RBcf: An VCF API for R.

Pierre Lindenbaum / @yokofakun/ Institut du Thorax . Nantes.

April 27, 2020

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

```
# 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.")</pre>
```

```
[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</pre>
```

```
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
                1 Float
      MQSB
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 v
IMF
                                                                                                                                              "Maximum_fraction_of_reads_
DP
               "Variant_Distance_Bias_for_filtering_splice-site_artefacts_in_RNA-seq_data
\verb|"Mann-Whitney|| U_{\sqcup} test_{\sqcup} of_{\sqcup} Mapping_{\sqcup} Quality_{\sqcup} Bias
MQB
BQB______Mann-Whitney U test of Base Quality Bias
                                                                     \verb|"Mann-Whitney|| U_{\sqcup} test_{\sqcup} of_{\sqcup} \verb|Mapping|| Quality_{\sqcup} vs_{\sqcup} Strand_{\sqcup} Bias
MQOF
                                                                                                                                                         "Fraction of MQ0 reads to model in the mode
ICB " Inbreeding Coefficient Binomial test
                                                                                                                         "Bias_in_the_number_of_HOMs_number_
"Total_{\square}number_{\square}of_{\square}alleles_{\square}
MQ
                                                                                                                                                                                                          "Aver
[1] LTRUE
```

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)</pre>
```

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

```
# 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)</pre>
```

```
TD
PASS
               PASS
AC0
                ACO
InbreedingCoeff InbreedingCoeff
LCR
RF
                RF
              SEGDUP
SEGDUP
PASS
AC0
         "Allele_Count_is_zero_(i.e._no_high-confidence_genotype_(Q_{\square}>=_2
LCR
                             "Failed<sub>□</sub>random<sub>□</sub>forests<sub>□</sub>filte
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)</pre>
```

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

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)</pre>
```

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

```
# 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)</pre>
```

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

2.9 Scanning the variants

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

```
./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

```
# load rbcf
library(rbcf)
# create a function counting variants in a VCF
count.variants<-function(filename,predicate) {</pre>
        # we don't need the index for this file
        fp <- bcf.open(filename,FALSE)</pre>
        # number of variants
        n<-0
        # loop while we can read a variant
        while(!is.null(vc<-bcf.next(fp))) {</pre>
                # 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(ct
        list("desc"="POS_{\sqcup}is_{\sqcup}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<sub>□</sub>is<sub>□</sub>'A'", "predicate"=function(ctx) {variant.reference(ct
        list("desc"="anyualleleuisu'A'", "predicate"=function(ctx) {"A" %in% var
        list("desc"="any LALT allele is 'A'", "predicate"=function(ctx) {"A" %in%
        list("desc"="No_QUAL", "predicate"=function(ctx) {!variant.has.qual(ctx)}
        list("desc"="variant<sub>\ull</sub>has<sub>\ull</sub>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)</pre>
                 # In SCN5A
                 predictions <- predictions[which(predictions$SYMBOL=="PLEKHN1"),</pre>
                 # Consequence must contain missense
                 predictions <- predictions[grep("missense_variant",predictions$C</pre>
                 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_{\sqcup \sqcup}filter:_{\sqcup}accept_{\sqcup}all_{\sqcup \sqcup}count:_{\sqcup}50_{\sqcup}\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf,,filter:,accept,none,,count:,0,\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_u_filter:_CHROM_is_'1'_u_count:_50_\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_{\sqcup \sqcup}filter:_{\sqcup}POS_{\sqcup}is_{\sqcup}even_{\sqcup \sqcup}count:_{\sqcup}24_{\sqcup}\n"
[1] "gnomad.exomes.r2.0.1.sites.bcfuufilter:uPASSufilteruucount:u48u\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_\( \sigma filter:\( \sigma count(FILTER) > 1 \) \( \sigma count:\( \sigma 2 \) \\ n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_{\sqcup\sqcup}filter:_{\sqcup}FILTER_{\sqcup}contains_{\sqcup}SEGDUP_{\sqcup\sqcup}count:_{\sqcup}1_{\sqcup}\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_u_filter:_not_diallelic_u_count:_\8_\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_ufilter:uREF_is_'A'u_count:u6u\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_{\sqcup \sqcup}filter:_{\sqcup}any_{\sqcup}allele_{\sqcup}is_{\sqcup}'A'_{\sqcup \sqcup}count:_{\sqcup}27_{\sqcup}\n"
[1] "gnomad.exomes.r2.0.1.sites.bcf_{\sqcup \sqcup}filter:_{\sqcup}any_{\sqcup}ALT_{\sqcup}allele_{\sqcup}is_{\sqcup}'A'_{\sqcup \sqcup}count:_{\sqcup}21_{\sqcup}\n
[1] "gnomad.exomes.r2.0.1.sites.bcf_{\sqcup \sqcup}filter:_{\sqcup}No_{\sqcup}QUAL_{\sqcup \sqcup}count:_{\sqcup}1_{\sqcup}\n"
[1] "gnomad.exomes.r2.0.1.sites.bcfuufilter:uvariantuhasuIDuucount:u34u\n"
[1] "gnomad.exomes.r2.0.1.sites.bcfullfilter:uvariantuIDumatchu'rs1*'uuucount:u2u
[1] "gnomad.exomes.r2.0.1.sites.bcfullfilter:uvariantuhasuINFO/AF_NFEulcount:u50u
[1] "gnomad.exomes.r2.0.1.sites.bcfullfilter:uvariantuhasuINF0/AF_NFEL>u1E-5ulcou
[1] "gnomad.exomes.r2.0.1.sites.bcf_{\sqcup\sqcup}filter:_{\sqcup}Missense_{\sqcup}in_{\sqcup}PLEKHN_{\sqcup\sqcup}(VEP)_{\sqcup\sqcup}count:_{\sqcup}1
```

2.11 Print a VEP table for a Variant

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

# dispose the vcf reader
bcf.close(fp)
# show
predictions</pre>
```

