

# RBcf: An VCF API for R.

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## 1 Abstract

RBcf uses the Htslib C API for parsing VCF and BCF files. This API was written by a regular user of the htsjdk library who doesn't like R.

A list of functions is available at: <https://github.com/lindenb/rbcf/blob/master/R/rbcf.R>

## 2 Examples

### 2.1 Show Htslib and Rbcf versions

Code:

```
# load the library
library(rbcf)
#print the version of the associated htslib
paste("HTSLIB:",htslib.version())
#print the version of rbcf
paste("RBCF:",rcbf.version())
```

Output:

```
[1] "HTSLIB: 1.10.2"
[1] "RBCF: 0.0-1"
```

### 2.2 Open and close a VCF file

Code:

```
# load rbcf
library(rbcf)
# we don't need the index for this file
```

```
fp <- bcf.open("./data/rotavirus_rf.01.vcf",FALSE)
# dispose the vcf reader
bcf.close(fp)
print("Done.")
```

Output:

```
[1] TRUE
[1] "Done."
```

## 2.3 Print the INFOs in the VCF header

Code:

```
# load rbcf
library(rbcf)
# we don't need the index for this file
fp <- bcf.open("./data/rotavirus_rf.01.vcf",FALSE)
bcf.infos(fp)
# dispose the vcf reader
bcf.close(fp)
# print the table
```

Output:

	ID	Number	Type
INDEL	INDEL	0	Flag
IDV	IDV	1	Integer
IMF	IMF	1	Float
DP	DP	1	Integer
VDB	VDB	1	Float
RPB	RPB	1	Float
MQB	MQB	1	Float
BQB	BQB	1	Float
MQSB	MQSB	1	Float
SGB	SGB	1	Float
MQOF	MQOF	1	Float
ICB	ICB	1	Float
HOB	HOB	1	Float
AC	AC	A	Integer
AN	AN	1	Integer
DP4	DP4	4	Integer
MQ	MQ	1	Integer

```

INDEL "Indicates that the variant is an indel"
IDV "Maximum number of reads"
IMF "Maximum fraction of reads"
DP
VDB "Variant Distance Bias for filtering splice-site artefacts in RNA-seq data"
RPB "Mann-Whitney U test of Read Position Bias"
MQB "Mann-Whitney U test of Mapping Quality Bias"
BQB "Mann-Whitney U test of Base Quality Bias"
MQSB "Mann-Whitney U test of Mapping Quality vs Strand Bias"
SGB "Segregation Bias"
MQOF "Fraction of MQ0 reads"
ICB "Inbreeding Coefficient Binomial test"
HOB "Bias in the number of HOMs"
AC "Allele count in genotypes for each ALT allele, in the s"
AN "Total number of alleles"
DP4 "Number of high-quality ref-forward , ref-reverse, alt-forward and alt-reverse"
MQ "Average mapping quality"
[1] TRUE

```

## 2.4 Print the FORMATS in the VCF header

Code:

```

# load rbcf
library(rbcf)
# we don't need the index for this file
fp <- bcf.open("./data/rotavirus_rf.01.vcf",FALSE)
bcf.formats(fp)
# dispose the vcf reader
bcf.close(fp)

```

Output:

ID	Number	Type	Description
PL	PL	G Integer	"List of Phred-scaled genotype likelihoods"
GT	GT	1 String	"Genotype"
[1]	TRUE		

## 2.5 Print the FILTERs in the VCF header

Code:

```
# load rbcf
library(rbcf)
# we don't need the index for this file
fp <- bcf.open("./data/gnomad.exomes.r2.0.1.sites.bcf",FALSE)
bcf.filters(fp)
# dispose the vcf reader
bcf.close(fp)
```

Output:

```

                                ID
PASS                                PASS
ACO                                ACO
InbreedingCoeff InbreedingCoeff
LCR                                LCR
RF                                RF
SEGDUP                            SEGDUP

PASS
ACO                                "Allele_Count_is_zero_(i.e._no_high-confidence_genotype_(GQ_>=_2
InbreedingCoeff_
LCR_
RF                                "Failed_random_forests_filt
SEGDUP_
[1] TRUE
```

## 2.6 Print the Samples in the VCF header

The samples are defined in the '#CHROM' line of the VCF **Code**:

```
# load rbcf
library(rbcf)
# we don't need the index for this file
fp <- bcf.open("./data/rotavirus_rf.01.vcf",FALSE)
# print the number of samples
paste("Number_of_samples:",bcf.nsamples(fp))
# get the name for the 1st sample
paste("First_sample:",bcf.sample.at(fp,1))
# get the 1-based index for the samples
bcf.sample2index(fp,c("S1","S2","S3","missing"))
# get all the samples
bcf.samples(fp)
```

```
# dispose the vcf reader
bcf.close(fp)
```

