

Package ‘vcfppR’

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Title Rapid Manipulation of the Variant Call Format (VCF)

Version 0.8.1

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Description The 'vcfpp.h' (<<https://github.com/Zilong-Li/vcfpp>>) provides an easy-to-use 'C++' 'API' of 'htlib', offering full functionality for manipulating Variant Call Format (VCF) files. The 'vcfppR' package serves as the R bindings of the 'vcfpp.h' library, enabling rapid processing of both compressed and uncompressed VCF files. Explore a range of powerful features for efficient VCF data manipulation.

Encoding UTF-8

Depends R (>= 3.6.0)

RoxygenNote 7.3.2

Suggests knitr, codetools, rmarkdown, testthat (>= 3.0.0)

Config/testthat/edition 3

SystemRequirements libcurl: libcurl-devel (rpm) or
libcurl4-openssl-dev (deb), GNU make.

Imports Rcpp, methods, stats, utils

LinkingTo Rcpp

URL <https://github.com/Zilong-Li/vcfppR>

BugReports <https://github.com/Zilong-Li/vcfppR/issues>

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VignetteBuilder knitr

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vcfppR-package	<i>vcfppR: Rapid Manipulation of the Variant Call Format (VCF)</i>
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Description

The 'vcfpp.h' (<https://github.com/Zilong-Li/vcfpp>) provides an easy-to-use 'C++' 'API' of 'htslib', offering full functionality for manipulating Variant Call Format (VCF) files. The 'vcfppR' package serves as the R bindings of the 'vcfpp.h' library, enabling rapid processing of both compressed and uncompressed VCF files. Explore a range of powerful features for efficient VCF data manipulation.

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See Also

- Useful links:
- <https://github.com/Zilong-Li/vcfppR>
 - Report bugs at <https://github.com/Zilong-Li/vcfppR/issues>

vcfcomp

*Compare two VCF/BCF files reporting various statistics***Description**

Compare two VCF/BCF files reporting various statistics

Usage

```
vcfcomp(
  test,
  truth,
  formats = c("DS", "GT"),
  stats = "r2",
  by.sample = FALSE,
  by.variant = FALSE,
  flip = FALSE,
  names = NULL,
  bins = NULL,
  af = NULL,
  out = NULL,
  choose_random_start = FALSE,
  return_pse_sites = FALSE,
  ...
)
```

Arguments

test	path to the comparison file (test), which can be a VCF/BCF file, vcftable object or saved RDS file.
truth	path to the baseline file (truth), which can be a VCF/BCF file, vcftable object or saved RDS file.
formats	character vector. the FORMAT tags to extract for the test and truth respectively. default c("DS", "GT") extracts 'DS' of the target and 'GT' of the truth.
stats	the statistics to be calculated. supports the following. "r2": the Pearson correlation coefficient square. "f1": the F1-score, good balance between sensitivity and precision. "nrc": the Non-Reference Concordance rate "pse": the Phasing Switch Error rate
by.sample	logical. calculate sample-wise concordance, which can be stratified by MAF bin.
by.variant	logical. calculate variant-wise concordance, which can be stratified by MAF bin. If both by.sample and by.variant are FALSE, then do calculations for all samples and variants together in a bin.
flip	logical. flip the ref and alt variants
names	character vector. reset samples' names in the test VCF.

bins	numeric vector. break statistics into allele frequency bins.
af	file path with allele frequency or a RDS file with a saved object for af. Format of the text file: a space-separated text file with five columns and a header named 'chr' 'pos' 'ref' 'alt' 'af'
out	output prefix for saving objects into RDS file
choose_random_start	choose random start for stats="pse"
return_pse_sites	boolean. return phasing switch error sites
...	options passed to vcftable

Details

vcfcomp implements various statistics to compare two VCF/BCF files, e.g. report genotype concordance, correlation stratified by allele frequency.

