAlleleFreq(f, AA, minor=TRUE))

# allele counts
head(seqAlleleCount(f, NULL, verbose=TRUE))
head(seqAlleleCount(f, 0L, verbose=TRUE))
head(seqAlleleCount(f, 0L, minor=TRUE, verbose=TRUE))
head(seqAlleleCount(f, AA, verbose=TRUE))
head(seqAlleleCount(f, AA, minor=TRUE, verbose=TRUE))

# allele frequencies, allele counts and missing proportions
v <- seqGetAF_AC_Missing(f, minor=TRUE)
head(v)

# close the GDS file
seqClose(f)

# A GDS file with imputed dosages
ds_fn <- seqExampleFileName("dosage")
ds_fn

str(seqGetAF_AC_Missing(ds_fn, alt=FALSE))
str(seqGetAF_AC_Missing(ds_fn, alt=TRUE))
str(seqGetAF_AC_Missing(ds_fn, minor=TRUE))

```

---

seqApply

*Apply Functions Over Array Margins*


---

## Description

Returns a vector or list of values obtained by applying a function to margins of genotypes and annotations.

## Usage

```

seqApply(gdsfile, var.name, FUN, margin=c("by.variant", "by.sample"),
  as.is=c("none", "list", "integer", "double", "character", "logical", "raw"),
  var.index=c("none", "relative", "absolute"), parallel=FALSE,
  .useraw=FALSE, .progress=FALSE, .list_dup=TRUE, .balancing=FALSE, ...)

```

**Arguments**

<code>gdsfile</code>	a <a href="#">SeqVarGDSClass</a> object or a GDS file name
<code>var.name</code>	the variable name(s), see details
<code>FUN</code>	the function to be applied
<code>margin</code>	giving the dimension which the function will be applied over; <code>margin="by.variant"</code> by default
<code>as.is</code>	returned value: a list, an integer vector, etc; return nothing by default <code>as.is="none"</code> ; <code>as.is</code> can be a <a href="#">connection</a> object, or a GDS node <a href="#">gdsn.class</a> object; if "unlist" is used, produces a vector which contains all the atomic components, via <code>unlist(..., recursive=FALSE)</code>
<code>var.index</code>	if "none" (by default), call <code>FUN(x, ...)</code> without variable index; if "relative" or "absolute", add an argument to the user-defined function <code>FUN</code> like <code>FUN(index, x, ...)</code> where <code>index</code> is an index of variant starting from 1 if <code>margin = "by.variant"</code> : "relative" for indexing in the selection defined by <a href="#">seqSetFilter</a> , "absolute" for indexing with respect to all data
<code>parallel</code>	FALSE (serial processing), TRUE (multicore processing), numeric value or other value; <code>parallel</code> is passed to the argument <code>cl</code> in <a href="#">seqParallel</a> , see <a href="#">seqParallel</a> for more details.
<code>.useraw</code>	TRUE, force to use RAW instead of INTEGER for genotypes and dosages; FALSE, use INTEGER; NA, use RAW for small numbers instead of INTEGER if possible, it is needed to detect data type (RAW or INTEGER) in the user-defined function; for genotypes, 0xFF is missing value if RAW is used
<code>.progress</code>	if TRUE, show progress information; or a file name for updating progress information
<code>.list_dup</code>	internal use only
<code>.balancing</code>	whether to perform workload balancing or not, only applicable when multiple cores are used; if NA, use TRUE as a default until <code>getOption("seqarray.balancing")</code> is set and not TRUE
<code>...</code>	optional arguments to <code>FUN</code>

**Details**

The variable name should be "sample.id", "variant.id", "position", "chromosome", "allele", "genotype", "annotation/id", "annotation/qual", "annotation/filter", "annotation/info/VARIABLE\_NAME" or "annotation/format/VARIABLE\_NAME".

"@genotype", "annotation/info/@VARIABLE\_NAME" or "annotation/format/@VARIABLE\_NAME" are used to obtain the index associated with these variables.

"\$dosage" is also allowed for the dosages of reference allele (integer: 0, 1, 2 and NA for diploid genotypes).

"\$dosage\_alt" returns a RAW/INTEGER matrix for the dosages of alternative allele without distinguishing different alternative alleles.

"\$num\_allele" returns an integer vector with the numbers of distinct alleles.

"\$ref" returns a character vector of reference alleles

"\$alt" returns a character vector of alternative alleles (delimited by comma)

"\$chrom\_pos" returns characters with the combination of chromosome and position, e.g., "1:1272721".

"\$chrom\_pos\_allele" returns characters with the combination of chromosome, position and alleles, e.g., "1:1272721\_A\_G" (i.e., chr:position\_REF\_ALT).

The algorithm is highly optimized by blocking the computations to exploit the high-speed memory instead of disk.

**Value**

A vector, a list of values or none.

**Author(s)**

Xiuwen Zheng

**See Also**

[seqBlockApply](#), [seqSetFilter](#), [seqGetData](#), [seqParallel](#), [seqGetParallel](#)

**Examples**

```
# the GDS file
(gds.fn <- seqExampleFileName("gds"))

# display
(f <- seqOpen(gds.fn))

# get 'sample.id'
(samp.id <- seqGetData(f, "sample.id"))
# "NA06984" "NA06985" "NA06986" ...

# get 'variant.id'
head(variant.id <- seqGetData(f, "variant.id"))

# set sample and variant filters
set.seed(100)
seqSetFilter(f, sample.id=samp.id[c(2,4,6,8,10)],
            variant.id=sample(variant.id, 10))

# read
seqApply(f, "genotype", FUN=print, margin="by.variant")
seqApply(f, "genotype", FUN=print, margin="by.variant", .useraw=TRUE)

seqApply(f, "genotype", FUN=print, margin="by.sample")
seqApply(f, "genotype", FUN=print, margin="by.sample", .useraw=TRUE)

# read multiple variables variant by variant
seqApply(f, c(geno="genotype", phase="phase", rsid="annotation/id",
            DP="annotation/format/DP"), FUN=print, as.is="none")

# get the numbers of alleles per variant
seqApply(f, "allele",
        FUN=function(x) length(unlist(strsplit(x,","))), as.is="integer")

# output to a file
fl <- file("tmp.txt", "wt")
seqApply(f, "genotype", FUN=sum, na.rm=TRUE, as.is=fl)
close(fl)
readLines("tmp.txt")
```

```

seqApply(f, "genotype", FUN=sum, na.rm=TRUE, as.is=stdout())
seqApply(f, "genotype", FUN=sum, na.rm=TRUE, as.is="integer")
# should be identical

#####
# with an index of variant

seqApply(f, c(geno="genotype", phase="phase", rsid="annotation/id"),
  FUN=function(index, x) { print(index); print(x); index },
  as.is="integer", var.index="relative")
# it is as the same as
which(seqGetFilter(f)$variant.sel)

#####
# reset sample and variant filters
seqResetFilter(f)

# calculate the frequency of reference allele,
# a faster version could be obtained by C coding
af <- seqApply(f, "genotype", FUN=function(x) mean(x=="L", na.rm=TRUE),
  as.is="double")
length(af)
summary(af)

#####
# apply the user-defined function sample by sample

# reset sample and variant filters
seqResetFilter(f)
summary(seqApply(f, "genotype", FUN=function(x) { mean(is.na(x)) },
  margin="by.sample", as.is="double"))

# set sample and variant filters
set.seed(100)
seqSetFilter(f, sample.id=samp.id[c(2,4,6,8,10)],
  variant.id=sample(variant.id, 10))

seqApply(f, "genotype", FUN=print, margin="by.variant", as.is="none")

seqApply(f, "genotype", FUN=print, margin="by.sample", as.is="none")

seqApply(f, c(sample.id="sample.id", genotype="genotype"), FUN=print,
  margin="by.sample", as.is="none")

# close the GDS file
seqClose(f)

# delete the temporary file

```

```
unlink("tmp.txt")
```

---

seqAsVCF

*VariantAnnotation objects*

---

## Description

Create a [VCF-class](#) object

## Usage

```
seqAsVCF(x, chr.prefix="", info=NULL, geno=NULL)
```

## Arguments

x	a <a href="#">SeqVarGDSClass</a> object
chr.prefix	prefix to add to seqlevels
info	which INFO fields to return
geno	which GENO fields to return

## Details

Coerces a [SeqVarGDSClass](#) object to a [VCF-class](#) object. Row names correspond to the variant.id. info and geno specify the 'INFO' and 'GENO' (FORMAT) fields to return, respectively. If not specified, all fields are returned; if 'NA' no fields are returned. Use [seqSetFilter](#) prior to calling seqAsVCF to specify samples and variants to return.

The **VariantAnnotation** package should be loaded to explore this object.

## Value

A [CollapsedVCF](#) object.

## Author(s)

Stephanie Gogarten, Xiuwen Zheng

## See Also

[VCF-class](#)

## Examples

```
gds <- seqOpen(seqExampleFileName("gds"))

## Not run:
library(VariantAnnotation)
seqAsVCF(gds)

## End(Not run)

seqClose(gds)
```

seqBED2GDS

*Conversion between PLINK BED and SeqArray GDS***Description**

Conversion between PLINK BED format and SeqArray GDS format.

**Usage**

```
seqBED2GDS(bed.fn, fam.fn, bim.fn, out.gdsfn, compress.geno="LZMA_RA",
           compress.annotation="LZMA_RA", chr.conv=TRUE, include.pheno=TRUE,
           optimize=TRUE, digest=TRUE, parallel=FALSE, verbose=TRUE)
seqGDS2BED(gdsfile, out.fn, write.rsid=c("auto", "annot_id", "chr_pos_ref_alt"),
           multi.row=FALSE, verbose=TRUE)
```

**Arguments**

bed.fn	the file name of PLINK binary file, genotype information
fam.fn	the file name of first six columns of ".ped", sample or family information; if missing, determine the file name using bed.fn
bim.fn	the file name of extended MAP file with 6 columns, variant information; if missing, determine the file name using bed.fn
gdsfile	character (a GDS file name), or a <a href="#">SeqVarGDSClass</a> object
out.gdsfn	the file name, output a file of SeqArray format
out.fn	the file name of PLINK binary format without extended names
compress.geno	the compression method for "genotype"; optional values are defined in the function <code>add.gdsn</code>
compress.annotation	the compression method for the GDS variables, except "genotype"; optional values are defined in the function <code>add.gdsn</code>
chr.conv	if TRUE, convert numeric chromosome codes 23 to X, 24 to Y, 25 to XY, and 26 to MT
include.pheno	if TRUE, add 'family', 'father', 'mother', 'sex' and 'phenotype' in the FAM file to the output GDS file; FALSE for no phenotype; or a character vector to specify which of the family, father, mother, sex and phenotype variables to be added
optimize	if TRUE, optimize the access efficiency by calling <a href="#">cleanup.gds</a>
digest	a logical value (TRUE/FALSE) or a character ("md5", "sha1", "sha256", "sha384" or "sha512"); add hash codes to the GDS file if TRUE or a digest algorithm is specified
parallel	FALSE (serial processing), TRUE (parallel processing), a numeric value indicating the number of cores, or a cluster object for parallel processing; <code>parallel</code> is passed to the argument <code>cl</code> in <a href="#">seqParallel</a> , see <a href="#">seqParallel</a> for more details
write.rsid	"annot_id": use the node "annotation/id" for the variant IDs; "chr_pos_ref_alt": use the format "chrom_position_ref_alt"; "auto": use "annotation/id" for the variant IDs if it is not a blank string or ".", otherwise use "chrom_position_ref_alt"
multi.row	if TRUE, a multiallelic site is converted to multiple rows in PLINK bim and bed files
verbose	if TRUE, show information

**Value**

Return the file name of SeqArray file with an absolute path.

**Author(s)**

Xiuwen Zheng

**See Also**

[seqSNP2GDS](#), [seqVCF2GDS](#)

**Examples**

```
library(SNPRelate)

# PLINK BED files
bed.fn <- system.file("extdata", "plinkhapmap.bed.gz", package="SNPRelate")
fam.fn <- system.file("extdata", "plinkhapmap.fam.gz", package="SNPRelate")
bim.fn <- system.file("extdata", "plinkhapmap.bim.gz", package="SNPRelate")

# convert bed to gds
seqBED2GDS(bed.fn, fam.fn, bim.fn, "tmp.gds")

seqSummary("tmp.gds")

# convert gds to bed
gdsfn <- seqExampleFileName("gds")
seqGDS2BED(gdsfn, "plink")

# remove the temporary file
unlink(c("tmp.gds", "plink.fam", "plink.bim", "plink.bed"), force=TRUE)
```

---

seqBlockApply

*Apply Functions Over Array Margins via Blocking*

---

**Description**

Returns a vector or list of values obtained by applying a function to margins of genotypes and annotations via blocking.

**Usage**