Output:

```
[1] "Number_of_samples: 5"
[1] "First_sample: S1"
      S1      S2      S3 missing
      1      2      3      0
[1] "S1" "S2" "S3" "S4" "S5"
[1] TRUE
```

## 2.7 Print the Dictionary in the VCF header

Code:

```
# load rbcf
library(rbcf)
# we don't need the index for this file
fp <- bcf.open("./data/rotavirus_rf.01.vcf",FALSE)
bcf.dictionary(fp)
# dispose the vcf reader
bcf.close(fp)
```

Output:

```
      chrom size
RF01  RF01 3302
RF02  RF02 2687
RF03  RF03 2592
RF04  RF04 2362
RF05  RF05 1579
RF06  RF06 1356
RF07  RF07 1074
RF08  RF08 1059
RF09  RF09 1062
RF10  RF10  751
RF11  RF11  666
[1] TRUE
```

## 2.8 Print the Indexed Chromosomes

Code:

```
# load rbcf
library(rbcf)
# Open the indexed VCF
fp <- bcf.open("./data/rotavirus_rf.02.vcf.gz")
# get the indexed contigs
bcf.contigs(fp)
# dispose the vcf reader
bcf.close(fp)
```

Output:

```
[1] "RF01" "RF02" "RF03" "RF04" "RF05" "RF06" "RF07" "RF08" "RF09" "RF10"
[11] "RF11"
[1] TRUE
```

## 2.9 Scanning the variants

Code:

```
# load rbcf
library(rbcf)

# create a function counting variants in a VCF
count.variants<-function(filename) {
  # we don't need the index for this file
  fp <- bcf.open(filename,FALSE)
  # number of variants
  n<-0
  # loop while we can read a variant
  while(!is.null(vc<-bcf.next(fp))) {
    # increment the count
    n<-n+1
  }
  # dispose the vcf reader
  bcf.close(fp)
  # return the number of variant
  n
}

# filenames
vcfs<-c(
  "./data/gnomad.exomes.r2.0.1.sites.bcf",
```

```

        "/data/rotavirus_rf.01.vcf",
        "/data/rotavirus_rf.02.vcf.gz",
        "/data/rotavirus_rf.03.vcf.gz",
        "/data/rotavirus_rf.04.bcf"
    )
# print the number of variants for each vcf
for(f in vcfs) {
    cat(paste(f,"_",count.variants(f),"\n"))
}

```

Output:

```

./data/gnomad.exomes.r2.0.1.sites.bcf    50
./data/rotavirus_rf.01.vcf    45
./data/rotavirus_rf.02.vcf.gz    45
./data/rotavirus_rf.03.vcf.gz    45
./data/rotavirus_rf.04.bcf    45

```

## 2.10 Scanning the variants

Code:

```

# load rbcf
library(rbcf)

# create a function counting variants in a VCF
count.variants<-function(filename,predicate) {
    # we don't need the index for this file
    fp <- bcf.open(filename,FALSE)
    # number of variants
    n<-0
    # loop while we can read a variant
    while(!is.null(vc<-bcf.next(fp))) {
        # test the variant
        if(predicate(vc)) {
            # increment the count
            n<-n+1
        }
    }
    # dispose the vcf reader
    bcf.close(fp)
    # return the number of variant
}

```

```

    n
}

# A vcf
filename <- "./data/gnomad.exomes.r2.0.1.sites.bcf"
# filters
filters<-list(
  list("desc"="accept_all","predicate"=function(ctx) {TRUE} ),
  list("desc"="accept_none","predicate"=function(ctx) {FALSE} ),
  list("desc"="CHROM_is_1","predicate"=function(ctx) { variant.contig(ctx)%
  list("desc"="POS_is_even","predicate"=function(ctx) { (variant.pos(ctx)%
  list("desc"="PASS_filter","predicate"=function(ctx) {!variant.is.filtere
  list("desc"="count(FILTER)>1","predicate"=function(ctx) {length(variant.
  list("desc"="FILTER_contains_SEGDUP","predicate"=function(ctx) {variant.
  list("desc"="SNP","predicate"=function(ctx) {variant.is.snp(ctx)} ),
  list("desc"="POS!=END","predicate"=function(ctx) { variant.pos(ctx)!=var
  list("desc"="not_diallelic","predicate"=function(ctx) {variant.nalleles(
  list("desc"="REF_is_A","predicate"=function(ctx) {variant.reference(ct
  list("desc"="any_allele_is_A","predicate"=function(ctx) {"A" %in% var
  list("desc"="any_ALT_allele_is_A","predicate"=function(ctx) {"A" %in%
  list("desc"="No_QUAL","predicate"=function(ctx) {!variant.has.qual(ctx)}
  list("desc"="variant_has_ID","predicate"=function(ctx) {variant.has.id(c
  list("desc"="variant_ID_match_rs1*'',"predicate"=function(ctx) {grepl(
  list("desc"="variant_has_INFO/AF_NFE","predicate"=function(ctx) {variant
  list("desc"="variant_has_INFO/AF_NFE_>1E-5","predicate"=function(ctx) {
  list("desc"="Missense_in_PLEKHN1(VEP)","predicate"=function(ctx) {
    # NO VEP annotation ?
    if(!variant.has.attribute(ctx,"CSQ")) return(FALSE);
    # get VEP annotation
    predictions <- variant.vep(ctx)
    # In SCN5A
    predictions <- predictions[which(predictions$SYMBOL=="PLEKHN1"),
    # Consequence must contain missense
    predictions <- predictions[grep("missense_variant",predictions$C
    nrow(predictions)>0
  })
)

# count the variant for each filter
for(flt in filters) {
  print(paste(basename(filename),"_filter:",flt[["desc"]],"_count:",count.