Value

a list of various statistics

Author(s)

Zilong Li <zilong.dk@gmail.com>

Examples

```
library('vcfppR')
test <- system.file("extdata", "imputed.gt.vcf.gz", package="vcfppR")
truth <- system.file("extdata", "imputed.gt.vcf.gz", package="vcfppR")
samples <- "HG00133,HG00143,HG00262"
res <- vcfcomp(test, truth, stats="f1", samples=samples, setid=TRUE)
str(res)
```

vcfinfo	<i>read a INFO tag in the VCF/BCF into R data structure</i>
---------	---

Description

read a INFO tag in the VCF/BCF into R data structure

Usage

```
vcfinfo(
  vcffile,
  tag,
  region = "",
  vartype = "all",
  ids = NULL,
  qual = 0,
  pass = FALSE,
  setid = FALSE
)
```

Arguments

<code>vcffile</code>	path to the VCF/BCF file
<code>tag</code>	the INFO tag to extract.
<code>region</code>	region to subset in bcftools-like style: "chr1", "chr1:1-100000000"
<code>vartype</code>	restrict to specific type of variants. supports "snps", "indels", "sv", "multisnps", "multiallelics"
<code>ids</code>	character vector. restrict to sites with ID in the given vector. default NULL won't filter any sites.
<code>qual</code>	numeric. restrict to variants with QUAL > qual.
<code>pass</code>	logical. restrict to variants with FILTER = "PASS".
<code>setid</code>	logical. reset ID column as CHR_POS_REF_ALT.

Details

`vcfinfo` uses the C++ API of `vcfpp`, which is a wrapper of `htslib`, to read VCF/BCF files. Thus, it has the full functionalities of `htslib`, such as restrict to specific variant types, samples and regions. For the memory efficiency reason, the `vcfinfo` is designed to parse only one tag at a time in the INFO column of the VCF. Currently it does not support parsing a vector of values for a given INFO tag.

Value

Return a list containing the following components:

chr : character vector;
the CHR column in the VCF file

pos : character vector;
the POS column in the VCF file

id : character vector;
the ID column in the VCF file

ref : character vector;
the REF column in the VCF file

alt : character vector;
the ALT column in the VCF file

qual : character vector;
the QUAL column in the VCF file

filter : character vector;
the FILTER column in the VCF file

tag : vector of either integer, numeric or character values depending on the tag to extract;
a specify tag in the INFO column to be extracted

Author(s)

Zilong Li <zilong.dk@gmail.com>

Examples

```
library('vcfppR')
vcffile <- system.file("extdata", "raw.gt.vcf.gz", package="vcfppR")
res <- vcfinfo(vcffile, "AF", region = "chr21:1-5050000", vartype = "snps", pass = TRUE)
str(res)
```

vcfplot

Make sensible and beautiful plots based on various objects in vcfppR

Description

Make sensible and beautiful plots based on various objects in vcfppR

Usage

```
vcfplot(obj, which.sample = NULL, variant = c("SNP", "INDEL"), pop = NULL, ...)
```

Arguments

obj	object returned by vcftable, vcfcamp, vcfsummary
which.sample	which sample to be plotted. NULL will aggregate all samples.
variant	which types of variant are desired
pop	file contains population information
...	parameters passed to graphics

vcfpopgen	<i>count the heterozygous sites per sample in the VCF/BCF</i>
-----------	---

Description

count the heterozygous sites per sample in the VCF/BCF

Usage

```
vcfpopgen(  
  vcffile,  
  region = "",  
  samples = "-",  
  pass = FALSE,  
  qual = 0,  
  fun = "heterozygosity"  
)
```

Arguments

vcffile	path to the VCF/BCF file
region	region to subset like bcftools
samples	samples to subset like bcftools
pass	restrict to variants with FILTER==PASS
qual	restrict to variants with QUAL > qual.
fun	which popgen function to run. available functions are "heterozygosity".

Value

vcfpopgen a list containing the following components:

samples : character vector;
the samples ids in the VCF file after subsetting

hets : integer vector;
the counts of heterozygous sites of each sample in the same order as samples

Author(s)