```
seqBlockApply(gdsfile, var.name, FUN, margin=c("by.variant"),
  as.is=c("none", "list", "unlist"), var.index=c("none", "relative", "absolute"),
  bsize=1024L, parallel=FALSE, .useraw=FALSE, .padNA=TRUE, .tolist=FALSE,
  .balancing=FALSE, .progress=FALSE, ...)
```

**Arguments**

<code>gdsfile</code>	a <a href="#">SeqVarGDSClass</a> object or a GDS file name
<code>var.name</code>	the variable name(s), see details
<code>FUN</code>	the function to be applied
<code>margin</code>	giving the dimension which the function will be applied over
<code>as.is</code>	returned value: a list, an integer vector, etc; return nothing by default as <code>is="none"</code> ; <code>as.is</code> can be a <a href="#">connection</a> object, or a GDS node <a href="#">gdsn.class</a> object; if "unlist" is used, produces a vector which contains all the atomic components, via <code>unlist(..., recursive=FALSE)</code>
<code>var.index</code>	if "none" (by default), call <code>FUN(x, ...)</code> without variable index; if "relative" or "absolute", add an argument to the user-defined function <code>FUN</code> like <code>FUN(index, x, ...)</code> where <code>index</code> is an index of variant starting from 1 if <code>margin="by.variant"</code> : "relative" for indexing in the selection defined by <a href="#">seqSetFilter</a> , "absolute" for indexing with respect to all data
<code>bsize</code>	block size
<code>parallel</code>	FALSE (serial processing), TRUE (multicore processing), numeric value or other value; <code>parallel</code> is passed to the argument <code>cl</code> in <a href="#">seqParallel</a> , see <a href="#">seqParallel</a> for more details.
<code>.useraw</code>	TRUE, force to use RAW instead of INTEGER for genotypes and dosages; FALSE, use INTEGER; NA, use RAW instead of INTEGER if possible; for genotypes, 0xFF is missing value if RAW is used
<code>.padNA</code>	TRUE, pad a variable-length vector with NA if the number of data points for each variant is not greater than 1
<code>.tolist</code>	if TRUE, return a list of vectors instead of the structure <code>list(length, data)</code> for variable-length data; NA, return a compressed List defined in <b>IRanges</b> when it is applicable
<code>.balancing</code>	whether to perform workload balancing or not, only applicable when multiple cores are used; if NA, use TRUE as a default until <code>getOption("seqarray.balancing")</code> is set and not TRUE
<code>.progress</code>	if TRUE, show progress information
<code>...</code>	optional arguments to <code>FUN</code>

**Details**

The variable name should be "sample.id", "variant.id", "position", "chromosome", "allele", "genotype", "annotation/id", "annotation/qual", "annotation/filter", "annotation/info/VARIABLE\_NAME" or "annotation/format/VARIABLE\_NAME".

"@genotype", "annotation/info/@VARIABLE\_NAME" or "annotation/format/@VARIABLE\_NAME" are used to obtain the index associated with these variables.

"\$chromosome" returns chromosome codes in an object of `S4Vectors::Rle`.

"\$dosage" is also allowed for the dosages of reference allele (integer: 0, 1, 2 and NA for diploid genotypes).

"\$dosage\_alt" returns a RAW/INTEGER matrix for the dosages of alternative allele without distinguishing different alternative alleles. "\$dosage\_alt2" allow the alleles are partially missing (e.g., genotypes on chromosome X for males)

"\$dosage\_sp" returns a sparse matrix (dgCMatrix) for the dosages of alternative allele without distinguishing different alternative alleles. "\$dosage\_sp2" allow the alleles are partially missing (e.g., genotypes on chromosome X for males)

"\$num\_allele" returns an integer vector with the numbers of distinct alleles.

"\$ref" returns a character vector of reference alleles. "\$alt" returns a character vector of alternative alleles (delimited by comma).

"\$chrom\_pos" returns characters with the combination of chromosome and position, e.g., "1:1272721".

"\$chrom\_pos2" is similar to "\$chrom\_pos", except the suffix "\_1" is added to the first duplicate following the variant, "\_2" is added to the second duplicate, and so on. "\$chrom\_pos\_allele" returns characters with the combination of chromosome, position and alleles, e.g., "1:1272721\_A\_G" (i.e., chr:position\_REF\_ALT).

"\$variant\_index" returns the indices of selected variants starting from 1, and "\$sample\_index" returns the indices of selected samples starting from 1.

The algorithm is highly optimized by blocking the computations to exploit the high-speed memory instead of disk.

### Value

A vector, a list of values or none.

### Author(s)

Xiuwen Zheng

### See Also

[seqApply](#), [seqSetFilter](#), [seqGetData](#), [seqParallel](#), [seqGetParallel](#)

### Examples

```
# the GDS file
(gds.fn <- seqExampleFileName("gds"))

# display
(f <- seqOpen(gds.fn))

# get 'sample.id'
(samp.id <- seqGetData(f, "sample.id"))
# "NA06984" "NA06985" "NA06986" ...

# get 'variant.id'
head(variant.id <- seqGetData(f, "variant.id"))

# set sample and variant filters
set.seed(100)
seqSetFilter(f, sample.id=samp.id[c(2,4,6,8,10)],
            variant.id=sample(variant.id, 10))

# read in block
seqGetData(f, "$dosage")
seqBlockApply(f, "$dosage", print, bsize=3)
seqBlockApply(f, "$dosage", function(x) x, as.is="list", bsize=3)
seqBlockApply(f, c(dos="$dosage", pos="position"), print, bsize=3)
```

```
# close the GDS file
seqClose(f)
```

---

seqCheck

*Data Integrity Checking*

---

### Description

Performs data integrity on a SeqArray GDS file.

### Usage

```
seqCheck(gdsfile, verbose=TRUE)
```

### Arguments

gdsfile	a <a href="#">SeqVarGDSClass</a> object, or a file name
verbose	if TRUE, display information

### Value

A list of the following components:

hash	a data.frame for hash checking, including algo for digest algorithms and ok for the checking states
dimension	a data.frame for checking the dimension of each variable, including ok for the checking states and info for the error messages

### Author(s)

Xiuwen Zheng

### Examples

```
# the GDS file
(gds.fn <- seqExampleFileName("gds"))

seqCheck(gds.fn)
```

---

seqClose-methods	<i>Close the SeqArray GDS File</i>
------------------	------------------------------------

---

**Description**

Closes a SeqArray GDS file which is open.

**Usage**

```
## S4 method for signature 'gds.class'
seqClose(object)
## S4 method for signature 'SeqVarGDSClass'
seqClose(object)
```

**Arguments**

object            a SeqArray object

**Details**

If object is

- [gds.class](#), close a general GDS file
- [SeqVarGDSClass](#), close the sequence GDS file.

**Value**

None.

**Author(s)**

Xiuwen Zheng

**See Also**

[seqOpen](#)

---

seqDelete	<i>Delete GDS Variables</i>
-----------	-----------------------------

---

**Description**

Deletes variables in the SeqArray GDS file.

**Usage**

```
seqDelete(gdsfile, info.var=character(), fmt.var=character(),
          samp.var=character(), verbose=TRUE)
```

**Arguments**

<code>gdsfile</code>	a <a href="#">SeqVarGDSClass</a> object
<code>info.var</code>	the variables in the INFO field, i.e., "annotation/info/VARIABLE_NAME"
<code>fmt.var</code>	the variables in the FORMAT field, i.e., "annotation/format/VARIABLE_NAME"
<code>samp.var</code>	the variables in the sample annotation field, i.e., "sample.annotation/VARIABLE_NAME"
<code>verbose</code>	if TRUE, show information

**Value**

None.

**Author(s)**

Xiuwen Zheng

**See Also**

[seqOpen](#), [seqClose](#)

**Examples**

```
# the file of GDS
gds.fn <- seqExampleFileName("gds")
file.copy(gds.fn, "tmp.gds", overwrite=TRUE)

# display
(f <- seqOpen("tmp.gds", FALSE))

seqDelete(f, info.var=c("HM2", "AA"), fmt.var="DP")
f

# close the GDS file
seqClose(f)

# clean up the fragments, reduce the file size
cleanup.gds("tmp.gds")

# remove the temporary file
unlink("tmp.gds", force=TRUE)
```

---

seqDigest

*Hash function digests*

---

**Description**

Create hash function digests for all or a subset of data

**Usage**

```
seqDigest(gdsfile, varname, algo=c("md5"), parallel=FALSE, verbose=FALSE)
```

## Arguments

gdsfile	a <a href="#">SeqVarGDSClass</a> object or a GDS file name
varname	the variable name(s), see details
algo	the digest hash algorithm: "md5"
parallel	FALSE (serial processing), TRUE (multicore processing), numeric value or other value; parallel is passed to the argument cl in <a href="#">seqParallel</a> , see <a href="#">seqParallel</a> for more details.
verbose	if TRUE, show progress information

## Details

The variable name should be "sample.id", "variant.id", "position", "chromosome", "allele", "annotation/id", "annotation/qual", "annotation/filter", "annotation/info/VARIABLE\_NAME", or "annotation/format/VARIABLE\_NAME".

Users can define a subset of data via [seqSetFilter](#) and create a hash digest for the subset only.

Note that the hashing algorithm is not parallelizable. When multiple processes are used, the hash codes are calculated for each chunk of data, and then call `digest()` to obtain the final hash code. Hence, the returned hash codes are different when comparing `parallel=FALSE` and a multicore implementation.

## Value

A hash character.

## Author(s)

Xiuwen Zheng

## See Also

[seqSetFilter](#), [seqApply](#)

## Examples

```
# the GDS file
(gds.fn <- seqExampleFileName("gds"))

# display
f <- seqOpen(gds.fn)

seqDigest(f, "genotype")
seqDigest(f, "annotation/filter")
seqDigest(f, "annotation/format/DP")

# close the GDS file
seqClose(f)
```

---

seqEmptyFile	<i>Empty GDS file</i>
--------------	-----------------------

---

**Description**

Create a new empty GDS file.

**Usage**

```
seqEmptyFile(outfn, sample.id=character(), numvariant=1L, verbose=TRUE)
```

**Arguments**

outfn	the output file name for a GDS file
sample.id	a list of sample IDs
numvariant	the number of variants
verbose	if TRUE, show information

**Value**

None.

**Author(s)**

Xiuwen Zheng

**See Also**

[seqVCF2GDS](#)

**Examples**

```
seqEmptyFile("tmp.gds")

(f <- seqOpen("tmp.gds"))
seqClose(f)

# remove the temporary file
unlink("tmp.gds", force=TRUE)
```

---

seqExampleFileName	<i>Example files</i>
--------------------	----------------------

---

**Description**

The example files of VCF and GDS format.

**Usage**

```
seqExampleFileName(type=c("gds", "vcf", "KG_Phase1", "dosage"))
```

**Arguments**

type "gds" (by default), "vcf", "KG\_Phase1" or "dosage"

**Value**

Return the path of a VCF file in the package if type="vcf", or the path of a GDS file if type="gds"; if type="KG\_Phase1", return the path of GDS file on Chromosome 22 of the 1000 Genomes Phase 1 project; if type="dosage", return the path of GDS file with imputed allele dosages.

**Author(s)**

Xiuwen Zheng

**Examples**

```
seqExampleFileName("gds")  
seqExampleFileName("vcf")  
seqExampleFileName("KG_Phase1")  
seqExampleFileName("dosage")
```

---

seqExport	<i>Export to a GDS File</i>
-----------	-----------------------------

---

**Description**

Exports to a GDS file with selected samples and variants, which are defined by seqSetFilter().

**Usage**

```
seqExport(gdsfile, out.fn, info.var=NULL, fmt.var=NULL, samp.var=NULL,  
optimize=TRUE, digest=TRUE, verbose=TRUE, verbose.clean=NA)
```

**Arguments**

<code>gdsfile</code>	a <a href="#">SeqVarGDSClass</a> object
<code>out.fn</code>	the file name of output GDS file
<code>info.var</code>	characters, the variable name(s) in the INFO field for import; or NULL for all variables
<code>fmt.var</code>	characters, the variable name(s) in the FORMAT field for import; or NULL for all variables
<code>samp.var</code>	characters, the variable name(s) in the folder "sample.annotation"
<code>optimize</code>	if TRUE, optimize the access efficiency by calling <a href="#">cleanup.gds</a>
<code>digest</code>	a logical value (TRUE/FALSE) or a character ("md5", "sha1", "sha256", "sha384" or "sha512"); add md5 hash codes to the GDS file if TRUE or a digest algorithm is specified
<code>verbose</code>	if TRUE, show information
<code>verbose.clean</code>	when <code>verbose.clean=NA</code> , set it to <code>verbose</code> ; whether display information when calling <code>cleanup.gds</code> or not; only applicable when <code>optimize=TRUE</code>

**Value**

Return the file name of GDS format with an absolute path.

**Author(s)**

Xiuwen Zheng

**See Also**

[seqVCF2GDS](#), [cleanup.gds](#)

**Examples**

```
# open the GDS file
(gds.fn <- seqExampleFileName("gds"))
(f <- seqOpen(gds.fn))

# get 'sample.id'
head(samp.id <- seqGetData(f, "sample.id"))

# get 'variant.id'
head(variant.id <- seqGetData(f, "variant.id"))

set.seed(100)
# set sample and variant filters
seqSetFilter(f, sample.id=samp.id[c(2,4,6,8,10,12,14,16)])
seqSetFilter(f, variant.id=sample(variant.id, 100))

# export
seqExport(f, "tmp.gds")
seqExport(f, "tmp.gds", info.var=character())
seqExport(f, "tmp.gds", fmt.var=character())
seqExport(f, "tmp.gds", samp.var=character())
```

```
# show file
(f1 <- seqOpen("tmp.gds")); seqClose(f1)

# close
seqClose(f)

# delete the temporary file
unlink("tmp.gds")
```

seqGDS2SNP

*Convert to a SNP GDS File***Description**

Converts a SeqArray GDS file to a SNP GDS file.

**Usage**