[1]	TRUE				
	Allele	Consequence	IMPACT	SYMBOL	Gene
1	C downs	tream_gene_variant	MODIFIER	KLHL17	ENSG00000187961
2	A downs	tream_gene_variant	MODIFIER	KLHL17	ENSG00000187961
3	C downs	tream_gene_variant	MODIFIER	C1orf170	ENSG00000187642
4	A downs	tream_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	С				ENSG00000187583
10	A				ENSG00000187583
11	C downs	tream_gene_variant	MODIFIER	C1orf170	ENSG00000187642
12		tream_gene_variant			
13	C downs	tream_gene_variant	MODIFIER	C1orf170	ENSG00000187642
14	A downs	tream_gene_variant	MODIFIER	C1orf170	ENSG00000187642
15	_	tream_gene_variant			
16	_	tream_gene_variant			
17	C ups	tream_gene_variant	MODIFIER	PLEKHN1	ENSG00000187583
18	A ups	tream_gene_variant			
	Feature_type	Feature	BIO	OTYPE EXO	N INTRON
1	-	ENST00000338591 I	protein_c	oding	
2	-	•	protein_c	_	
3	-		protein_c		
4	Transcript	ENST00000341290 I	protein_c	oding	
5	-	•	protein_c	oding	2/14
6	Transcript	ENST00000379407 I	protein_c	oding	2/14
7	Transcript	ENST00000379409 I	protein_c	oding	2/14
8	Transcript	ENST00000379409 I	protein_c	oding	2/14
9	Transcript	ENST00000379410 I	protein_c	oding	2/15
10	Transcript	ENST00000379410 I	protein_c	oding	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
                           HGVSc HGVSp cDNA_position CDS_position
1
2
3
4
5
  ENST00000379407.3:c.184-51G>C
6
  ENST00000379407.3:c.184-51G>A
  ENST00000379409.2:c.184-51G>C
8 ENST00000379409.2:c.184-51G>A
  ENST00000379410.3:c.184-51G>C
10 ENST00000379410.3:c.184-51G>A
11
12
13
14
15
16
17
18
   Protein_position Amino_acids Codons Existing_variation ALLELE_NUM DISTANCE
1
                                               rs540662886
                                                                     1
                                                                           4511
2
                                               rs540662886
                                                                     2
                                                                           4511
3
                                               rs540662886
                                                                     1
                                                                           4978
4
                                               rs540662886
                                                                     2
                                                                           4978
5
                                               rs540662886
                                                                     1
6
                                                                     2
                                               rs540662886
7
                                               rs540662886
                                                                     1
8
                                                                     2
                                               rs540662886
                                               rs540662886
                                                                     1
10
                                               rs540662886
                                                                     2
11
                                               rs540662886
                                                                     1
                                                                           4973
12
                                                                     2
                                               rs540662886
                                                                           4973
13
                                               rs540662886
                                                                     1
                                                                           4979
```

4.4				E 4.0	00000		0 4070
14					662886		2 4979
15				rs540662886			1 649
16		rs540662886					2 649
17					rs540662886		1 3286
18					662886		2 3286
	STRAND	FLAGS V	VARIANT_CLASS MIN	IMISED SYMB			
1	1		SNV		HGNC	2402	
2	1		SNV		HGNC	2402	
3	-1		SNV		HGNC	2820)8
4	-1		SNV		HGNC	2820	08
5	1		SNV		HGNC	2528	34
6	1		SNV		HGNC	2528	34
7	1		SNV		HGNC	2528	34
8	1		SNV		HGNC	2528	34
9	1		SNV		HGNC	2528	34 YES
10	1		SNV		HGNC	2528	34 YES
11	-1		SNV		HGNC	2820	98 YES
12	-1		SNV		HGNC	2820	98 YES
13	-1		SNV		HGNC	2820)8
14	-1		SNV		HGNC	2820)8
15	1		SNV		HGNC	2528	34
16	1		SNV		HGNC	2528	34
17	1	cds_start_NF	SNV		HGNC	2528	34
18	1	cds_start_NF	SNV		HGNC	2528	34
	TSL APP	RIS CCI	OS ENS	P SWISSPROT	TR	EMBL	UNIPARC
1		CCDS30550	.1 ENSP0000034393	O Q6TDP4	QOVGE6&B3	KXL7 U	JPI00001DFBF0
2		CCDS30550	.1 ENSP0000034393				JPI00001DFBF0
3			ENSP0000034386				JPI000022DAF4
4			ENSP0000034386	4		Ţ	JPI000022DAF4
5		CCDS53256	.1 ENSP0000036871		J3	KSM5 U	PI00005764FF
6		CCDS53256	.1 ENSP0000036871	•			PI00005764FF
7			ENSP0000036871	•			JPI0000D61E06
8			ENSP0000036871	•			JPI0000D61E06
9		CCDS4	.1 ENSP0000036872	•			PI00001416D8
10			.1 ENSP0000036872	•			PI00001416D8
11		00202	ENSP0000041402	•			JPI0000418FB0
12			ENSP0000041402	•			JPI0000418FB0
13			2 0000011102	_ 400701			- 10000 1101 00
14							
15							
16							