```



```
}
```

Output:

```
[1] "gnomad.exomes.r2.0.1.sites.bcf_filter:accept_all_count:50\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_filter:accept_none_count:0\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_filter:CHROM_is_1_count:50\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_filter:POS_is_even_count:24\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_filter:PASS_filter_count:48\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_filter:count(FILTER)>1_count:2\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_filter:FILTER_contains_SEGDUP_count:1\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_filter:SNP_count:47\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_filter:POS!=END_count:3\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_filter:not_diallelic_count:8\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_filter:REF_is_A_count:6\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_filter:any_allele_is_A_count:27\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_filter:any_ALT_allele_is_A_count:21\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_filter:No_QUAL_count:1\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_filter:variant_has_ID_count:34\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_filter:variant_ID_match_rs1*_count:2\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_filter:variant_has_INFO/AF_NFE_count:50\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_filter:variant_has_INFO/AF_NFE>1E-5_count:1\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_filter:Missense_in_PLEKHN1(VEP)_count:1"
```

## 2.11 Print a VEP table for a Variant

Code:

```
# load rbcf
library(rbcf)
# A vcf
filename <- "./data/gnomad.exomes.r2.0.1.sites.bcf"
# we don't need the index for this file
fp <- bcf.open(filename,FALSE)
# current variant
vc <- NULL
while(!is.null(vc<-bcf.next(fp))) {
  #find the first variant having an INFO/CSQ attribute
  if(variant.has.attribute(vc,"CSQ")) break;
}

if(!is.null(vc)) {
```

```

# get the VEP table for the variant
predictions<-variant.vep(vc)
}

# dispose the vcf reader
bcf.close(fp)
# show
predictions

```

Output:

```

[1] TRUE

```

	Allele	Consequence	IMPACT	SYMBOL	Gene
1	C	downstream_gene_variant	MODIFIER	KLHL17	ENSG00000187961
2	A	downstream_gene_variant	MODIFIER	KLHL17	ENSG00000187961
3	C	downstream_gene_variant	MODIFIER	C1orf170	ENSG00000187642
4	A	downstream_gene_variant	MODIFIER	C1orf170	ENSG00000187642
5	C	intron_variant	MODIFIER	PLEKHN1	ENSG00000187583
6	A	intron_variant	MODIFIER	PLEKHN1	ENSG00000187583
7	C	intron_variant	MODIFIER	PLEKHN1	ENSG00000187583
8	A	intron_variant	MODIFIER	PLEKHN1	ENSG00000187583
9	C	intron_variant	MODIFIER	PLEKHN1	ENSG00000187583
10	A	intron_variant	MODIFIER	PLEKHN1	ENSG00000187583
11	C	downstream_gene_variant	MODIFIER	C1orf170	ENSG00000187642
12	A	downstream_gene_variant	MODIFIER	C1orf170	ENSG00000187642
13	C	downstream_gene_variant	MODIFIER	C1orf170	ENSG00000187642
14	A	downstream_gene_variant	MODIFIER	C1orf170	ENSG00000187642
15	C	upstream_gene_variant	MODIFIER	PLEKHN1	ENSG00000187583
16	A	upstream_gene_variant	MODIFIER	PLEKHN1	ENSG00000187583
17	C	upstream_gene_variant	MODIFIER	PLEKHN1	ENSG00000187583
18	A	upstream_gene_variant	MODIFIER	PLEKHN1	ENSG00000187583

  

	Feature_type	Feature	BIOTYPE	EXON	INTRON
1	Transcript	ENST00000338591	protein_coding		
2	Transcript	ENST00000338591	protein_coding		
3	Transcript	ENST00000341290	protein_coding		
4	Transcript	ENST00000341290	protein_coding		
5	Transcript	ENST00000379407	protein_coding		2/14
6	Transcript	ENST00000379407	protein_coding		2/14
7	Transcript	ENST00000379409	protein_coding		2/14
8	Transcript	ENST00000379409	protein_coding		2/14
9	Transcript	ENST00000379410	protein_coding		2/15
10	Transcript	ENST00000379410	protein_coding		2/15