```
seqGDS2SNP(gdsfile, out.gdsfn, dosage=FALSE, compress.geno="LZMA_RA",
  compress.annotation="LZMA_RA", ds.type=c("packedreal16", "float", "double"),
  optimize=TRUE, verbose=TRUE)
```

**Arguments**

<code>gdsfile</code>	character (a GDS file name), or a <a href="#">SeqVarGDSClass</a> object
<code>out.gdsfn</code>	the file name, output a file of VCF format
<code>dosage</code>	a logical value, or characters for the variable name of dosage in the SeqArray file; if FALSE exports genotypes, otherwise exports dosages
<code>compress.geno</code>	the compression method for "genotype"; optional values are defined in the function <a href="#">add.gdsn</a>
<code>compress.annotation</code>	the compression method for the GDS variables, except "genotype"; optional values are defined in the function <a href="#">add.gdsn</a>
<code>ds.type</code>	applicable when import dosages, the data type for storing dosages; see <a href="#">add.gdsn</a> ; <code>ds.type="packedreal16"</code> by default
<code>optimize</code>	if TRUE, optimize the access efficiency by calling <a href="#">cleanup.gds</a>
<code>verbose</code>	if TRUE, show information

**Details**

[seqSetFilter](#) can be used to define a subset of data for the conversion.

**Value**

Return the file name of VCF file with an absolute path.

**Author(s)**

Xiuwen Zheng

**See Also**

[seqSNP2GDS](#), [seqVCF2GDS](#), [seqGDS2VCF](#)

**Examples**

```
# the GDS file
gds.fn <- seqExampleFileName("gds")

seqGDS2SNP(gds.fn, "tmp.gds")

# delete the temporary file
unlink("tmp.gds")
```

---

seqGDS2VCF

*Convert to a VCF File*


---

**Description**

Converts a SeqArray GDS file to a Variant Call Format (VCF) file.

**Usage**

```
seqGDS2VCF(gdsfile, vcf.fn, info.var=NULL, fmt.var=NULL, chr_prefix="",
  header=TRUE, no_sample=NA, use_Rsamtools=TRUE, verbose=TRUE,
  verbose.progress=NA)
```

**Arguments**

<code>gdsfile</code>	a <a href="#">SeqVarGDSClass</a> object
<code>vcf.fn</code>	the file name, output a file of VCF format; or a <a href="#">connection</a> object
<code>info.var</code>	a list of variable names in the INFO field, or NULL for using all variables; character(0) for no variable in the INFO field
<code>fmt.var</code>	a list of variable names in the FORMAT field, or NULL for using all variables; character(0) for no variable in the FORMAT field
<code>chr_prefix</code>	the prefix of chromosome, e.g., "chr"; no prefix by default
<code>header</code>	TRUE for exporting the metadata header and the header line containing mandatory column names
<code>no_sample</code>	TRUE to exclude the sample-level data and create a site-only VCF file
<code>use_Rsamtools</code>	TRUE for loading the Rsamtools package, see details
<code>verbose</code>	if TRUE, show information
<code>verbose.progress</code>	if NA, set it to verbose; if TRUE, display the progress for each variant

**Details**

[seqSetFilter](#) can be used to define a subset of data for the export.

If the filename extension is "gz" or "bgz", the gzip compression algorithm will be used to compress the output data. When the Rsamtools package is installed and `use_Rsamtools=TRUE`, the exported file utilizes the bgzf format ([bgzip](#), a variant of gzip format) allowing for fast indexing. `bzfile` or `xzfile` will be used, if the filename extension is "bz" or "xz".

**Value**

Return the file name of VCF file with an absolute path.

**Author(s)**

Xiuwen Zheng

**References**

Danecek, P., Auton, A., Abecasis, G., Albers, C.A., Banks, E., DePristo, M.A., Handsaker, R.E., Lunter, G., Marth, G.T., Sherry, S.T., et al. (2011). The variant call format and VCFtools. *Bioinformatics* 27, 2156-2158.

**See Also**

[seqVCF2GDS](#)

**Examples**

```
# the GDS file
(gds.fn <- seqExampleFileName("gds"))

# display
(f <- seqOpen(gds.fn))

# output the first 10 samples
samp.id <- seqGetData(f, "sample.id")
seqSetFilter(f, sample.id=samp.id[1:5])

# convert
seqGDS2VCF(f, "tmp.vcf.gz")

# no INFO and FORMAT
seqGDS2VCF(f, "tmp1.vcf.gz", info.var=character(), fmt.var=character())

# output BN,GP,AA,DP,HM2 in INFO (the variables are in this order), no FORMAT
seqGDS2VCF(f, "tmp2.vcf.gz", info.var=c("BN","GP","AA","DP","HM2"),
  fmt.var=character())

# read
(txt <- readLines("tmp.vcf.gz", n=20))
(txt <- readLines("tmp1.vcf.gz", n=20))
(txt <- readLines("tmp2.vcf.gz", n=20))

#####
# Users could compare the new VCF file with the original VCF file
# call "diff" in Unix (a command line tool comparing files line by line)

# using all samples and variants
seqResetFilter(f)
```

```

# convert
seqGDS2VCF(f, "tmp.vcf.gz")

# file.copy(seqExampleFileName("vcf"), "old.vcf.gz", overwrite=TRUE)
# system("diff <(gunzip -c old.vcf.gz) <(gunzip -c tmp.vcf.gz)")

# 1a2,3
# > ##fileDate=20130309
# > ##source=SeqArray_RPackage_v1.0

# LOOK GOOD!

# delete temporary files
unlink(c("tmp.vcf.gz", "tmp.vcf.gz.tbi", "tmp1.vcf.gz", "tmp1.vcf.gz.tbi",
        "tmp2.vcf.gz", "tmp2.vcf.gz.tbi"), force=TRUE)

# close the GDS file
seqClose(f)

```

---

seqGet2bGeno

*Get packed genotypes*


---

## Description

Gets a RAW matrix of genotypes in a packed 2-bit format.

## Usage

```
seqGet2bGeno(gdsfile, samp_by_var=TRUE, ext_nbyte=0L, parallel=FALSE,
             verbose=FALSE)
```

## Arguments

gdsfile	a <a href="#">SeqVarGDSClass</a> object or a GDS file name
samp_by_var	if TRUE, return a sample-by-variant matrix; otherwise, return a variant-by-sample matrix
ext_nbyte	additional ext_nbyte row(s) with missing genotypes
parallel	FALSE (serial processing), TRUE (multicore processing), numeric value or other value; parallel is passed to the argument cl in <a href="#">seqParallel</a> , see <a href="#">seqParallel</a> for more details.
verbose	if TRUE, show progress information

## Details

If `samp_by_var=TRUE`, the function returns a sample-by-variant RAW matrix (`nrow = ceiling(# of samples / 4)`); otherwise, it returns a variant-by-sample RAW matrix (`nrow = ceiling(# of variants / 4)`). The RAW matrix consists of a 2-bit array, with 0, 1 and 2 for dosage, and 3 for missing genotype.

**Value**

Return a RAW matrix.

**Author(s)**

Xiuwen Zheng

**See Also**

[seqGetData](#)

**Examples**

```
# open a GDS file
f <- seqOpen(seqExampleFileName("gds"))

str(seqGet2bGeno(f))

str(seqGet2bGeno(f, samp_by_var=FALSE))

# close the GDS file
seqClose(f)
```

---

seqGetData

*Get Data*

---

**Description**

Gets data from a SeqArray GDS file.

**Usage**

```
seqGetData(gdsfile, var.name, .useraw=FALSE, .padNA=TRUE, .tolist=FALSE,
           .envir=NULL)
```

**Arguments**

<code>gdsfile</code>	a <a href="#">SeqVarGDSClass</a> object or a GDS file name
<code>var.name</code>	a variable name or a character vector, see details; if <code>character()</code> , return <code>NULL</code>
<code>.useraw</code>	<code>TRUE</code> , force to use RAW instead of <code>INTEGER</code> for genotypes and dosages; <code>FALSE</code> , use <code>INTEGER</code> ; <code>NA</code> , use RAW for small numbers instead of <code>INTEGER</code> if possible; <code>0xFF</code> is missing value if RAW is used
<code>.padNA</code>	<code>TRUE</code> , pad a variable-length vector with <code>NA</code> if the number of data points for each variant is not greater than 1
<code>.tolist</code>	if <code>TRUE</code> , return a list of vectors instead of the structure <code>list(length, data)</code> for variable-length data; <code>NA</code> , return a compressed List defined in <b>IRanges</b> when it is applicable
<code>.envir</code>	<code>NULL</code> , an environment object, a list or a <code>data.frame</code>

## Details

The variable name should be "sample.id", "variant.id", "position", "chromosome", "allele", "genotype", "annotation/id", "annotation/qual", "annotation/filter", "annotation/info/VARIABLE\_NAME" or "annotation/format/VARIABLE\_NAME".

"@genotype", "annotation/info/@VARIABLE\_NAME" or "annotation/format/@VARIABLE\_NAME" are used to obtain the index associated with these variables.

"\$chromosome" returns chromosome codes in an object of `S4Vectors::Rle`.

"\$dosage" is also allowed for the dosages of reference allele (integer: 0, 1, 2 and NA for diploid genotypes).

"\$dosage\_alt" returns a RAW/INTEGER matrix for the dosages of alternative allele without distinguishing different alternative alleles. "\$dosage\_alt2" allow the alleles are partially missing (e.g., genotypes on chromosome X for males)

"\$dosage\_sp" returns a sparse matrix (`dgCMatrix`) for the dosages of alternative allele without distinguishing different alternative alleles. "\$dosage\_sp2" allow the alleles are partially missing (e.g., genotypes on chromosome X for males)

"\$num\_allele" returns an integer vector with the numbers of distinct alleles.

"\$ref" returns a character vector of reference alleles. "\$alt" returns a character vector of alternative alleles (delimited by comma).

"\$chrom\_pos" returns characters with the combination of chromosome and position, e.g., "1:1272721".

"\$chrom\_pos2" is similar to "\$chrom\_pos", except the suffix "\_1" is added to the first duplicate following the variant, "\_2" is added to the second duplicate, and so on. "\$chrom\_pos\_allele" returns characters with the combination of chromosome, position and alleles, e.g., "1:1272721\_A\_G" (i.e., chr:position\_REF\_ALT).

"\$variant\_index" returns the indices of selected variants starting from 1, and "\$sample\_index" returns the indices of selected samples starting from 1.

"\$:VAR" return the variable "VAR" from `.envir` according to the selected variants.

## Value

Return vectors, matrices or lists (with length and data components) with a class name `SeqVarDataList`.

## Author(s)

Xiuwen Zheng

## See Also

[seqSetFilter](#), [seqApply](#), [seqNewVarData](#), [seqListVarData](#)

## Examples

```
# the GDS file
(gds.fn <- seqExampleFileName("gds"))

# display
(f <- seqOpen(gds.fn))

# get 'sample.id'
(samp.id <- seqGetData(f, "sample.id"))
# "NA06984" "NA06985" "NA06986" ...
```

```

# get 'variant.id'
head(variant.id <- seqGetData(f, "variant.id"))

# get 'chromosome'
table(seqGetData(f, "chromosome"))
seqGetData(f, "$chromosome")

# get 'allele'
head(seqGetData(f, "allele"))
# "T,C" "G,A" "G,A" ...

# get '$chrom_pos'
head(seqGetData(f, "$chrom_pos"))

# get '$dosage'
seqGetData(f, "$dosage")[1:6, 1:10]

# get a sparse matrix of dosages
seqGetData(f, "$dosage_sp")[1:6, 1:10]

# get '$num_allele'
head(seqGetData(f, "$num_allele"))

# set sample and variant filters
set.seed(100)
seqSetFilter(f, sample.id=samp.id[c(2,4,6,8,10)])
seqSetFilter(f, variant.id=sample(variant.id, 10))

# get a list
seqGetData(f, c(chr="chromosome", pos="position", allele="allele"))

# get the indices of selected variants/samples
seqGetData(f, "$variant_index")
seqGetData(f, "$sample_index")

# get genotypic data
seqGetData(f, "genotype")

# get annotation/info/DP
seqGetData(f, "annotation/info/DP")

# get annotation/info/AA, a variable-length dataset
seqGetData(f, "annotation/info/AA", .padNA=FALSE)
# $length          <- indicating the length of each variable-length data
# [1] 1 1 1 1 1 1 ...
# $data            <- the data according to $length
# [1] "T" "C" "T" "C" "G" "C" ...

# or return a simplified vector
seqGetData(f, "annotation/info/AA", .padNA=TRUE)

# return a compressed list (CharacterList)
seqGetData(f, "annotation/info/AA", .padNA=FALSE, .tolist=NA)

```

```

# get annotation/format/DP, a variable-length dataset
seqGetData(f, "annotation/format/DP")
# $length          <- indicating the length of each variable-length data
# [1] 1 1 1 1 1 1 ...
# $data           <- the data according to $length
#      variant
# sample [1] [2] [3] [4] [5] [6] ...
# [1,] 25 25 22 3 4 17 ...