```
17
                            ENSP00000462558
                                                                 J3KSM5 UPI000268AE1F
18
                                                                 J3KSM5 UPI000268AE1F
                            ENSP00000462558
   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
4
                                                     C:0.0008
                                                                   C:0
                                                                            C:0
                                                                            C:0
                                                     C:0.0008
                                                                   C:0
5
                                                                   C:0
                                                     C:0.0008
                                                                            C:0
6
                                                     C:0.0008
                                                                   C:0
                                                                            C:0
                                                     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
                                                                            C:0
                                                     C:0.0008
12
                                                     C:0.0008
                                                                   C:0
                                                                            C:0
13
                                                     C:0.0008
                                                                   C:0
                                                                            C:0
14
                                                                   C:0
                                                                            C:0
                                                     C:0.0008
15
                                                                   C:0
                                                                            C:0
                                                     C:0.0008
16
                                                     C:0.0008
                                                                   C:0
                                                                            C:0
17
                                                                   C:0
                                                                            C:0
                                                     C:0.0008
18
                                                                            C:0
                                                     C:0.0008
                                                                   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:2.146e-04
                         C:0
2
       C:0 C:0.004
                                                                      C:2.146e-04
                         C:0
                                 C:0
                                                                 C:0
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:2.146e-04
                         C:0
                                                                 C:0
5
       C:0 C:0.004
                                 C:0
                                                                 C:0
                                                                      C:2.146e-04
                         C:0
6
                                                                      C:2.146e-04
       C:0 C:0.004
                         C:0
                                 C:0
                                                                 C:0
       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:2.146e-04
                         C:0
9
                                                                      C:2.146e-04
       C:0 C:0.004
                         C:0
                                 C:0
                                                                 C:0
                         C:0
10
       C:0 C:0.004
                                 C:0
                                                                 C:0
                                                                      C:2.146e-04
                                                                      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
13
       C:0 C:0.004
                                                                 C:0
                                                                      C:2.146e-04
                         C:0
                                 C:0
14
       C:0 C:0.004
                         C:0
                                 C:0
                                                                 C:0
                                                                      C:2.146e-04
15
       C:0 C:0.004
                                                                      C:2.146e-04
                         C:0
                                 C:0
                                                                 C:0
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:2.146e-04
                         C:0
                                                                 C:0
   ExaC_AMR_MAF ExaC_EAS_MAF ExaC_FIN_MAF ExaC_NFE_MAF ExaC_OTH_MAF
```