11	Transcript	ENST00000433179	protein_coding
12	Transcript	ENST00000433179	protein_coding
13	Transcript	ENST00000479361	retained_intron
14	Transcript	ENST00000479361	retained_intron
15	Transcript	ENST00000480267	retained_intron
16	Transcript	ENST00000480267	retained_intron
17	Transcript	ENST00000491024	protein_coding
18	Transcript	ENST00000491024	protein_coding

HGVSg HGVSs cDNA\_position CDS\_position

```
5  ENST00000379407.3:c.184-51G>C
6  ENST00000379407.3:c.184-51G>A
7  ENST00000379409.2:c.184-51G>C
8  ENST00000379409.2:c.184-51G>A
9  ENST00000379410.3:c.184-51G>C
10 ENST00000379410.3:c.184-51G>A
```

14					rs540662886	2	4979
15					rs540662886	1	649
16					rs540662886	2	649
17					rs540662886	1	3286
18					rs540662886	2	3286
	STRAND	FLAGS	VARIANT_CLASS	MINIMISED	SYMBOL_SOURCE	HGNC_ID	CANONICAL
1	1		SNV		HGNC	24023	YES
2	1		SNV		HGNC	24023	YES
3	-1		SNV		HGNC	28208	
4	-1		SNV		HGNC	28208	
5	1		SNV		HGNC	25284	
6	1		SNV		HGNC	25284	
7	1		SNV		HGNC	25284	
8	1		SNV		HGNC	25284	
9	1		SNV		HGNC	25284	YES
10	1		SNV		HGNC	25284	YES
11	-1		SNV		HGNC	28208	YES
12	-1		SNV		HGNC	28208	YES
13	-1		SNV		HGNC	28208	
14	-1		SNV		HGNC	28208	
15	1		SNV		HGNC	25284	
16	1		SNV		HGNC	25284	
17	1	cds_start_NF	SNV		HGNC	25284	
18	1	cds_start_NF	SNV		HGNC	25284	
	TSL	APPRIS	CCDS	ENSP	SWISSPROT	TREMBL	UNIPARC
1			CCDS30550.1	ENSP00000343930	Q6TDP4	Q0VGE6&B3KXL7	UPI00001DFBF0
2			CCDS30550.1	ENSP00000343930	Q6TDP4	Q0VGE6&B3KXL7	UPI00001DFBF0
3				ENSP00000343864			UPI000022DAF4
4				ENSP00000343864			UPI000022DAF4
5			CCDS53256.1	ENSP00000368717	Q494U1	J3KSM5	UPI00005764FF
6			CCDS53256.1	ENSP00000368717	Q494U1	J3KSM5	UPI00005764FF
7				ENSP00000368719	Q494U1	J3KSM5	UPI0000D61E06
8				ENSP00000368719	Q494U1	J3KSM5	UPI0000D61E06
9			CCDS4.1	ENSP00000368720	Q494U1	J3KSM5	UPI00001416D8
10			CCDS4.1	ENSP00000368720	Q494U1	J3KSM5	UPI00001416D8
11				ENSP00000414022	Q5SV97		UPI0000418FB0
12				ENSP00000414022	Q5SV97		UPI0000418FB0
13							
14							
15							
16							

17	ENSP00000462558						J3KSM5	UPI000268AE1F
18	ENSP00000462558						J3KSM5	UPI000268AE1F
	GENE_PHENO	SIFT	PolyPhen	DOMAINS	HGVS_OFFSET	GMAF	AFR_MAF	AMR_MAF
1						C:0.0008	C:0	C:0
2						C:0.0008	C:0	C:0
3						C:0.0008	C:0	C:0
4						C:0.0008	C:0	C:0
5						C:0.0008	C:0	C:0
6						C:0.0008	C:0	C:0
7						C:0.0008	C:0	C:0
8						C:0.0008	C:0	C:0
9						C:0.0008	C:0	C:0
10						C:0.0008	C:0	C:0
11						C:0.0008	C:0	C:0
12						C:0.0008	C:0	C:0
13						C:0.0008	C:0	C:0
14						C:0.0008	C:0	C:0
15						C:0.0008	C:0	C:0
16						C:0.0008	C:0	C:0
17						C:0.0008	C:0	C:0
18						C:0.0008	C:0	C:0
	EAS_MAF	EUR_MAF	SAS_MAF	AA_MAF	EA_MAF	ExAC_MAF	ExAC_Adj_MAF	ExAC_AFR_MAF
1	C:0	C:0.004	C:0	C:0			C:0	C:2.146e-04
2	C:0	C:0.004	C:0	C:0			C:0	C:2.146e-04
3	C:0	C:0.004	C:0	C:0			C:0	C:2.146e-04
4	C:0	C:0.004	C:0	C:0			C:0	C:2.146e-04
5	C:0	C:0.004	C:0	C:0			C:0	C:2.146e-04
6	C:0	C:0.004	C:0	C:0			C:0	C:2.146e-04
7	C:0	C:0.004	C:0	C:0			C:0	C:2.146e-04
8	C:0	C:0.004	C:0	C:0			C:0	C:2.146e-04
9	C:0	C:0.004	C:0	C:0			C:0	C:2.146e-04
10	C:0	C:0.004	C:0	C:0			C:0	C:2.146e-04
11	C:0	C:0.004	C:0	C:0			C:0	C:2.146e-04
12	C:0	C:0.004	C:0	C:0			C:0	C:2.146e-04
13	C:0	C:0.004	C:0	C:0			C:0	C:2.146e-04
14	C:0	C:0.004	C:0	C:0			C:0	C:2.146e-04
15	C:0	C:0.004	C:0	C:0			C:0	C:2.146e-04
16	C:0	C:0.004	C:0	C:0			C:0	C:2.146e-04
17	C:0	C:0.004	C:0	C:0			C:0	C:2.146e-04
18	C:0	C:0.004	C:0	C:0			C:0	C:2.146e-04
	ExAC_AMR_MAF	ExAC_EAS_MAF	ExAC_FIN_MAF	ExAC_NFE_MAF	ExAC_OTH_MAF			