# get values from R environment
env <- new.env()
env$x <- 1:1348 / 10
env$x[seqGetData(f, "$variant_index")]
seqGetData(f, "$:x", .envir=env)

# close the GDS file
seqClose(f)

```

---

seqGetFilter

*Get the Filter of GDS File*


---

## Description

Gets the filter of samples and variants.

## Usage

```
seqGetFilter(gdsfile, .useraw=FALSE)
```

## Arguments

gdsfile	a <a href="#">SeqVarGDSClass</a> object
.useraw	returns logical vectors if FALSE, and returns raw vectors if TRUE

## Value

Return a list:

sample.sel	a logical/raw vector indicating selected samples
variant.sel	a logical/raw vector indicating selected variants

## Author(s)

Xiuwen Zheng

## See Also

[seqSetFilter](#)

**Examples**

```

# the GDS file
(gds.fn <- seqExampleFileName("gds"))

# display
(f <- seqOpen(gds.fn))

# get 'sample.id'
(samp.id <- seqGetData(f, "sample.id"))
# "NA06984" "NA06985" "NA06986" ...

# get 'variant.id'
head(variant.id <- seqGetData(f, "variant.id"))

# set sample and variant filters
seqSetFilter(f, sample.id=samp.id[c(2,4,6,8,10)])
set.seed(100)
seqSetFilter(f, variant.id=sample(variant.id, 10))

# get filter
z <- seqGetFilter(f)

# the number of selected samples
sum(z$sample.sel)
# the number of selected variants
sum(z$variant.sel)

z <- seqGetFilter(f, .useraw=TRUE)
head(z$sample.sel)
head(z$variant.sel)

# close the GDS file
seqClose(f)

```

seqMerge

*Merge Multiple SeqArray GDS Files***Description**

Merges multiple SeqArray GDS files.

**Usage**

```
seqMerge(gds.fn, out.fn, storage.option="LZMA_RA", info.var=NULL, fmt.var=NULL,
         samp.var=NULL, optimize=TRUE, digest=TRUE, geno.pad=TRUE, verbose=TRUE)
```

**Arguments**

gds.fn	the file names of multiple GDS files
out.fn	the output file name

storage.option	specify the storage and compression option, "ZIP_RA" ( <a href="#">seqStorageOption("ZIP_RA")</a> ); or "LZMA_RA" to use LZMA compression algorithm with higher compression ratio (by default)
info.var	characters, the variable name(s) in the INFO field; NULL for all variables, or <code>character()</code> excludes all INFO variables
fmt.var	characters, the variable name(s) in the FORMAT field; NULL for all variables, or <code>character()</code> excludes all FORMAT variables
samp.var	characters, the variable name(s) in 'sample.annotation'; or NULL for all variables
optimize	if TRUE, optimize the access efficiency by calling <a href="#">cleanup.gds</a>
digest	a logical value (TRUE/FALSE) or a character ("md5", "sha1", "sha256", "sha384" or "sha512"); add md5 hash codes to the GDS file if TRUE or a digest algorithm is specified
geno.pad	TRUE, pad a 2-bit genotype array in bytes to avoid recompressing genotypes if possible
verbose	if TRUE, show information

### Details

The function merges multiple SeqArray GDS files. Users can specify the compression method and level for the new GDS file. If `gds.fn` contains one file, users can change the storage type to create a new file.

WARNING: the functionality of `seqMerge()` is limited.

### Value

Return the file name of GDS format with an absolute path.

### Author(s)

Xiuwen Zheng

### See Also

[seqVCF2GDS](#), [seqExport](#)

### Examples

```
# the VCF file
vcf.fn <- seqExampleFileName("vcf")

# the number of variants
total.count <- seqVCF_Header(vcf.fn, getnum=TRUE)$num.variant

split.cnt <- 5
start <- integer(split.cnt)
count <- integer(split.cnt)

s <- (total.count+1) / split.cnt
st <- 1L
for (i in 1:split.cnt)
{
  z <- round(s * i)
```

```

    start[i] <- st
    count[i] <- z - st
    st <- z
  }

  fn <- paste0("tmp", 1:split.cnt, ".gds")

  # convert to 5 gds files
  for (i in 1:split.cnt)
  {
    seqVCF2GDS(vcf.fn, fn[i], storage.option="ZIP_RA",
              start=start[i], count=count[i])
  }

  # merge different variants
  seqMerge(fn, "tmp.gds", storage.option="ZIP_RA")
  seqSummary("tmp.gds")

  ##### merging different samples #####

  vcf.fn <- seqExampleFileName("gds")
  file.copy(vcf.fn, "test.gds", overwrite=TRUE)

  # modify 'sample.id'
  seqAddValue("test.gds", "sample.id", paste0("S", 1:90), replace=TRUE)

  # merging
  seqMerge(c(vcf.fn, "test.gds"), "output.gds", storage.option="ZIP_RA")

  # delete the temporary files
  unlink(c("tmp.gds", "test.gds", "output.gds"), force=TRUE)
  unlink(fn, force=TRUE)

```

---

seqMissing

*Missing genotype percentage*


---

### Description

Calculates the missing rates per variant or per sample.

### Usage

```
seqMissing(gdsfile, per.variant=TRUE, parallel=seqGetParallel(), balancing=NA,
           verbose=FALSE)
```

### Arguments

gdsfile	a <a href="#">SeqVarGDSClass</a> object
per.variant	missing rate per variant if TRUE, missing rate per sample if FALSE, or calculating missing rates for variants and samples if NA

parallel	FALSE (serial processing), TRUE (multicore processing), numeric value or other value; parallel is passed to the argument <code>cl</code> in <a href="#">seqParallel</a> , see <a href="#">seqParallel</a> for more details.
balancing	whether to perform workload balancing or not, only applicable when multiple cores are used; if NA, use TRUE as a default until <code>getOption("seqarray.balancing")</code> is set and not TRUE
verbose	if TRUE, show progress information

### Details

If the `gds` node 'genotype/data' (integer genotypes) is not available, the node 'annotation/format/DS' (numeric genotype dosages for alternative alleles) will be used to calculate allele frequencies. At a site, it assumes 'annotation/format/DS' stores the dosage of the 1st alternative allele in the 1st column, 2nd alt. allele in the 2nd column if it is multi-allelic, and so on.

### Value

A vector of missing rates, or a `list(variant, sample)` for both variants and samples.

### Author(s)

Xiuwen Zheng

### See Also

[seqAlleleFreq](#), [seqNumAllele](#), [seqParallel](#), [seqGetParallel](#)

### Examples

```
# the GDS file
(gds.fn <- seqExampleFileName("gds"))

# display
(f <- seqOpen(gds.fn))

summary(m1 <- seqMissing(f, TRUE, verbose=TRUE))
summary(m2 <- seqMissing(f, FALSE, verbose=TRUE))

str(m <- seqMissing(f, NA, verbose=TRUE))
identical(m1, m$variant) # should be TRUE
identical(m2, m$sample) # should be TRUE

# close the GDS file
seqClose(f)
```

---

seqNewVarData

*Variable-length data*

---

### Description

Gets a variable-length data object.

**Usage**

```
seqNewVarData(len, data)
seqListVarData(obj, useList=FALSE)
```

**Arguments**

len                    a non-negative vector for variable lengths  
 data                   a vector of data according to len  
 obj                    a SeqVarDataList object or a compressed list (defined in **IRanges**)  
 useList                if TRUE, return a compressed List defined in **IRanges** (e.g., [seqGetData](#))

**Details**

seqNewVarData() creates a SeqVarDataList object for variable-length data, and seqListVarData() converts the SeqVarDataList object to a list. seqGetData() returns a SeqVarDataList object for variable-length data; seqAddValue() can add a SeqVarDataList object to a GDS file.

**Value**

Return a SeqVarDataList object or a CompressedAtomicList object (e.g., IntegerList()).

**Author(s)**

Xiuwen Zheng

**See Also**

[seqGetData](#), [seqAddValue](#), [IntegerList](#)

**Examples**

```
obj <- seqNewVarData(c(1,2,1,0,2), c("A", "B", "B", "C", "E", "E"))
obj

seqListVarData(obj) # a list
(a <- seqListVarData(obj, useList=TRUE)) # CharacterList
seqListVarData(a) # a list
```

---

seqNumAllele	<i>Number of alleles</i>
--------------	--------------------------

---

**Description**

Returns the numbers of alleles for each site.

**Usage**

```
seqNumAllele(gdsfile)
```

**Arguments**

gdsfile                a [SeqVarGDSClass](#) object

**Value**

The numbers of alleles for each site.

**Author(s)**

Xiuwen Zheng

**See Also**

[seqAlleleFreq](#), [seqMissing](#)

**Examples**

```
# the GDS file
(gds.fn <- seqExampleFileName("gds"))

# display
f <- seqOpen(gds.fn)

table(seqNumAllele(f))

# close the GDS file
seqClose(f)
```

---

seqOpen

*Open a SeqArray GDS File*

---

**Description**

Opens a SeqArray GDS file.

**Usage**

```
seqOpen(gds.fn, readonly=TRUE, allow.duplicate=FALSE)
```

**Arguments**

gds.fn	the file name
readonly	whether read-only or not
allow.duplicate	if TRUE, it is allowed to open a GDS file with read-only mode when it has been opened in the same R session

**Details**

It is strongly suggested to call seqOpen instead of [openfn.gds](#), since seqOpen will perform internal checking for data integrity.

**Value**

Return an object of class SeqVarGDSClass inherited from [gds.class](#).

**Author(s)**

Xiuwen Zheng

**See Also**[seqClose](#), [seqGetData](#), [seqApply](#)**Examples**

```
# the GDS file
(gds.fn <- seqExampleFileName("gds"))

# open the GDS file
gdsfile <- seqOpen(gds.fn)

# display the contents of the GDS file in a hierarchical structure
gdsfile

# close the GDS file
seqClose(gdsfile)
```

seqOptimize

*Optimize the Storage of Data Array***Description**

Transpose data array or matrix for possibly higher-speed access.

**Usage**

```
seqOptimize(gdsfile, target=c("chromosome", "by.sample"), format.var=TRUE,
            cleanup=TRUE, verbose=TRUE)
```

**Arguments**

gdsfile	a <a href="#">SeqVarGDSClass</a> object or a GDS file name
target	"chromosome", "by.sample"; see details
format.var	a character vector for selected variable names, or TRUE for all variables, according to "annotation/format"
cleanup	call <code>link{cleanup.gds}</code> if TRUE
verbose	if TRUE, show information

**Details**

"chromosome": adding or updating two additional nodes '@chrom\_rle\_val' and '@chrom\_rle\_len' for faster chromosome indexing, requiring `SeqArray>=v1.20.0`.

"by.sample": optimizing GDS file for `seqApply(..., margin="by.sample")`. Warning: optimizing GDS file for reading data by sample may increase file size by up to 2X as genotype data and all format data are duplicated.

**Value**

None.

**Author(s)**

Xiuwen Zheng

**See Also**

[seqGetData](#), [seqApply](#)

**Examples**

```
# the file name of VCF
(vcf.fn <- seqExampleFileName("vcf"))
# or vcf.fn <- "C:/YourFolder/Your_VCF_File.vcf"

# convert
seqVCF2GDS(vcf.fn, "tmp.gds", storage.option="ZIP_RA")

# prepare data
seqOptimize("tmp.gds", target="by.sample")

# list the structure of GDS variables
(f <- seqOpen("tmp.gds", readonly=FALSE))
seqOptimize(f, "chromosome")

# close
seqClose(f)

# delete the temporary file
unlink("tmp.gds")
```

---

seqParallel

*Apply Functions in Parallel*

---

**Description**

Applies a user-defined function in parallel.

**Usage**

```
seqParallel(cl=seqGetParallel(), gdsfile, FUN,
  split=c("by.variant", "by.sample", "none"), .combine="unlist",
  .selection.flag=FALSE, .initialize=NULL, .finalize=NULL, .initparam=NULL,
  .balancing=FALSE, .bl_size=NA_integer_, .bl_progress=FALSE,
  .status_file=FALSE, .proc_time=FALSE, ...)
seqParApply(cl=seqGetParallel(), x, FUN, .balancing=TRUE, ...)
```

**Arguments**

<code>cl</code>	NULL or FALSE: serial processing; TRUE: multicore processing (the maximum number of cores minor one); a numeric value: the number of cores to be used; a cluster object for parallel processing, created by the functions in the package <a href="#">parallel</a> , like <a href="#">makeCluster</a> ; a <code>BiocParallelParam</code> object from the <code>BiocParallel</code> package. See details
<code>gdsfile</code>	a <a href="#">SeqVarGDSClass</a> object, or NULL
<code>FUN</code>	the function to be applied, should be like <code>FUN(gdsfile, ...)</code> if <code>gdsfile</code> is given, or <code>FUN(...)</code> if <code>gdsfile=NULL</code>
<code>split</code>	split the dataset by variant or sample according to multiple processes, or "none" for no split; <code>split="by.variant"</code> by default
<code>.combine</code>	define a function for combining results from different processes; by default, "unlist" is used, to produce a vector which contains all the atomic components, via <code>unlist(..., recursive=FALSE)</code> ; "list", return a list of results created by child processes; "none", no return; or a function with one or two arguments, like "+"
<code>.selection.flag</code>	TRUE – passes a logical vector of selection to the second argument of <code>FUN(gdsfile, selection, ...)</code>
<code>.initialize</code>	a user-defined function for initializing workers, should have two arguments ( <code>process_id, param</code> )
<code>.finalize</code>	a user-defined function for finalizing workers, should have two arguments ( <code>process_id, param</code> )
<code>.initparam</code>	parameters passed to <code>.initialize</code> and <code>.initialize</code>
<code>.balancing</code>	whether to perform workload balancing or not, only applicable when multiple cores are used; if NA, use TRUE as a default until <code>getOption("seqarray.balancing")</code> is set and not TRUE
<code>.bl_size</code>	chunk size (# of variants or samples), the increment for load balancing, only applicable if <code>.balancing=TRUE</code> ; if NA_integer_, automatically determined by the number of cores
<code>.bl_progress</code>	if TRUE and <code>.balancing=TRUE</code> , show progress information
<code>.status_file</code>	if TRUE, create empty files for saving the status of child processes, which can be used in the user-defined function
<code>.proc_time</code>	if TRUE, show the elapsed time calling the function
<code>x</code>	a vector (atomic or list), passed to <code>FUN</code>
<code>...</code>	optional arguments to <code>FUN</code>

**Details**

When `cl` is TRUE or a numeric value, forking techniques are used to create a new child process as a copy of the current R process, see `?parallel:mcfork`. However, forking is not available on Windows, and [makeCluster](#) is called to make a cluster which will be deallocated after calling `FUN`.

It is strongly suggested to use `seqParallel` together with `seqParallelSetup`. `seqParallelSetup` could work around the problem of forking on Windows, without allocating clusters frequently.

The user-defined function could use two predefined variables `SeqArray:::process_count` and `SeqArray:::process_index` to tell the total number of cluster nodes and which cluster node being used.

`seqParallel(, gdsfile=NULL, FUN=..., split="none")` could be used to setup multiple streams of pseudo-random numbers, and see [nextRNGStream](#) or [nextRNGSubStream](#) in the package `parallel`.

**Value**

A vector or list of values.

**Author(s)**

Xiuwen Zheng

**See Also**

[seqSetFilter](#), [seqGetData](#), [seqApply](#), [seqParallelSetup](#), [seqGetParallel](#)

**Examples**