```
C:0
                  C:0.0002281
                                  C:0.002986
                                                        C:0
                                                             C:1.606e-05
2
3
             C:0
                  C:0.0002281
                                  C:0.002986
                                                        C:0
                                                             C:1.606e-05
                  C:0.0002281
                                  C:0.002986
                                                             C:1.606e-05
             C:0
                                                        C:0
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
             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
                                                             C:1.606e-05
15
             C:0
                  C:0.0002281
                                  C:0.002986
                                                        C:0
16
             C:0
                  C:0.0002281
                                  C:0.002986
                                                        C:0
                                                             C:1.606e-05
17
                                                             C:1.606e-05
             C:0
                  C:0.0002281
                                  C:0.002986
                                                        C:0
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
             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
```

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)</pre>
# current variant
vc <- NULL
while(!is.null(vc<-bcf.next(fp))) {</pre>
        #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)</pre>
# dispose the vcf reader
bcf.close(fp)
# show
predictions
```

```
[1] TRUE
 Allele
               Annotation Annotation_Impact
                                               Gene_Name
                                                               Gene_ID
      C missense_variant
                                   MODERATE Gene_18_3284 Gene_18_3284
 Feature_Type Feature_ID Transcript_BioType Rank
                                                    HGVS.c
                                                                HGVS.p
  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
                952/3267
                                     952/3267
                                                        318/1088
 ERRORS / WARNINGS / INFO'"
```

2.13 Query the indexed vcf using intervals

```
# load rbcf
library(rbcf)
# create a function counting variants in a VCF, in some intervals
count.variants<-function(filename,intervals) {</pre>
        # open the indexed VCF
        fp <- bcf.open(filename)</pre>
        # 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))) {</pre>
                                 # increment the count
                                 n < -n+1
                         print(paste("Number of variants in ", basename(filename),
                # query failed
                else {
                         print(paste("Cannot query ", basename(filename), "/',", inte
                }
        # 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")</pre>
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'"

2.14 Attribute in INFO

[1] "Number_of_variants_using_collect:_8"

Code:

[1] TRUE

[1] TRUE

```
# load rbcf
library(rbcf)