1	C:0	C:0.0002281	C:0.002986	C:0	C:1.606e-05			
2	C:0	C:0.0002281	C:0.002986	C:0	C:1.606e-05			
3	C:0	C:0.0002281	C:0.002986	C:0	C:1.606e-05			
4	C:0	C:0.0002281	C:0.002986	C:0	C:1.606e-05			
5	C:0	C:0.0002281	C:0.002986	C:0	C:1.606e-05			
6	C:0	C:0.0002281	C:0.002986	C:0	C:1.606e-05			
7	C:0	C:0.0002281	C:0.002986	C:0	C:1.606e-05			
8	C:0	C:0.0002281	C:0.002986	C:0	C:1.606e-05			
9	C:0	C:0.0002281	C:0.002986	C:0	C:1.606e-05			
10	C:0	C:0.0002281	C:0.002986	C:0	C:1.606e-05			
11	C:0	C:0.0002281	C:0.002986	C:0	C:1.606e-05			
12	C:0	C:0.0002281	C:0.002986	C:0	C:1.606e-05			
13	C:0	C:0.0002281	C:0.002986	C:0	C:1.606e-05			
14	C:0	C:0.0002281	C:0.002986	C:0	C:1.606e-05			
15	C:0	C:0.0002281	C:0.002986	C:0	C:1.606e-05			
16	C:0	C:0.0002281	C:0.002986	C:0	C:1.606e-05			
17	C:0	C:0.0002281	C:0.002986	C:0	C:1.606e-05			
18	C:0	C:0.0002281	C:0.002986	C:0	C:1.606e-05			
	ExAC_SAS_MAF	CLIN_SIG	SOMATIC	PHENO	PUBMED	MOTIF_NAME	MOTIF_POS	HIGH_INF_POS
1	C:0							
2	C:0							
3	C:0							
4	C:0							
5	C:0							
6	C:0							
7	C:0							
8	C:0							
9	C:0							
10	C:0							
11	C:0							
12	C:0							
13	C:0							
14	C:0							
15	C:0							
16	C:0							
17	C:0							
18	C:0							
	MOTIF_SCORE_CHANGE	LoF	LoF_filter	LoF_flags	LoF_info"			
1								
2								
3								

```
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
```

## 2.12 Print a SNPEFF table for a Variant

Code:

```
# load rbcf
library(rbcf)
# A vcf
filename <- "./data/rotavirus_rf.ann.vcf.gz"
# we don't need the index for this file
fp <- bcf.open(filename,FALSE)
# current variant
vc <- NULL
while(!is.null(vc<-bcf.next(fp))) {
  #find the first variant having an INFO/ANN attribute
  if(variant.has.attribute(vc,"ANN")) break;
}
if(!is.null(vc)) {
  # get SNPEFF table
  predictions<-variant.snpeff(vc)
}
# dispose the vcf reader
bcf.close(fp)
# show
predictions
```

Output:

```
[1] TRUE
  Allele      Annotation Annotation_Impact   Gene_Name   Gene_ID
1      C missense_variant             MODERATE Gene_18_3284 Gene_18_3284
  Feature_Type Feature_ID Transcript_BioType Rank   HGVS.c      HGVS.p
1   transcript AAA47319.1    protein_coding  1/1 c.952A>C p.Lys318Gln
  cDNA.pos / cDNA.length CDS.pos / CDS.length AA.pos / AA.length Distance
1              952/3267              952/3267              318/1088
  ERRORS / WARNINGS / INFO'"
1
```