```
library(parallel)

# choose an appropriate cluster size or number of cores
seqParallelSetup(2)

# the GDS file
(gds.fn <- seqExampleFileName("gds"))

# display
(gdsfile <- seqOpen(gds.fn))

# the uniprocessor version
afreq1 <- seqParallel(, gdsfile, FUN = function(f) {
  seqApply(f, "genotype", as.is="double",
    FUN=function(x) mean(x==0, na.rm=TRUE))
}, split="by.variant")

length(afreq1)
summary(afreq1)

# run in parallel
afreq2 <- seqParallel(, gdsfile, FUN = function(f) {
  seqApply(f, "genotype", as.is="double",
    FUN=function(x) mean(x==0, na.rm=TRUE))
}, split="by.variant")

length(afreq2)
summary(afreq2)

# check
length(afreq1) # 1348
all(afreq1 == afreq2)

#####
# check -- variant splits

seqParallel(, gdsfile, FUN = function(f) {
  v <- seqGetFilter(f)
  sum(v$variant.sel)
}, split="by.variant")
```

```

# [1] 674 674

#####

seqParallel(, NULL, FUN = function() {
  paste(SeqArray:::process_index, SeqArray:::process_count, sep=" / ")
}, split="none")

seqParallel(, NULL, FUN = function() {
  SeqArray:::process_index
}, split="none", .combine=function(i) print(i))

seqParallel(, NULL, FUN = function() {
  SeqArray:::process_index
}, split="none", .combine="+")

#####

# close the GDS file
seqClose(gdsfile)

# clear the parallel cluster
seqParallelSetup(FALSE)

```

---

seqParallelSetup

*Setup/Get a Parallel Environment*


---

## Description

Sets up a parallel environment in R for the current session.

## Usage

```

seqParallelSetup(cluster=TRUE, verbose=TRUE)
seqGetParallel()
seqMulticoreSetup(num, type=c("psock", "fork"), verbose=TRUE)

```

## Arguments

cluster	NULL or FALSE: serial processing; TRUE: parallel processing with the maximum number of cores minor one; a numeric value: the number of cores to be used; a cluster object for parallel processing, created by the functions in the package <a href="#">parallel</a> , like <a href="#">makeCluster</a> . See details
num	the maximum number of cores used for the user-defined multicore setting; FALSE, NA or any value less than 2, to disable the pre-defined multicore cluster; see details
type	either PSOCK or Fork cluster setup for the multicore setting, the resulting parallel cluster will be used if 'parallel' is a number greater than one in associated functions
verbose	if TRUE, show information

## Details

When `cl` is `TRUE` or a numeric value, forking techniques are used to create a new child process as a copy of the current R process, see `?parallel::mcfork`. However, forking is not available on Windows, so multiple processes created by `makeCluster` are used instead. The R environment option `seqarray.parallel` will be set according to the value of `cluster`. Using `seqParallelSetup(FALSE)` removes the registered cluster, as does stopping the registered cluster.

`seqMulticoreSetup` sets a multicore cluster and can avoid large memory usage when forking is used for the number of cores specified in the argument `'parallel'`. It is recommended to call `seqMulticoreSetup` before running any memory-intensive functions, as forking within such functions can significantly increase memory usage.

## Value

`seqParallelSetup()` has no return, and `seqGetParallel()` returns `getOption("seqarray.parallel", FALSE)`.

## Author(s)

Xiuwen Zheng

## See Also

[seqParallel](#), [seqApply](#)

## Examples

```
library(parallel)

seqParallelSetup(2L)

# the GDS file
(gds.fn <- seqExampleFileName("gds"))

# display
(f <- seqOpen(gds.fn))

# run in parallel
summary(seqMissing(f))

# close the GDS file
seqClose(f)

seqParallelSetup(FALSE)
```

---

seqRecompress

*Recompress the GDS file*

---

## Description

Recompress the SeqArray GDS file.

**Usage**

```
seqRecompress(gds.fn,
              compress=c("ZIP", "LZ4", "LZMA", "Ultra", "UltraMax", "none"),
              exclude=character(), optimize=TRUE, digest=TRUE, verbose=TRUE)
```

**Arguments**

<code>gds.fn</code>	the file name of SeqArray file
<code>compress</code>	the compression method, <code>compress="ZIP"</code> by default
<code>exclude</code>	a list of GDS nodes to be excluded, see details
<code>optimize</code>	if TRUE, optimize the access efficiency by calling <a href="#">cleanup.gds</a>
<code>digest</code>	a logical value (TRUE/FALSE), whether adding md5 hash codes to the GDS file
<code>verbose</code>	if TRUE, show information

**Details**

This function requires `gdsfmt` ( $\geq$  v1.17.2). [seqVCF2GDS](#) usually takes lots of memory when the compression method `"LZMA_RA.max"`, `"Ultra"` or `"UltraMax"` is specified. So users could call `seqVCF2GDS(, storage.option="ZIP_RA")` first, and then recompress the GDS file with a higher compression option, e.g., `"UltraMax"`. `seqRecompress()` takes much less memory than `seqVCF2GDS()`, since it recompresses data in a GDS node each time.

`"UltraMax"` might be not better than `"Ultra"`, and its behavior is similar to `xz -9 --extreme`: use a slower variant of the selected compression preset level (-9) to hopefully get a little bit better compression ratio, but with bad luck this can also make it worse.

`ls.gdsn(gdsfile, include.hidden=TRUE, recursive=TRUE)` returns a list of GDS nodes to be re-compressed, and users can specify the excluded nodes in the argument `exclude`.

**Value**

None.

**Author(s)**

Xiuwen Zheng

**See Also**

[seqVCF2GDS](#), [seqStorageOption](#)

**Examples**

```
gds.fn <- seqExampleFileName("gds")
file.copy(gds.fn, "tmp.gds")

seqRecompress("tmp.gds", "LZMA")

unlink("tmp.gds")
```

---

seqResetVariantID      *Reset Variant ID in SeqArray GDS Files*

---

### Description

Resets the variant IDs in multiple SeqArray GDS files.

### Usage

```
seqResetVariantID(gds.fn, start=1L, set=NULL, digest=TRUE, optimize=TRUE,
  verbose=TRUE)
```

### Arguments

gds.fn	a character vector of multiple GDS file names
start	the starting number of the sequence of variant IDs
set	NULL or a logical vector; NULL for resetting all files, or TRUE for resetting variant.id for that GDS file
digest	a logical value, if TRUE, add a md5 hash code
optimize	if TRUE, optimize the access efficiency by calling <a href="#">cleanup.gds</a>
verbose	if TRUE, show information

### Details

The variant IDs will be replaced by the numbers in sequential order and adjacent to each file. The variant ID starts from start (1 by default) in the first GDS file.

### Value

None.

### Author(s)

Xiuwen Zheng

### See Also

[seqVCF2GDS](#)

### Examples

```
fn <- seqExampleFileName("gds")

file.copy(fn, "tmp1.gds", overwrite=TRUE)
file.copy(fn, "tmp2.gds", overwrite=TRUE)

gds.fn <- c("tmp1.gds", "tmp2.gds")
seqResetVariantID(gds.fn)

f <- seqOpen("tmp1.gds")
head(seqGetData(f, "variant.id"))
```

```
seqClose(f)

f <- seqOpen("tmp2.gds")
head(seqGetData(f, "variant.id"))
seqClose(f)

# delete the temporary files
unlink(gds.fn, force=TRUE)
```

---

seqSetFilter-methods *Set a Filter to Sample or Variant*

---

## Description

Sets a filter to sample and/or variant.

## Usage

```
## S4 method for signature 'SeqVarGDSCClass,ANY'
seqSetFilter(object, variant.sel,
             sample.sel=NULL, variant.id=NULL, sample.id=NULL,
             action=c("set", "intersect", "push", "push+set", "push+intersect", "pop"),
             ret.idx=FALSE, warn=TRUE, verbose=TRUE)
## S4 method for signature 'SeqVarGDSCClass,GRanges'
seqSetFilter(object, variant.sel,
             rm.txt="chr", intersect=FALSE, verbose=TRUE)
## S4 method for signature 'SeqVarGDSCClass,GRangesList'
seqSetFilter(object, variant.sel,
             rm.txt="chr", intersect=FALSE, verbose=TRUE)
## S4 method for signature 'SeqVarGDSCClass,IRanges'
seqSetFilter(object, variant.sel,
             chr, intersect=FALSE, verbose=TRUE)
seqResetFilter(object, sample=TRUE, variant=TRUE, verbose=TRUE)
seqSetFilterChrom(object, include=NULL, is.num=NA, from.bp=NULL, to.bp=NULL,
                 intersect=FALSE, verbose=TRUE)
seqSetFilterPos(object, chr, pos, ref=NULL, alt=NULL, intersect=FALSE,
                multi.pos=TRUE, ret.idx=FALSE, verbose=TRUE)
seqSetFilterAnnotID(object, id, ret.idx=FALSE, verbose=TRUE)
seqFilterPush(object) # store the current filter
seqFilterPop(object) # restore the last filter
```

## Arguments

object	a <a href="#">SeqVarGDSCClass</a> object
variant.sel	a logical/raw/index vector indicating the selected variants; <a href="#">GRanges</a> , a <a href="#">GRanges</a> object for the genomic locations; <a href="#">GRangesList</a> , a <a href="#">GRangesList</a> object for storing a collection of <a href="#">GRanges</a> objects; <a href="#">IRanges</a> , a <a href="#">IRanges</a> object for storing a collection of range objects
sample.sel	a logical/raw/index vector indicating the selected samples
variant.id	ID of selected variants

sample.id	ID of selected samples
action	"set" – set the current filter via sample.id, variant.id, samp.sel or variant.sel; "intersect" – set the current filter to the intersection of selected samples and/or variants; "push" – push the current filter to the stack, and it could be recovered by "pop" later, no change on the current filter; "push+set" – push the current filter to the stack, and changes the current filter via sample.id, variant.id, samp.sel or variant.sel; "push+intersect" – push the current filter to the stack, and set the current filter to the intersection of selected samples and/or variants; "pop" – pop up the last filter
ret.idx	if TRUE, return the index in the output array according to the order of 'sample.id', 'sample.sel', 'variant.id' or 'variant.sel'
rm.txt	a character, the characters will be removed from seqnames(variant.sel)
chr	a vector of character for chromosome coding
pos	a vector of numeric values for genome coordinate
sample	logical, if TRUE, include all samples
variant	logical, if TRUE, include all variants
include	NULL, or a vector of characters for specified chromosome(s)
is.num	a logical variable: TRUE, chromosome code is numeric; FALSE, chromosome is not numeric; is.num=TRUE is usually used to exclude non-autosomes
from.bp	NULL, no limit; a numeric vector, the lower bound of position
to.bp	NULL, no limit; a numeric vector, the upper bound of position
intersect	if FALSE, the candidate samples/variants for selection are all samples/variants (by default); if TRUE, the candidate samples/variants are from the selected samples/variants defined via the previous call
ref	a character vector of reference alleles for additional matching, or NULL (no reference allele filter); the length should be the same as pos, and NA for entries that match any reference allele. Must be used together with alt
alt	a character vector of alternative alleles for additional matching, or NULL (no alternative allele filter); the length should be the same as pos, and NA for entries that match any alternative allele. Must be used together with ref
multi.pos	FALSE, use the first matched position; TRUE, allow multiple variants at the same position
id	a character vector for RS IDs (stored in "annotation/id")
warn	if TRUE, show a warning when the input sample.sel or variant.sel is not ordered as the GDS file or there is any duplicate
verbose	if TRUE, show information

## Details

seqResetFilter(file) is equivalent to seqSetFilter(file), where the selection arguments in seqSetFilter are NULL.

If from.bp and to.bp has values, they should be equal-size as include. A trio of include, from.bp and to.bp indicates a region on human genomes. NA in from.bp is treated as 0, and NA in to.bp is treated as the maximum of integer ( $2^{31} - 1$ ).

**Value**

If `ret.idx=TRUE`, `seqSetFilter()` returns a list with two components `sample_idx` and `variant_idx` to indicate the indices of the output array according to the input `'sample.id'`, `'sample.sel'`, `'variant.id'` or `'variant.sel'`; if `ret.idx=TRUE`, `seqSetFilterAnnotID()` return an index vector; otherwise no return.

**Author(s)**

Xiuwen Zheng

**See Also**

[seqSetFilterCond](#), [seqGetFilter](#), [seqGetData](#), [seqApply](#)

**Examples**

```
# the GDS file
(gds.fn <- seqExampleFileName("gds"))