Zilong Li <zilong.dk@gmail.com>

Examples

```
library('vcfppR')  
vcffile <- system.file("extdata", "raw.gt.vcf.gz", package="vcfppR")  
res <- vcfpopgen(vcffile)  
str(res)
```

vcfreader

*API for manipulating the VCF/BCF.***Description**

Type the name of the class to see the details and methods

Value

A C++ class with the following fields/methods for manipulating the VCF/BCF

Fields

new Constructor given a vcf file

- Parameter: vcffile - The path of a vcf file

new Constructor given a vcf file and the region

- Parameter: vcffile - The path of a vcf file
- Parameter: region - The region to be constrained

new Constructor given a vcf file, the region and the samples

- Parameter: vcffile - The path of a vcf file
- Parameter: region - The region to be constrained
- Parameter: samples - The samples to be constrained. Comma separated list of samples to include (or exclude with "^" prefix).

setRegion try to set specific region to work with. will throw errors if no index or region found. Use getStatus to check if the region is valid or empty!

getStatus return 1: region is valid and not empty. 0: region is valid but empty. -1: no index file. -2: region not found or invalid region form

variant Try to get next variant record. return FALSE if there are no more variants or hit the end of file, otherwise TRUE.

chr Return the CHROM field of current variant

pos Return the POS field of current variant

id Return the CHROM field of current variant

ref Return the REF field of current variant

alt Return the ALT field of current variant

qual Return the QUAL field of current variant

filter Return the FILTER field of current variant

info Return the INFO field of current variant

infoInt Return the tag value of integer type in INFO field of current variant

- Parameter: tag - The tag name to retrieve in INFO

infoFloat Return the tag value of float type in INFO field of current variant

- Parameter: tag - The tag name to retrieve in INFO

infoStr Return the tag value of string type in INFO field of current variant

- Parameter: tag - The tag name to retrieve in INFO

infoIntVec Return the tag value in a vector of integer type in INFO field of current variant

- Parameter: tag - The tag name to retrieve in INFO

infoFloatVec Return the tag value in a vector of float type in INFO field of current variant

- Parameter: tag - The tag name to retrieve in INFO

genotypes Return the genotype values in a vector of integers

- Parameter: collapse - Boolean value indicates wheather to collapse the size of genotypes, eg, return diploid genotypes.

formatInt Return the tag value of integer type for each sample in FORAMT field of current variant

- Parameter: tag - The tag name to retrieve in FORAMT

formatFloat Return the tag value of float type for each sample in FORAMT field of current variant

- Parameter: tag - The tag name to retrieve in FORAMT

formatStr Return the tag value of string type for each sample in FORAMT field of current variant

- Parameter: tag - The tag name to retrieve in FORAMT

isSNP Test if current variant is exculsively a SNP or not

isIndel Test if current variant is exculsively a INDEL or not

isSV Test if current variant is exculsively a SV or not

isMultiAllelics Test if current variant is exculsively a Multi Allelics or not

isMultiAllelicSNP Test if current variant is exculsively a Multi Biallelics (SNPs) or not

hasSNP Test if current variant has a SNP or not

hasINDEL Test if current variant has a INDEL or not

hasINS Test if current variant has a INS or not

hasDEL Test if current variant has a DEL or not

hasMNP Test if current variant has a MNP or not

hasBND Test if current variant has a BND or not

hasOTHER Test if current variant has a OTHER or not

hasOVERLAP Test if current variant has a OVERLAP or not

nsamples Return the number of samples

samples Return a vector of samples id

header Return the raw string of the vcf header

string Return the raw string of current variant including newline

line Return the raw string of current variant without newline

output Init an output object for streaming out the variants to another vcf

updateSamples update samples name in the output VCF

- Parameter: s - A comma-seperated string for new samples names

`write` Streaming out current variant the output vcf

`close` Close the connection to the output vcf

`setCHR` Modify the CHR of current variant

- Parameter: `s` - A string for CHR

`setID` Modify the ID of current variant

- Parameter: `s` - A string for ID

`setPOS` Modify the POS of current variant

- Parameter: `pos` - An integer for POS

`setRefAlt` Modify the REF and ALT of current variant

- Parameter: `s` - A string repeated by comma

`setInfoInt` Modify the given tag of INT type in the INFO of current variant

- Parameter: `tag` - A string for the tag name
- Parameter: `v` - An integer for the tag value

`setInfoFloat` Modify the given tag of FLOAT type in the INFO of current variant

- Parameter: `tag` - A string for the tag name
- Parameter: `v` - A double for the tag value

`setInfoStr` Modify the given tag of STRING type in the INFO of current variant

- Parameter: `tag` - A string for the tag name
- Parameter: `s` - A string for the tag value

`setPhasing` Modify the phasing status of each sample

- Parameter: `v` - An integer vector with size of the number of samples. only 1s and 0s are valid.