## 2.13 Query the indexed vcf using intervals

Code:

```
# load rbcf
library(rbcf)

# create a function counting variants in a VCF, in some intervals
count.variants<-function(filename,intervals) {
  # open the indexed VCF
  fp <- bcf.open(filename)
  # loop over the intervals
  for(interval in intervals) {
    # try query the interval
    if(bcf.query(fp,interval)) {
      # number of variants
      n<-0
      # loop while we can read a variant
      while(!is.null(vc<-bcf.next(fp))) {
        # increment the count
        n<-n+1
      }
      print(paste("Number of variants in ",basename(filename),
        })
    # query failed
    else {
      print(paste("Cannot query ",basename(filename),"/'",interval))
    }
  }
  # dispose the vcf reader
  bcf.close(fp)
}
```



```

}

some_intervals <-c("", "RF03", "RF03:2000-3000", "1:1-10000000", "chr1")
count.variants("./data/rotavirus_rf.02.vcf.gz", some_intervals)
count.variants("./data/1000G.ALL.2of4intersection.20100804.genotypes.bcf", some_i

# another way to query is set collect=TRUE to return a vector of variant
fp <- bcf.open("./data/rotavirus_rf.02.vcf.gz")
print(paste("Number_of_variants_using_collect:", length(bcf.query(fp, "RF03", colle
bcf.close(fp)

```

**Output:**

```

[1] "Cannot_query_rotavirus_rf.02.vcf.gz/' '"
[1] "Number_of_variants_in_rotavirus_rf.02.vcf.gz/'RF03':8"
[1] "Number_of_variants_in_rotavirus_rf.02.vcf.gz/'RF03:2000-3000':4"
[1] "Cannot_query_rotavirus_rf.02.vcf.gz/'1:1-10000000'"
[1] "Cannot_query_rotavirus_rf.02.vcf.gz/'chr1'"
[1] TRUE
[1] "Cannot_query_1000G.ALL.2of4intersection.20100804.genotypes.bcf/' '"
[1] "Cannot_query_1000G.ALL.2of4intersection.20100804.genotypes.bcf/'RF03'"
[1] "Cannot_query_1000G.ALL.2of4intersection.20100804.genotypes.bcf/'RF03:2000-3"
[1] "Number_of_variants_in_1000G.ALL.2of4intersection.20100804.genotypes.bcf/'1:"
[1] "Cannot_query_1000G.ALL.2of4intersection.20100804.genotypes.bcf/'chr1'"
[1] TRUE
[1] "Number_of_variants_using_collect:8"
[1] TRUE

```

## 2.14 Attribute in INFO

**Code:**

```

# load rbcf
library(rbcf)

# find given variant
find.variant<-function(fp, contig, pos) {
  if(!bcf.query(fp, paste(contig, ":", pos, "-", pos, sep=""))) return(NULL)
  # loop while we can read a variant
  while(!is.null(vc<-bcf.next(fp))) {
    return(vc)
  }
}

```

```

        return(NULL)
    }
    filename<-"./data/gnomad.exomes.r2.0.1.sites.bcf"
    # open the VCF with index
    fp <- bcf.open(filename)
    ctx <-find.variant(fp,"1",905608)
    stopifnot(variant.has.attribute(ctx,"CSQ"))
    print(paste("CSQ(no_split)",variant.string.attribute(ctx,"CSQ",split=FALSE)))
    print(paste("CSQ(split)",variant.string.attribute(ctx,"CSQ")))
    stopifnot(variant.has.attribute(ctx,"AN_POPMAX"))
    print(paste("AN_POPMAX:",variant.int.attribute(ctx,"AN_POPMAX")))
    stopifnot(variant.has.attribute(ctx,"AF_POPMAX"))
    print(paste("AF_POPMAX:",variant.float.attribute(ctx,"AF_POPMAX")))
    print(paste("flag:VQSR_NEGATIVE_TRAIN_SITE:",variant.flag.attribute(ctx,"VQSR_NE
    # dispose the vcf reader
    bcf.close(fp)