# display
(f <- seqOpen(gds.fn))

# get 'sample.id'
(samp.id <- seqGetData(f, "sample.id"))
# "NA06984" "NA06985" "NA06986" ...

# get 'variant.id'
head(variant.id <- seqGetData(f, "variant.id"))

# get 'chromosome'
table(seqGetData(f, "chromosome"))

# get 'allele'
head(seqGetData(f, "allele"))
# "T,C" "G,A" "G,A" ...

# set sample filters
seqSetFilter(f, sample.id=samp.id[c(2,4,6,8)])
seqSetFilter(f, sample.id=samp.id[c(2,4,6,8)], ret.idx=TRUE)

(v <- seqSetFilter(f, sample.id=samp.id[c(8,2,6,4)], ret.idx=TRUE))
all(seqGetData(f, "sample.id")[v$sample_idx] == samp.id[c(8,2,6,4)])

# set variant filters
seqSetFilter(f, variant.id=variant.id[c(2,4,6,8,10,12)], ret.idx=TRUE)
(v <- seqSetFilter(f, variant.id=variant.id[c(12,4,6,10,8,12)], ret.idx=TRUE))
all(variant.id[c(12,4,6,10,8,12)] == seqGetData(f, "variant.id")[v$variant_idx])

set.seed(100)
seqSetFilter(f, variant.id=sample(variant.id, 5))

# get genotypic data
seqGetData(f, "genotype")
```

```

## OR
# set sample and variant filters
seqSetFilter(f, sample.sel=c(2,4,6,8))
set.seed(100)
seqSetFilter(f, variant.sel=sample.int(length(variant.id), 5))

# get genotypic data
seqGetData(f, "genotype")

## set the intersection

seqResetFilter(f)
seqSetFilterChrom(f, 10L)
seqSummary(f, "genotype", check="none")

AF <- seqAlleleFreq(f)
table(AF <= 0.9)

seqSetFilter(f, variant.sel=(AF<=0.9), action="intersect")
seqSummary(f, "genotype", check="none")

## chromosome

seqResetFilter(f)

seqSetFilterChrom(f, is.num=TRUE)
seqSummary(f, "genotype", check="none")

seqSetFilterChrom(f, is.num=FALSE)
seqSummary(f, "genotype", check="none")

seqSetFilterChrom(f, 1:4)
seqSummary(f, "genotype", check="none")
table(seqGetData(f, "chromosome"))

# HLA region
seqSetFilterChrom(f, 6, from.bp=29719561, to.bp=32883508)
seqSummary(f, "genotype", check="none")

# two regions
seqSetFilterChrom(f, c(1, 6), from.bp=c(1000000, 29719561),
  to.bp=c(90000000, 32883508))
seqSummary(f, "genotype", check="none")
seqGetData(f, "chromosome")

## intersection option

seqResetFilter(f)
seqSetFilterChrom(f, 6, from.bp=29719561, to.bp=32883508) # MHC
seqSetFilterChrom(f, include=6) # chromosome 6

seqResetFilter(f)

```

```

seqSetFilterChrom(f, 6, from.bp=29719561, to.bp=32883508) # MHC
seqSetFilterChrom(f, include=6, intersect=TRUE) # MHC region only

## set filter by position
seqSetFilterPos(f, chr=c(5,14,19,19,1,20),
  pos=c(7945937,93982552,57310283,42747452,158593598,52208935))
as.data.frame(seqGetData(f, c("chromosome", "position", "allele")))

seqSetFilterPos(f, chr=c(5,14,19,19,1,20),
  pos=c(7945937,93982552,57310283,42747452,158593598,52208935),
  ref=c("C","G","G","G","T","C"), alt=c("T","A","A","A","C","T"),
  ret.idx=TRUE)
# NA 1 NA NA NA 2 (NA for ref/alt not matched)
as.data.frame(seqGetData(f, c("chromosome", "position", "allele")))

# close the GDS file
seqClose(f)

```

---

seqSetFilterCond      *Set a Filter to Variant with Allele Count/Freq*

---

## Description

Sets a filter to variant with specified allele count/frequency and missing rate.

## Usage

```

seqSetFilterCond(gdsfile, maf=NaN, mac=1L, missing.rate=NaN,
  parallel=seqGetParallel(), balancing=NA, .progress=FALSE, verbose=TRUE)

```

## Arguments

gdsfile	a <a href="#">SeqVarGDSCClass</a> object
maf	minimum minor reference allele frequency, or a range of MAF maf[1] <= ... < maf[2]
mac	minimum minor reference allele count, or a range of MAC mac[1] <= ... < mac[2]
missing.rate	maximum missing genotype rate
parallel	FALSE (serial processing), TRUE (multicore processing), numeric value or other value; parallel is passed to the argument cl in <a href="#">seqParallel</a> , see <a href="#">seqParallel</a> for more details.
balancing	whether to perform workload balancing or not, only applicable when multiple cores are used; if NA, use TRUE as a default until <code>getOption("seqarray.balancing")</code> is set and not TRUE
.progress	if TRUE, show progress information
verbose	if TRUE, show the number of selected variants

## Value

None.

**Author(s)**

Xiuwen Zheng

**See Also**[seqSetFilter](#), [seqSetFilterChrom](#), [seqAlleleFreq](#), [seqAlleleCount](#), [seqMissing](#)**Examples**

```
# the GDS file
(gds.fn <- seqExampleFileName("gds"))

# display
(f <- seqOpen(gds.fn))

seqSetFilterChrom(f, c(1, 6))
seqSetFilterCond(f, maf=0.05, .progress=TRUE)

seqSetFilterChrom(f, c(1, 6))
seqSetFilterCond(f, maf=c(0.01, 0.05), .progress=TRUE)

# close the GDS file
seqClose(f)
```

seqSNP2GDS

*Convert SNPRelate Format to SeqArray Format***Description**

Converts a SNP GDS file to a SeqArray GDS file.

**Usage**

```
seqSNP2GDS(gds.fn, out.fn, storage.option="LZMA_RA", major.ref=TRUE,
           ds.type=c("packedreal16", "float", "double"), optimize=TRUE, digest=TRUE,
           verbose=TRUE)
```

**Arguments**

<code>gds.fn</code>	the file name of SNP format
<code>out.fn</code>	the file name, output a file of SeqArray format
<code>storage.option</code>	specify the storage and compression options, "LZMA_RA" to use LZMA compression algorithm with higher compression ratio compared to "ZIP_RA"
<code>major.ref</code>	if TRUE, use the major allele as a reference allele; otherwise, use A allele in SNP GDS file as a reference allele
<code>ds.type</code>	applicable when import dosages, the data type for storing dosages; see <a href="#">add.gdsn</a> ; <code>ds.type="packedreal16"</code> by default
<code>optimize</code>	if TRUE, optimize the access efficiency by calling <a href="#">cleanup.gds</a>

digest	a logical value (TRUE/FALSE) or a character ("md5", "sha1", "sha256", "sha384" or "sha512"); add hash codes to the GDS file if TRUE or a digest algorithm is specified
verbose	if TRUE, show information

**Value**

Return the file name of SeqArray file with an absolute path. If the input file is genotype dosage, the dosage matrix is stored in the node annotation/format/DS with the estimated dosage of alternative alleles. Any value less than 0 or greater than 2 will be replaced by NaN.

**Author(s)**

Xiuwen Zheng

**See Also**

[seqGDS2SNP](#), [seqVCF2GDS](#), [seqGDS2VCF](#), [seqBED2GDS](#)

**Examples**

```
library(SNPRelate)

# the GDS file
gds.fn <- snpgdsExampleFileName()

seqSNP2GDS(gds.fn, "tmp.gds")

seqSummary("tmp.gds")

# remove the temporary file
unlink("tmp.gds", force=TRUE)
```

---

seqStorageOption

*Storage and Compression Options*


---

**Description**

Storage and compression options for GDS import and merging.

**Usage**

```
seqStorageOption(compression=c("ZIP_RA", "ZIP_RA.fast", "ZIP_RA.max", "LZ4_RA",
  "LZ4_RA.fast", "LZ4_RA.max", "LZMA_RA", "LZMA_RA.fast", "LZMA_RA.max",
  "Ultra", "UltraMax", "none"), mode=NULL, float.mode="float32",
  geno.compress=NULL, info.compress=NULL, format.compress=NULL,
  index.compress=NULL, ...)
```

**Arguments**

compression	the default compression level ("ZIP_RA"), see <a href="#">add.gdsn</a> for the description of compression methods
mode	a character vector, specifying storage type for corresponding variable, e.g., c('annotation/info/HM'='int', 'annotation/format/PL'='int')
float.mode	specify the storage mode for read numbers, e.g., "float32", "float64", "packedreal16"; the additional parameters can follow by colon, like "packedreal16:scale=0.0001"
geno.compress	NULL for the default value, or the compression method for genotypic data
info.compress	NULL for the default value, or the compression method for data sets stored in the INFO field (i.e., "annotation/info")
format.compress	NULL for the default value, or the compression method for data sets stored in the FORMAT field (i.e., "annotation/format")
index.compress	NULL for the default value, or the compression method for data index variables (e.g., "annotation/info/@HM")
...	other specified storage compression for corresponding variable, e.g., 'annotation/info/HM'="ZIP_MAX"

**Details**

The compression modes "Ultra" and "UltraMax" attempt to maximize the compression ratio using gigabyte-sized or even terabyte-sized virtual memory, according to "LZMA\_RA.ultra" and "LZMA\_RA.ultra\_max" in [compression.gdsn](#). These features require gdsfmt ( $\geq v1.16.0$ ). "Ultra" and "UltraMax" may not increase the compression ratio much compared with "LZMA\_RA.max", and these options are designed for the users who want to exhaust the computational resources.

**Value**

Return a list with a class name "SeqGDSStorageClass", contains the compression algorithm for each data type.

**Author(s)**

Xiuwen Zheng

**See Also**

[seqVCF2GDS](#), [seqRecompress](#), [seqMerge](#)

**Examples**

```
# the file of VCF
(vcf.fn <- seqExampleFileName("vcf"))

# convert
seqVCF2GDS(vcf.fn, "tmp1.gds", storage.option=seqStorageOption())
(f1 <- seqOpen("tmp1.gds"))

# convert (maximize the compression ratio)
seqVCF2GDS(vcf.fn, "tmp2.gds", storage.option=seqStorageOption("ZIP_RA.max"))
(f2 <- seqOpen("tmp2.gds"))
```

```

# does not compress the genotypic data
seqVCF2GDS(vcf.fn, "tmp3.gds", storage.option=
  seqStorageOption("ZIP_RA", geno.compress=""))
(f3 <- seqOpen("tmp3.gds"))

# compress with LZ4
seqVCF2GDS(vcf.fn, "tmp4.gds", storage.option=seqStorageOption("LZ4_RA"))
(f4 <- seqOpen("tmp4.gds"))

# close and remove the files
seqClose(f1)
seqClose(f2)
seqClose(f3)
seqClose(f4)

unlink(c("tmp1.gds", "tmp2.gds", "tmp3.gds", "tmp4.gds"))

```

seqSummary

*Summarize a SeqArray GDS File***Description**

Gets the summary of SeqArray GDS file.

**Usage**

```
seqSummary(gdsfile, varname=NULL, check=c("default", "none", "full"),
  verbose=TRUE)
```

**Arguments**

gdsfile	a <a href="#">SeqVarGDSClass</a> object, or a file name
varname	if NULL, check the whole GDS file; or a character specifying variable name, and return a description of that variable. See details
check	should be one of "default", "none", "full"; check="default" by default
verbose	if TRUE, display information

**Details**

If check="default", the function performs regular checking, like variable dimensions. If check="full", it performs more checking, e.g., unique sample id, unique variant id, whether genotypic data are in a valid range or not.

**Value**

If varname=NULL, the function returns a list:

filename	the file name
version	the version of SeqArray format
reference	genome reference, a character vector (0-length for undefined)

ploidy	the number of sets of chromosomes
num.sample	the total number of samples
num.variant	the total number of variants
allele	allele information, see <code>seqSummary(gdsfile, "allele")</code>
annot_qual	the total number of "annotation/qual" if <code>check="none"</code> , or a summary object including min, max, median, mean
filter	filter information, see <code>seqSummary(gdsfile, "annotation/filter")</code>
info	a <code>data.frame</code> of INFO field: ID, Number, Type, Description, Source and Version
format	a <code>data.frame</code> of FORMAT field: ID, Number, Type and Description
sample.annot	a <code>data.frame</code> of sample annotation with ID, Type and Description

— `seqSummary(gdsfile, "genotype", check="none", verbose=FALSE)` returns a list with components:

dim	an integer vector: ploidy, # of samples, # of variants
seldim	an integer vector: ploidy, # of selected samples, # of selected variants

— `seqSummary(gdsfile, "allele")` returns a `data.frame` with ID and descriptions (`check="none"`), or a list with components:

value	a <code>data.frame</code> with ID and Description
table	cross tabulation for the number of alleles per site

— `seqSummary(gdsfile, "$alt")` returns a `data.frame` with ID and Description for describing the alternative alleles.

— `seqSummary(gdsfile, "annotation/filter")` or `seqSummary(gdsfile, "$filter")` returns a `data.frame` with ID and description (`check="none"`), or a list with components: value (a `data.frame` with ID and Description), table (cross tabulation for the variable 'filter').

— `seqSummary(gdsfile, "annotation/info")` or `seqSummary(gdsfile, "$info")` returns a `data.frame` describing the variables in the folder "annotation/info" with ID, Number, Type, Description, Source and Version.

— `seqSummary(gdsfile, "annotation/format")` returns a `data.frame` describing the variables in the folder "annotation/format" with ID, Number, Type and Description.

— `seqSummary(gdsfile, "sample.annotation")` returns a `data.frame` describing sample annotation with ID, Type and Description.

— `seqSummary(gdsfile, "$reference")` returns the genome reference if it is defined (a 0-length character vector if undefined).

— `seqSummary(gdsfile, "$contig")` returns the contig information, a `data.frame` including ID.

— `seqSummary(gdsfile, "$format")` returns a `data.frame` describing VCF FORMAT header with ID, Number, Type and Description. The first row is used for genotypes.

— `seqSummary(gdsfile, "$digest")` returns a `data.frame` with the full names of GDS variables, digest codes and validation (FALSE/TRUE).

### Author(s)

Xiuwen Zheng

**See Also**

[seqGetData](#), [seqApply](#)

**Examples**