`setGenotypes` Modify the genotypes of current variant

- Parameter: `v` - An integer vector for genotypes. Use NA or -9 for missing value.

`setFormatInt` Modify the given tag of INT type in the FORMAT of current variant

- Parameter: `tag` - A string for the tag name
- Parameter: `v` - An integer for the tag value

`setFormatFloat` Modify the given tag of FLOAT type in the FORMAT of current variant

- Parameter: `tag` - A string for the tag name
- Parameter: `v` - A double for the tag value

`setFormatStr` Modify the given tag of STRING type in the FORMAT of current variant

- Parameter: `tag` - A string for the tag name
- Parameter: `s` - A string for the tag value

`rmInfoTag` Remove the given tag from the INFO of current variant

- Parameter: `s` - A string for the tag name

`rmFormatTag` Remove the given tag from the FORMAT of current variant

- Parameter: `s` - A string for the tag name

`setVariant` Modify current variant by adding a vcf line

- Parameter: s - A string for one line in the VCF

addINFO Add a INFO in the header of the vcf

- Parameter: id - A string for the tag name
- Parameter: number - A string for the number
- Parameter: type - A string for the type
- Parameter: desc - A string for description of what it means

addFORMAT Add a FORMAT in the header of the vcf

- Parameter: id - A string for the tag name
- Parameter: number - A string for the number
- Parameter: type - A string for the type
- Parameter: desc - A string for description of what it means

Examples

```
vcffile <- system.file("extdata", "raw.gt.vcf.gz", package="vcfppR")
br <- vcfreader$new(vcffile)
res <- rep(0L, br$nsamples())
while(br$variant()) {
  if(br$isSNP()) {
    gt <- br$genotypes(TRUE) == 1
    gt[is.na(gt)] <- FALSE
    res <- res + gt
  }
}
```

vcfsummary

summarize the various variant types at both variant level and sample level.

Description

summarize the various variant types at both variant level and sample level.

Usage

```
vcfsummary(
  vcffile,
  region = "",
  samples = "-",
  pass = FALSE,
  qual = 0,
  svtype = FALSE
)
```

Arguments

vcffile	path to the VCF/BCF file
region	region to subset like bcftools
samples	samples to subset like bcftools
pass	restrict to variants with FILTER==PASS
qual	restrict to variants with QUAL > qual.
svtype	summarize the variants with SVTYPE

Details

```
bcftools view -s "id01,id02" input.bcf.gz chr1:100000-20000
```

Value

vcfsummary a list containing the following components:

- summary** : named integer vector;
summarize the counts of each variant type
- samples** : character vector;
the samples ids in the VCF file after subsetting
- vartype** : integer vector;
the counts of the variant type at sample level in the same order as samples

Author(s)

Zilong Li <zilong.dk@gmail.com>

Examples

```
library('vcfppR')
svfile <- system.file("extdata", "sv.vcf.gz", package="vcfppR")
res <- vcfsummary(svfile, region = "chr21:1-100000000", svtype = TRUE)
str(res)
```

vcftable

read VCF/BCF contents into R data structure

Description

The swiss army knife for reading VCF/BCF into R data types rapidly and easily.

Usage

```
vcftable(
  vcffile,
  region = "",
  samples = "-",
  vartype = "all",
  format = "GT",
  ids = NULL,
  qual = 0,
  pass = FALSE,
  info = TRUE,
  collapse = TRUE,
  setid = FALSE,
  mac = 0,
  rmdup = FALSE
)
```