```

### Output:

```

[1] "CSQ(no_split)_T|downstream_gene_variant|MODIFIER|KLHL17|ENSG00000187961|Tr
[1] "CSQ(split) T|downstream_gene_variant|MODIFIER|KLHL17|ENSG00000187961|Tran
[2] "CSQ(split)_A|downstream_gene_variant|MODIFIER|KLHL17|ENSG00000187961|Tran
[3] "CSQ(split) T|downstream_gene_variant|MODIFIER|C1orf170|ENSG00000187642|Tr
[4] "CSQ(split)_A|downstream_gene_variant|MODIFIER|C1orf170|ENSG00000187642|Tr
[5] "CSQ(split) T|intron_variant|MODIFIER|PLEKHN1|ENSG00000187583|Transcript|E
[6] "CSQ(split)_A|intron_variant|MODIFIER|PLEKHN1|ENSG00000187583|Transcript|E
[7] "CSQ(split) T|intron_variant|MODIFIER|PLEKHN1|ENSG00000187583|Transcript|E
[8] "CSQ(split)_A|intron_variant|MODIFIER|PLEKHN1|ENSG00000187583|Transcript|E
[9] "CSQ(split) T|intron_variant|MODIFIER|PLEKHN1|ENSG00000187583|Transcript|E
[10] "CSQ(split)_A|intron_variant|MODIFIER|PLEKHN1|ENSG00000187583|Transcript|E
[11] "CSQ(split) T|downstream_gene_variant|MODIFIER|C1orf170|ENSG00000187642|Tr
[12] "CSQ(split)_A|downstream_gene_variant|MODIFIER|C1orf170|ENSG00000187642|Tr
[13] "CSQ(split) T|downstream_gene_variant|MODIFIER|C1orf170|ENSG00000187642|Tr
[14] "CSQ(split)_A|downstream_gene_variant|MODIFIER|C1orf170|ENSG00000187642|Tr
[15] "CSQ(split) T|upstream_gene_variant|MODIFIER|PLEKHN1|ENSG00000187583|Trans
[16] "CSQ(split)_A|upstream_gene_variant|MODIFIER|PLEKHN1|ENSG00000187583|Trans
[17] "CSQ(split) T|upstream_gene_variant|MODIFIER|PLEKHN1|ENSG00000187583|Trans
[18] "CSQ(split)_A|upstream_gene_variant|MODIFIER|PLEKHN1|ENSG00000187583|Trans
[1] "AN_POPMAX: 106408" "AN_POPMAX: 106408"
[1] "AF_POPMAX: 1.87955993169453e-05" "AF_POPMAX: 9.39778965403093e-06"
[1] "flag:VQSR_NEGATIVE_TRAIN_SITE: FALSE"
[1] TRUE

```

## 2.15 Working with Genotypes

Code:

```
# load rbcf
library(rbcf)

# find given variant
find.variant<-function(fp,contig,pos) {
  if(!bcf.query(fp,paste(contig,":",pos,"-",pos,sep=""))) return(NULL)
  # loop while we can read a variant
  while(!is.null(vc<-bcf.next(fp))) {
    return(vc)
  }
  return(NULL)
}

filename<-"/data/1000G.ALL.2of4intersection.20100804.genotypes.bcf"
# open the VCF with index
fp <- bcf.open(filename)
# find a variant
ctx <-find.variant(fp,"1",10583)
print(paste("Number_of_genotypes_",variant.nsamples(ctx)))
# get 10-th genotype
gt<-variant.genotype(ctx,10)
print(paste("sample_",genotype.sample(gt)))
# get genotype by name
gt<-variant.genotype(ctx,"NA18997")
print(paste("sample_",genotype.sample(gt)))
print(paste("alleles_",genotype.alleles.idx0(gt)))
print(paste("genotype_ploidy_",genotype.ploidy(gt)))
print(paste("genotype_is_hom_ref_",genotype.homref(gt)))
print(paste("genotype_is_het_",genotype.het(gt)))
print(paste("genotype_is_het-non-ref_",genotype.hetnonref(gt)))
print(paste("genotype_is_phased_",genotype.phased(gt)))
print(paste("genotype_is_nocall_",genotype.nocall(gt)))
print(paste("genotype_FORMAT/OG_",genotype.string.attribute(gt,"OG")))
print(paste("genotype_FORMAT/GQ_",genotype.int.attribute(gt,"GQ")))# hum spec
print(paste("genotype_has_GQ_",genotype.has.gq(gt)))
print(paste("genotype_GQ_",genotype.gq(gt)))
print(paste("genotype_has_DP_",genotype.has.dp(gt)))
print(paste("genotype_DP_",genotype.int.attribute(gt,"DP")))
print(paste("genotype_DP_",genotype.dp(gt)))
```

```

print(paste("genotype_has_PL_", genotype.has.pl(gt)))
print(paste("genotype_PL_", genotype.pl(gt)))
print(paste("genotype_has_AD_", genotype.has.ad(gt)))
print(paste("genotype_AD_", genotype.ad(gt)))

# dispose the vcf reader
bcf.close(fp)

```

**Output:**

```

[1] "Number_of_genotypes_629"
[1] "sample_HG00120"
[1] "sample_NA18997"
[1] "alleles_0" "alleles_1"
[1] "genotype_ploidy_2"
[1] "genotype_is_hom_ref_FALSE"
[1] "genotype_is_het_TRUE"
[1] "genotype_is_het-non-ref_FALSE"
[1] "genotype_is_phased_TRUE"
[1] "genotype_is_no_call_FALSE"
[1] "genotype_FORMAT/OG_1/1"
[1] "genotype_FORMAT/GQ_"
[1] "genotype_has_GQ_FALSE"
[1] "genotype_GQ_-1"
[1] "genotype_has_DP_TRUE"
[1] "genotype_DP_1"
[1] "genotype_DP_1"
[1] "genotype_has_PL_FALSE"
[1] "genotype_PL_"
[1] "genotype_has_AD_TRUE"
[1] "genotype_AD_4" "genotype_AD_1"
[1] TRUE