```
# the GDS file
(gds.fn <- seqExampleFileName("gds"))

seqSummary(gds.fn)

ans <- seqSummary(gds.fn, check="full")
ans

seqSummary(gds.fn, "genotype")
seqSummary(gds.fn, "allele")
seqSummary(gds.fn, "annotation/filter")
seqSummary(gds.fn, "annotation/info")
seqSummary(gds.fn, "annotation/format")
seqSummary(gds.fn, "sample.annotation")

seqSummary(gds.fn, "$reference")
seqSummary(gds.fn, "$contig")
seqSummary(gds.fn, "$filter")
seqSummary(gds.fn, "$alt")
seqSummary(gds.fn, "$info")
seqSummary(gds.fn, "$format")
seqSummary(gds.fn, "$digest")

# open a GDS file
f <- seqOpen(gds.fn)

# get 'sample.id'
samp.id <- seqGetData(f, "sample.id")
# get 'variant.id'
variant.id <- seqGetData(f, "variant.id")

# set sample and variant filters
seqSetFilter(f, sample.id=samp.id[c(2,4,6,8,10)])
set.seed(100)
seqSetFilter(f, variant.id=sample(variant.id, 10))

seqSummary(f, "genotype")

# close a GDS file
seqClose(f)
```

---

seqSystem

*Get the parameters in the GDS system*

---

**Description**

Get a list of parameters in the GDS system

**Usage**

```
seqSystem()
```

**Value**

A list including

```
num.logical.core
```

the number of logical cores

```
compiler.flag
```

SIMD instructions supported by the compiler

```
options
```

list all options associated with SeqArray GDS format or packages

**Author(s)**

Xiuwen Zheng

**Examples**

```
seqSystem()
```

---

```
seqTranspose
```

*Transpose Data Array*

---

**Description**

Transpose data array or matrix for possibly higher-speed access.

**Usage**

```
seqTranspose(gdsfile, var.name, compress=NULL, digest=TRUE, verbose=TRUE)
```

**Arguments**

```
gdsfile
```

a [SeqVarGDSClass](#) object

```
var.name
```

the variable name with '/' as a separator

```
compress
```

the compression option used in [add.gdsn](#); or determine automatically if NULL

```
digest
```

a logical value (TRUE/FALSE) or a character ("md5", "sha1", "sha256", "sha384" or "sha512"); add md5 hash codes to the GDS file if TRUE or a digest algorithm is specified

```
verbose
```

if TRUE, show information

**Details**

It is designed for possibly higher-speed access. More details will be provided in the future version.

**Value**

None.

**Author(s)**

Xiuwen Zheng

**See Also**[seqGetData](#), [seqApply](#)**Examples**

```
# the VCF file
(vcf.fn <- seqExampleFileName("vcf"))

# convert
seqVCF2GDS(vcf.fn, "tmp.gds", storage.option="ZIP_RA")

# list the structure of GDS variables
f <- seqOpen("tmp.gds", FALSE)
f

seqTranspose(f, "genotype/data")
f

# the original array
index.gdsn(f, "genotype/data")
# the transposed array
index.gdsn(f, "genotype/~data")

# close
seqClose(f)

# delete the temporary file
unlink("tmp.gds")
```

seqUnitApply

*Apply Function Over Variant Units***Description**

Applies a user-defined function to each variant unit.

**Usage**

```
seqUnitApply(gdsfile, units, var.name, FUN, as.is=c("none", "list", "unlist"),
             parallel=FALSE, ..., .bl_size=256L, .progress=FALSE, .useraw=FALSE,
             .padNA=TRUE, .tolist=FALSE, .envir=NULL)
```

**Arguments**

gdsfile	a <a href="#">SeqVarGDSClass</a> object
units	a list of units of selected variants, with S3 class <a href="#">SeqUnitListClass</a>
var.name	the variable name(s), see details
FUN	the function to be applied

as.is	returned value: a list, an integer vector, etc; return nothing by default as.is="none"; as.is can be a <a href="#">connection</a> object, or a GDS node <a href="#">gdsn.class</a> object; if "unlist" is used, produces a vector which contains all the atomic components, via <code>unlist(..., recursive=FALSE)</code>
parallel	FALSE (serial processing), TRUE (multicore processing), numeric value or other value; parallel is passed to the argument <code>cl</code> in <a href="#">seqParallel</a> , see <a href="#">seqParallel</a> for more details.
.bl_size	chunk size, the increment for load balancing, 256 for units
.progress	if TRUE, show progress information
.useraw	TRUE, force to use RAW instead of INTEGER for genotypes and dosages; FALSE, use INTEGER; NA, use RAW instead of INTEGER if possible; for genotypes, 0xFF is missing value if RAW is used
.padNA	TRUE, pad a variable-length vector with NA if the number of data points for each variant is not greater than 1
.tolist	if TRUE, return a list of vectors instead of the structure <code>list(length, data)</code> for variable-length data
.envir	NULL, an environment object, or a list/data.frame
...	optional arguments to FUN

### Details

The variable name should be "sample.id", "variant.id", "position", "chromosome", "allele", "genotype", "annotation/id", "annotation/qual", "annotation/filter", "annotation/info/VARIABLE\_NAME" or "annotation/format/VARIABLE\_NAME".

"@genotype", "annotation/info/@VARIABLE\_NAME" or "annotation/format/@VARIABLE\_NAME" are used to obtain the index associated with these variables.

"\$dosage" is also allowed for the dosages of reference allele (integer: 0, 1, 2 and NA for diploid genotypes).

"\$dosage\_alt" returns a RAW/INTEGER matrix for the dosages of alternative allele without distinguishing different alternative alleles.

"\$dosage\_sp" returns a sparse matrix (dgCMatrix) for the dosages of alternative allele without distinguishing different alternative alleles.

"\$num\_allele" returns an integer vector with the numbers of distinct alleles.

"\$ref" returns a character vector of reference alleles

"\$alt" returns a character vector of alternative alleles (delimited by comma)

"\$chrom\_pos" returns characters with the combination of chromosome and position, e.g., "1:1272721".

"\$chrom\_pos\_allele" returns characters with the combination of chromosome, position and alleles, e.g., "1:1272721\_A\_G" (i.e., chr:position\_REF\_ALT).

"\$variant\_index" returns the indices of selected variants starting from 1, and "\$sample\_index" returns the indices of selected samples starting from 1.

### Value

A vector, a list of values or none.

### Author(s)

Xiuwen Zheng

**See Also**

[seqUnitSlidingWindows](#), [seqUnitFilterCond](#)

**Examples**

```
# open the GDS file
gdsfile <- seqOpen(seqExampleFileName("gds"))

# variant units via sliding windows
units <- seqUnitSlidingWindows(gdsfile)

v1 <- seqUnitApply(gdsfile, units, "genotype", function(x) dim(x)[3L],
  as.is="unlist", .progress=TRUE)
v2 <- seqUnitApply(gdsfile, units, "genotype", function(x) dim(x)[3L],
  as.is="unlist", parallel=2, .progress=TRUE)

all(v1 == lengths(units$index))
all(v1 == v2)

# call with an external R variable
ext <- list(x=1:1348/10)
v3 <- seqUnitApply(gdsfile, units, "$:x", function(x) x,
  as.is="list", .progress=TRUE, .envir=ext)
head(units$index)
head(v3)

table(sapply(seq_along(units$index), function(i) all(units$index[[i]] == v3[[i]]*10)))
# all TRUE

# close the GDS file
seqClose(gdsfile)
```

---

seqUnitCreate

*Subset and merge the units*


---

**Description**

Subset and merge the variant unit(s).

**Usage**

```
seqUnitCreate(idx, desp=NULL)
seqUnitSubset(units, i)
seqUnitMerge(ut1, ut2)
```

**Arguments**

idx	a list of numeric indexing vectors for specifying variants
desp	a data.frame for annotating the variant sets
units	a list of units of selected variants, with S3 class SeqUnitListClass

ut1                    a list of units of selected variants, with S3 class SeqUnitListClass  
 ut2                    a list of units of selected variants, with S3 class SeqUnitListClass  
 i                      a numeric or logical vector for indices specifying elements

**Value**

The variant unit of SeqUnitListClass.

**Author(s)**

Xiuwen Zheng

**See Also**

[seqUnitSlidingWindows](#), [seqUnitFilterCond](#)

**Examples**

```
# open the GDS file
gdsfile <- seqOpen(seqExampleFileName("gds"))

# variant units via sliding windows
units <- seqUnitSlidingWindows(gdsfile)

(u1 <- seqUnitSubset(units, 1:10))
(u2 <- seqUnitSubset(units, 30:39))

seqUnitMerge(u1, u2)

seqUnitCreate(list(1:10, 20:30), data.frame(gene=c("g1", "g2")))

# close the GDS file
seqClose(gdsfile)
```

---

seqUnitFilterCond      *Filter unit variants*

---

**Description**

Filters out the unit variants according to MAF, MAC and missing rates.

**Usage**

```
seqUnitFilterCond(gdsfile, units, maf=NaN, mac=1L, missing.rate=NaN,
  minsize=1L, parallel=seqGetParallel(), balancing=NA, verbose=TRUE)
```

**Arguments**

<code>gdsfile</code>	a <a href="#">SeqVarGDSClass</a> object
<code>units</code>	a list of units of selected variants, with S3 class <a href="#">SeqUnitListClass</a>
<code>maf</code>	minimum minor reference allele frequency, or a range of MAF <code>maf[1] &lt;= ... &lt; maf[2]</code>
<code>mac</code>	minimum minor reference allele count, or a range of MAC <code>mac[1] &lt;= ... &lt; mac[2]</code>
<code>missing.rate</code>	maximum missing genotype rate
<code>minsize</code>	the minimum of unit size
<code>parallel</code>	FALSE (serial processing), TRUE (multicore processing), numeric value or other value; <code>parallel</code> is passed to the argument <code>cl</code> in <a href="#">seqParallel</a> , see <a href="#">seqParallel</a> for more details.
<code>balancing</code>	whether to perform workload balancing or not, only applicable when multiple cores are used; if NA, use TRUE as a default until <code>getOption("seqarray.balancing")</code> is set and not TRUE
<code>verbose</code>	if TRUE, show information

**Value**

A S3 object with the class name "SeqUnitListClass" and two components (desp and index): the first is a data.frame with columns "chr", "start" and "end", and the second is list of integer vectors (the variant indices).

**Author(s)**

Xiuwen Zheng

**See Also**

[seqUnitApply](#), [seqUnitCreate](#), [seqUnitSubset](#), [seqUnitMerge](#)

**Examples**

```
# open the GDS file
gdsfile <- seqOpen(seqExampleFileName("gds"))

unit1 <- seqUnitSlidingWindows(gdsfile)
unit1 # "desp" "index"

# only rare variants
newunit <- seqUnitFilterCond(gdsfile, unit1, maf=c(0, 0.01))
newunit

# excluded variants
exvar <- setdiff(unique(unlist(unit1$index)), unique(unlist(newunit$index)))

seqSetFilter(gdsfile, variant.sel=exvar)
maf <- seqAlleleFreq(gdsfile, minor=TRUE)
table(maf > 0)
summary(maf[maf > 0]) # > 0.01
```

```
# close the GDS file
seqClose(gdsfile)
```

---

seqUnitSlidingWindows *Sliding units of selected variants*

---

### Description

Generates units of selected variants via sliding windows.

### Usage

```
seqUnitSlidingWindows(gdsfile, win.size=5000L, win.shift=2500L, win.start=0L,
  dup.rm=TRUE, verbose=TRUE)
```

### Arguments

gdsfile	a <a href="#">SeqVarGDSClass</a> object
win.size	window size in basepair
win.shift	the shift of sliding window in basepair
win.start	the start position in basepair
dup.rm	if TRUE, remove duplicate and zero-length windows
verbose	if TRUE, display information

### Value

A S3 object with the class name "SeqUnitListClass" and two components (desp and index): the first is a data.frame with columns "chr", "start" and "end", and the second is list of integer vectors (the variant indices).

### Author(s)

Xiuwen Zheng

### See Also

[seqUnitApply](#), [seqUnitFilterCond](#)

### Examples

```
# open the GDS file
gdsfile <- seqOpen(seqExampleFileName("gds"))

v <- seqUnitSlidingWindows(gdsfile)
v # "desp" "index"

# close the GDS file
seqClose(gdsfile)
```

---

`SeqVarGDSClass`*SeqVarGDSClass*

---

## Description

A `SeqVarGDSClass` object provides access to a GDS file containing Variant Call Format (VCF) data. It extends [gds.class](#).

## Details

A `SeqArray` GDS file is created from a VCF file with [seqVCF2GDS](#). This file can be opened with [seqOpen](#) to create a `SeqVarGDSClass` object.

## Accessors

In the following code snippets `x` is a `SeqVarGDSClass` object.

`granges(x)` Returns the chromosome and position of variants as a [GRanges](#) object. Names correspond to the `variant.id`.

`ref(x)` Returns the reference alleles as a [DNAStrngSet](#).

`alt(x)` Returns the alternate alleles as a [DNAStrngSetList](#).

`qual(x)` Returns the quality scores.

`filt(x)` Returns the filter data.

`fixed(x)` Returns the fixed fields (`ref`, `alt`, `qual`, `filt`).

`header(x)` Returns the header as a [DataFrameList](#).

`rowRanges(x)` Returns a [GRanges](#) object with metadata.

`colData(x)` Returns a [DataFrame](#) with sample identifiers and any information in the 'sample.annotation' node.

`info(x, info=NULL)` Returns the info fields as a [DataFrame](#). `info` is a character vector with the names of fields to return (default is to return all).

`geno(x, geno=NULL)` Returns the geno (format) fields as a [SimpleList](#). `geno` is a character vector with the names of fields to return (default is to return all).

Other data can be accessed with [seqGetData](#).

## Coercion methods

In the following code snippets `x` is a `SeqVarGDSClass` object.