Arguments

vcffile	path to the VCF/BCF file
region	region to subset in bcftools-like style: "chr1", "chr1:1-10000000"
samples	samples to subset in bcftools-like style. comma separated list of samples to include (or exclude with "^" prefix). e.g. "id01,id02", "^id01,id02".
vartype	restrict to specific type of variants. supports "snps", "indels", "sv", "multisnps", "multiallelics"
format	the FORMAT tag to extract. default "GT" is extracted.
ids	character vector. restrict to sites with ID in the given vector. default NULL won't filter any sites.
qual	numeric. restrict to variants with QUAL > qual.
pass	logical. restrict to variants with FILTER = "PASS".
info	logical. drop INFO column in the returned list.
collapse	logical. It acts on the FORMAT. If the FORMAT to extract is "GT", the dim of raw genotypes matrix of diploid is (M, 2 * N), where M is #markers and N is #samples. default TRUE will collapse the genotypes for each sample such that the matrix is (M, N). Set this to FALSE if one wants to maintain the phasing order, e.g. "110" is parsed as c(1, 0) with collapse=FALSE. If the FORMAT to extract is not "GT", then with collapse=TRUE it will try to turn a list of the extracted vector into a matrix. However, this raises issues when one variant is mutliallelic resulting in more vaules than others.
setid	logical. reset ID column as CHR_POS_REF_ALT.
mac	integer. restrict to variants with minor allele count higher than the value.
rmdup	logical. remove duplicated sites by keeping the first occurrence of POS. (default: FALSE)

Details

vcftable uses the C++ API of vcfpp, which is a wrapper of htlib, to read VCF/BCF files. Thus, it has the full functionalities of htlib, such as restrict to specific variant types, samples and regions. For the memory efficiency reason, the vcftable is designed to parse only one tag at a time in the FORMAT column of the VCF. In default, only the matrix of genotypes, i.e. "GT" tag, are returned by vcftable, but there are many other tags supported by the format option.

Value

Return a list containing the following components:

samples : character vector;
the samples ids in the VCF file after subsetting

chr : character vector;
the CHR column in the VCF file

pos : character vector;
the POS column in the VCF file

id : character vector;
the ID column in the VCF file

ref : character vector;
the REF column in the VCF file

alt : character vector;
the ALT column in the VCF file

qual : character vector;
the QUAL column in the VCF file

filter : character vector;
the FILTER column in the VCF file

info : character vector;
the INFO column in the VCF file

format : matrix of either integer or numeric values depending on the tag to extract;
a specify tag in the FORMAT column to be extracted

Author(s)

Zilong Li <zilong.dk@gmail.com>

Examples

```
library('vcfppR')
vcffile <- system.file("extdata", "raw.gt.vcf.gz", package="vcfppR")
res <- vcftable(vcffile, "chr21:1-5050000", vartype = "snps")
str(res)
```

Description

Type the name of the class to see the details and methods

Value

A C++ class with the following fields/methods for writing the VCF/BCF

Fields

`new` Constructor given a vcf file

- Parameter: `vcffile` - The path of a vcf file. don't start with "~"
- Parameter: `version` - The version of VCF specification

`addContig` Add a Contig in the header of the vcf

- Parameter: `str` - A string for the CONTIG name

`addFILTER` Add a FILTER in the header of the vcf

- Parameter: `id` - A string for the FILTER name
- Parameter: `desc` - A string for description of what it means

`addINFO` Add a INFO in the header of the vcf

- Parameter: `id` - A string for the tag name
- Parameter: `number` - A string for the number
- Parameter: `type` - A string for the type
- Parameter: `desc` - A string for description of what it means

`addFORMAT` Add a FORMAT in the header of the vcf

- Parameter: `id` - A string for the tag name
- Parameter: `number` - A string for the number
- Parameter: `type` - A string for the type
- Parameter: `desc` - A string for description of what it means

`addSample` Add a SAMPLE in the header of the vcf

- Parameter: `str` - A string for a SAMPLE name

`addLine` Add a line in the header of the vcf

- Parameter: `str` - A string for a line in the header of VCF

`writeline` Write a variant record given a line

- Parameter: `line` - A string for a line in the variant of VCF. Not ended with "newline"

`close` Close and save the vcf file

Examples

```
outvcf <- file.path(paste0(tempfile(), ".vcf.gz"))
bw <- vcfwriter$new(outvcf, "VCF4.1")
bw$addContig("chr20")
bw$addFORMAT("GT", "1", "String", "Genotype");
bw$addSample("NA12878")
s1 <- "chr20\t2006060\t.\tG\tC\t100\tPASS\t.\tGT\t1|0"
bw$writeline(s1)
bw$close()
```

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