```

## 2.16 Writing variants to a new VCF/BCF file

**Code:**

```

# load rbcf
library(rbcf)
# vcf input filename
filenamein = "./data/rotavirus_rf.01.vcf"
# output vcf filename. "-" is standard output

```

```

filenameout = "-"

fp <- bcf.open(filenamein,FALSE)
# create a new VCF writer using the header from 'fp'
out <- bcf.new.writer(fp,filenameout);
# loop while we can read a variant
while(!is.null(vc<-bcf.next(fp))) {
  # only write POS%10==0
  if(variant.pos(vc)%%10==0) {
    # write variant
    bcf.write.variant(out,vc);
  }
}
# dispose the vcf reader
bcf.close(fp)
# dispose the vcf rwriter
bcf.close(out);

```

Output:

```

[1] TRUE
##fileformat=VCFv4.2
##FILTER=<ID=PASS,Description="All filters passed">
##samtoolsVersion=1.3.1+htslib-1.3.1
##samtoolsCommand=samtools mpileup -Ou -f rotavirus_rf.fa S1.bam S2.bam S3.bam
##reference=file://rotavirus_rf.fa
##contig=<ID=RF01,length=3302>
##contig=<ID=RF02,length=2687>
##contig=<ID=RF03,length=2592>
##contig=<ID=RF04,length=2362>
##contig=<ID=RF05,length=1579>
##contig=<ID=RF06,length=1356>
##contig=<ID=RF07,length=1074>
##contig=<ID=RF08,length=1059>
##contig=<ID=RF09,length=1062>
##contig=<ID=RF10,length=751>
##contig=<ID=RF11,length=666>
##ALT=<ID=*,Description="Represents allele(s) other than observed.">
##INFO=<ID=INDEL,Number=0,Type=Flag,Description="Indicates that the variant is
##INFO=<ID=IDV,Number=1,Type=Integer,Description="Maximum number of reads supp
##INFO=<ID=IMF,Number=1,Type=Float,Description="Maximum fraction of reads supp
##INFO=<ID=DP,Number=1,Type=Integer,Description="Raw read depth">

```

```

##INFO=<ID=VDB,Number=1,Type=Float,Description="Variant Distance Bias for filt
##INFO=<ID=RPB,Number=1,Type=Float,Description="Mann-Whitney U test of Read Po
##INFO=<ID=MQB,Number=1,Type=Float,Description="Mann-Whitney U test of Mapping
##INFO=<ID=BQB,Number=1,Type=Float,Description="Mann-Whitney U test of Base Qu
##INFO=<ID=MQSB,Number=1,Type=Float,Description="Mann-Whitney U test of Mapping
##INFO=<ID=SGB,Number=1,Type=Float,Description="Segregation based metric.">
##INFO=<ID=MQOF,Number=1,Type=Float,Description="Fraction of MQO reads (smaller
##FORMAT=<ID=PL,Number=G,Type=Integer,Description="List of Phred-scaled genotyp
##FORMAT=<ID=GT,Number=1,Type=String,Description="Genotype">
##INFO=<ID=ICB,Number=1,Type=Float,Description="Inbreeding Coefficient Binomia
##INFO=<ID=HOB,Number=1,Type=Float,Description="Bias in the number of HOMs num
##INFO=<ID=AC,Number=A,Type=Integer,Description="Allele count in genotypes for
##INFO=<ID=AN,Number=1,Type=Integer,Description="Total number of alleles in ca
##INFO=<ID=DP4,Number=4,Type=Integer,Description="Number of high-quality ref-f
##INFO=<ID=MQ,Number=1,Type=Integer,Description="Average mapping quality">
##bcftools_callVersion=1.3-10-g820e1d6+htslib-1.2.1-267-g87141ea
##bcftools_callCommand=call -vm -Oz -o rotavirus_rf.vcf.gz -
##bcftools_viewVersion=1.10-6-g2782d9f+htslib-1.2.1-1336-g7c16b56-dirty
##bcftools_viewCommand=view /home/lindenb/src/jvarkit/src/test/resources/rotavi
#CHROM POS ID REF ALT QUAL FILTER INFO FORMAT S1
RF01 970 . A C 48.6696 . DP=36;VDB=0.693968;SGB=1
RF03 2150 . T A 6.90687 . DP=37;VDB=0.557348;SGB=-
RF04 1900 . A C 36.8224 . DP=39;VDB=0.706942;SGB=7
RF04 1920 . A T 42.014 . DP=39;VDB=0.966939;SGB=0

```