```
. seqAsVCF(x, chr.prefix="", info=NULL, geno=NULL):
```

## Author(s)

Stephanie Gogarten, Xiuwen Zheng

## See Also

[gds.class](#), [seqOpen](#)

**Examples**

```

gds <- seqOpen(seqExampleFileName("gds"))
gds

## sample ID
head(seqGetData(gds, "sample.id"))

## variants
granges(gds)

## Not run:
## alleles as comma-separated character strings
head(seqGetData(gds, "allele"))

## alleles as DNASTringSet or DNASTringSetList
ref(gds)
v <- alt(gds)

## genotype
geno <- seqGetData(gds, "genotype")
dim(geno)
## dimensions are: allele, sample, variant
geno[1,1:10,1:5]

## rsID
head(seqGetData(gds, "annotation/id"))

## alternate allele count
head(seqGetData(gds, "annotation/info/AC"))

## individual read depth
depth <- seqGetData(gds, "annotation/format/DP")
names(depth)
## VCF header defined DP as variable-length data
table(depth$length)
## all length 1, so depth$data should be a sample by variant matrix
dim(depth$data)
depth$data[1:10,1:5]

## End(Not run)

seqClose(gds)

```

---

seqVCF2GDS

*Reformat VCF Files*


---

**Description**

Reformats Variant Call Format (VCF) files.

**Usage**

```

seqVCF2GDS(vcf.fn, out.fn, header=NULL, storage.option="LZMA_RA",
           info.import=NULL, fmt.import=NULL, genotype.var.name="GT",

```

```

ploidy=NA_integer_, ignore.chr.prefix="chr",
scenario=c("general", "imputation"), reference=NULL, start=1L, count=-1L,
variant_count=NA_integer_, optimize=TRUE, raise.error=TRUE, digest=TRUE,
use.Rsamtools=NA, parallel=FALSE, verbose=TRUE)
seqBCF2GDS(bcf.fn, out.fn, header=NULL, storage.option="LZMA_RA",
info.import=NULL, fmt.import=NULL, genotype.var.name="GT",
ploidy=NA_integer_, ignore.chr.prefix="chr",
scenario=c("general", "imputation"), reference=NULL, optimize=TRUE,
raise.error=TRUE, digest=TRUE, bcftools="bcftools", verbose=TRUE)

```

## Arguments

<code>vcf.fn</code>	the file name(s) of VCF format; or a <a href="#">connection</a> object
<code>bcf.fn</code>	a file name of binary VCF format (BCF)
<code>out.fn</code>	the file name of output GDS file
<code>header</code>	if NULL, header is set to be <a href="#">seqVCF_Header</a> ( <code>vcf.fn</code> )
<code>storage.option</code>	specify the storage and compression option, "ZIP_RA" ( <a href="#">seqStorageOption</a> ("ZIP_RA")); or "LZMA_RA" to use LZMA compression algorithm with higher compression ratio by default; or "LZ4_RA" to use an extremely fast compression and decompression algorithm. "ZIP_RA.max", "LZMA_RA.max" and "LZ4_RA.max" correspond to the algorithms with a maximum compression level; the suffix "_RA" indicates that fine-level random access is available; see more details at <a href="#">seqStorageOption</a>
<code>info.import</code>	characters, the variable name(s) in the INFO field for import; or NULL for all variables
<code>fmt.import</code>	characters, the variable name(s) in the FORMAT field for import; or NULL for all variables
<code>genotype.var.name</code>	the ID for genotypic data in the FORMAT column; "GT" by default (in VCF v4)
<code>ploidy</code>	specify the ploidy by users or NA to determine the ploidy from the VCF header
<code>ignore.chr.prefix</code>	a vector of character, indicating the prefix of chromosome which should be ignored, e.g., "chr"; it is not case-sensitive
<code>scenario</code>	"general": use float32 to store floating-point numbers (by default); "imputation": use packedreal16 to store DS and GP in the FORMAT field with four decimal place accuracy
<code>reference</code>	genome reference, like "hg19", "GRCh37"; if the genome reference is not available in VCF files, users could specify the reference here
<code>start</code>	the starting variant if importing part of VCF files
<code>count</code>	the maximum count of variant if importing part of VCF files, -1 indicates importing to the end
<code>variant_count</code>	NA_integer_ (default) or a numeric vector specifying the numbers of variants in the VCF file(s) in <code>vcf.fn</code> ; only applicable when multiple cores are used; if the number of variants is known, the conversion can skip counting the variants before splitting the file(s); <code>variant_count</code> could be an approximate
<code>optimize</code>	if TRUE, optimize the access efficiency by calling <a href="#">cleanup.gds</a>
<code>raise.error</code>	TRUE: throw an error if numeric conversion fails; FALSE: get missing value if numeric conversion fails

digest	a logical value (TRUE/FALSE) or a character ("md5", "sha1", "sha256", "sha384" or "sha512"); add md5 hash codes to the GDS file if TRUE or a digest algorithm is specified
use_Rsamtools	only applicable when multiple cores are used; use_Rsamtools is passed to <a href="#">seqVCF_Header</a> to get the total number of variants; NA: using Rsamtools when it is installed; FALSE: not use the Rsamtools package; TRUE: to use Rsamtools, if it is not installed, the function fails
parallel	FALSE (serial processing), TRUE (parallel processing), a numeric value indicating the number of cores, or a cluster object for parallel processing; parallel is passed to the argument c1 in <a href="#">seqParallel</a> , see <a href="#">seqParallel</a> for more details
verbose	if TRUE, show information
bcftools	the path of the program bcftools

### Details

If there are more than one files in `vcf.fn`, `seqVCF2GDS` will merge all VCF files together if they contain the same samples. It is useful to merge multiple VCF files if variant data are split by chromosomes.

The real numbers in the VCF file(s) are stored in 32-bit floating-point format by default. Users can set `storage.option=seqStorageOption(float.mode="float64")` to switch to 64-bit floating point format. Or packed real numbers can be adopted by setting `storage.option=seqStorageOption(float.mode="pa`

By default, the compression method is "LZMA\_RA" (<https://tukaani.org/xz/>, LZMA algorithm with default compression level + independent data blocks for fine-level random access). Users can maximize the compression ratio by `storage.option="LZMA_RA.max"` or `storage.option=seqStorageOption("LZMA_RA.max")`. LZMA is known to have higher compression ratio than the zlib algorithm. LZ4 (<https://github.com/lz4/lz4>) is an option via `storage.option="LZ4_RA"` or `storage.option=seqStorageOption("LZ4_RA")`.

If multiple cores/processes are specified in `parallel`, all VCF files are scanned to calculate the total number of variants before format conversion, and then split by the number of cores/processes.

`storage.option="Ultra"` and `storage.option="UltraMax"` need much larger memory than other compression methods. Users may consider using [seqRecompress](#) to recompress the GDS file after calling `seqVCF2GDS()` with `storage.option="ZIP_RA"`, since `seqRecompress()` compresses data nodes one by one, taking much less memory than "Ultra" and "UltraMax".

If `storage.option="LZMA_RA"` runs out of memory (e.g., there are too many annotation fields in the VCF file), users could use `storage.option="ZIP_RA"` and then call `seqRecompress(, compress="LZMA")`.

### Value

Return the file name of GDS format with an absolute path.

### Author(s)

Xiuwen Zheng

### References

Danecek, P., Auton, A., Abecasis, G., Albers, C.A., Banks, E., DePristo, M.A., Handsaker, R.E., Lunter, G., Marth, G.T., Sherry, S.T., et al. (2011). The variant call format and VCFtools. *Bioinformatics* 27, 2156-2158.

**See Also**

[seqVCF\\_Header](#), [seqStorageOption](#), [seqMerge](#), [seqGDS2VCF](#), [seqRecompress](#)

**Examples**

```
# the VCF file
vcf.fn <- seqExampleFileName("vcf")

# conversion
seqVCF2GDS(vcf.fn, "tmp.gds", storage.option="ZIP_RA")

# conversion in parallel
seqVCF2GDS(vcf.fn, "tmp_p2.gds", storage.option="ZIP_RA", parallel=2L)

# display
(f <- seqOpen("tmp.gds"))
seqClose(f)

# convert without the INFO fields
seqVCF2GDS(vcf.fn, "tmp.gds", storage.option="ZIP_RA",
  info.import=character(0))

# display
(f <- seqOpen("tmp.gds"))
seqClose(f)

# convert without the INFO and FORMAT fields
seqVCF2GDS(vcf.fn, "tmp.gds", storage.option="ZIP_RA",
  info.import=character(0), fmt.import=character(0))

# display
(f <- seqOpen("tmp.gds"))
seqClose(f)

# delete the temporary file
unlink(c("tmp.gds", "tmp_p2.gds"), force=TRUE)
```

---

seqVCF\_Header

---

*Parse the Header of a VCF/BCF File*


---

**Description**

Parses the meta-information lines of a VCF or BCF file.

**Usage**

```
seqVCF_Header(vcf.fn, getnum=FALSE, use_Rsamtools=NA, parallel=FALSE,
  verbose=TRUE)
```

**Arguments**

<code>vcf.fn</code>	the file name of VCF or BCF format; or a <a href="#">connection</a> object for VCF format
<code>getnum</code>	if TRUE, return the total number of variants
<code>use_Rsamtools</code>	only applicable when <code>getnum=TRUE</code> and multiple cores are used; NA: using Rsamtools when it is installed; FALSE: not use the Rsamtools package; TRUE: to use Rsamtools, if it is not installed, the function fails
<code>parallel</code>	FALSE (serial processing), TRUE (parallel processing), a numeric value indicating the number of cores, or a cluster object for parallel processing; <code>parallel</code> is passed to the argument <code>c1</code> in <a href="#">seqParallel</a> , see <a href="#">seqParallel</a> for more details
<code>verbose</code>	when <code>getnum=TRUE</code> and <code>verbose=TRUE</code> , show the progress information for scanning the file

**Details**

The ID description contains four columns: ID – variable name; Number – the number of elements, see the webpage of the 1000 Genomes Project; Type – data type; Description – a variable description.

If multiple cores are used to get the total number of variants, the Rsamtools package should be installed, and the indexing file (i.e., `.csi` or `.tbi`) should be available along with `vcf.fn`.

**Value**

Return a list (with a class name "SeqVCFHeaderClass", S3 object):

<code>fileformat</code>	the file format
<code>info</code>	the ID description in the INFO field
<code>filter</code>	the ID description in the FILTER field
<code>format</code>	the ID description in the FORMAT field
<code>alt</code>	the ID description in the ALT field
<code>contig</code>	the description in the contig field
<code>assembly</code>	the link of assembly
<code>reference</code>	genome reference, or NULL if unknown
<code>header</code>	the other header lines
<code>ploidy</code>	ploidy, two for humans
<code>num.sample</code>	the number of samples
<code>num.variant</code>	the number of variants, applicable only if <code>getnum=TRUE</code>
<code>sample.id</code>	a vector of sample IDs in the VCF/BCF file

**Author(s)**

Xiuwen Zheng

**References**

Danecek, P., Auton, A., Abecasis, G., Albers, C.A., Banks, E., DePristo, M.A., Handsaker, R.E., Lunter, G., Marth, G.T., Sherry, S.T., et al. (2011). The variant call format and VCFtools. *Bioinformatics* 27, 2156-2158.

**See Also**

[seqVCF\\_SampID](#), [seqVCF2GDS](#)

**Examples**

```
# the VCF file
(vcf.fn <- seqExampleFileName("vcf"))
# or vcf.fn <- "C:/YourFolder/Your_VCF_File.vcf"

# get sample id
seqVCF_Header(vcf.fn, getnum=TRUE)

# use a connection object
f <- file(vcf.fn, "r")
seqVCF_Header(f, getnum=TRUE)
close(f)
```

---

seqVCF_SampID	<i>Get the Sample IDs</i>
---------------	---------------------------

---

**Description**

Returns the sample IDs of a VCF file.

**Usage**

```
seqVCF_SampID(vcf.fn)
```

**Arguments**

vcf.fn            the file name, output a file of VCF format; or a [connection](#) object

**Author(s)**

Xiuwen Zheng

**References**

Danecek, P., Auton, A., Abecasis, G., Albers, C.A., Banks, E., DePristo, M.A., Handsaker, R.E., Lunter, G., Marth, G.T., Sherry, S.T., et al. (2011). The variant call format and VCFtools. *Bioinformatics* 27, 2156-2158.

**See Also**

[seqVCF\\_Header](#), [seqVCF2GDS](#)

**Examples**

```
# the VCF file
(vcf.fn <- seqExampleFileName("vcf"))

# get sample id
seqVCF_SampID(vcf.fn)
```

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