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

```
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)
```

```
[1] "CSQ(no_{\sqcup}split)_{\sqcup\sqcup}T|downstream\_gene\_variant|MODIFIER|KLHL17|ENSG00000187961|Trightspace | Trightspace | 
_{\sqcup}[1]_{\sqcup}"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] LTRUE
```

2.15 Working with Genotypes

```
# load rbcf
library(rbcf)
# find given variant
find.variant<-function(fp,contig,pos) {</pre>
         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))) {</pre>
                 return(vc)
        return(NULL)
filename <- "./data/1000G.ALL.2of4intersection.20100804.genotypes.bcf"
# open the VCF with index
fp <- bcf.open(filename)</pre>
# 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<sub>□</sub>",genotype.sample(gt)))
# get genotype by name
gt<-variant.genotype(ctx,"NA18997")</pre>
print(paste("sample<sub>□</sub>",genotype.sample(gt)))
print(paste("alleles<sub>□</sub>",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<sub>□</sub>is<sub>□</sub>het-non-ref<sub>□</sub>?<sub>□</sub>",genotype.hetnonref(gt)))
print(paste("genotype_is_phased_?_",genotype.phased(gt)))
print(paste("genotype_is_no_call_?, genotype.nocall(gt)))
print(paste("genotype<sub>□</sub>FORMAT/OG<sub>□</sub>?<sub>□</sub>",genotype.string.attribute(gt,"OG")))
print(paste("genotype_FORMAT/GQ_?_",genotype.int.attribute(gt,"GQ")))# hum spec
print(paste("genotype_{l}has_{l}GQ_{l}?_{l}",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)
```

```
[1] "Number of genotypes 629"
[1] "sample__HG00120"
[1] "sample__NA18997"
[1] "alleles___0" "alleles___1"
[1] "genotype_ploidy_?__2"
[1] "genotype_{\sqcup}is_{\sqcup}hom_{\sqcup}ref_{\sqcup}?_{\sqcup\sqcup}FALSE"
[1] "genotype_is_het_?__TRUE"
[1] "genotype_is_het-non-ref_?_LFALSE"
[1] "genotype_is_phased_?__TRUE"
[1] "genotype_is_no_call_?__FALSE"
[1] "genotype_FORMAT/OG_?__1/1"
[1] "genotype_FORMAT/GQ_?___"
[1] "genotype_{\sqcup}has_{\sqcup}GQ_{\sqcup}?_{\sqcup\sqcup}FALSE"
[1] "genotype_GQ__-1"
[1] "genotype_has_DP_?__TRUE"
[1] "genotype_DP_U_1"
[1] "genotype_DP_U_1"
[1] "genotype_has_PL_?__FALSE"
[1] "genotype_PL__"
[1] "genotype_has_AD_?_LLTRUE"
[1] "genotype_AD_U4" "genotype_AD_U1"
[1] TRUE
```

2.16 Writing variants to a new VCF/BCF file

```
# load rbcf
library(rbcf)
# vcf input filename
filenamein = "./data/rotavirus_rf.01.vcf"
# output vcf filename. "-" is standard 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>
##contiq=<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 support
##INFO=<ID=IMF, Number=1, Type=Float, Description="Maximum fraction of reads support
##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 Pos
##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 Mappin
##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 genoty"
##FORMAT=<ID=GT, Number=1, Type=Strinq, 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 number"
##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-fe
##INFO=<ID=MQ, Number=1, Type=Integer, Description="Average mapping quality">
\#\#bcftools\_callVersion=1.3-10-q820e1d6+htslib-1.2.1-267-q87141ea
##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/rotava
#CHROM POS
                 ID
                         REF
                                  ALT
                                          QUAL
                                                  FILTER INFO
                                                                    FORMAT S1
RF01
        970
                                 C
                                         48.6696 .
                                                          DP=36; VDB=0.693968; SGB=1
                         Α
RF03
                                                          DP=37; VDB=0.557348; SGB=-
        2150
                         Τ
                                 Α
                                         6.90687 .
RF04
        1900
                                 C
                                         36.8224 .
                                                          DP=39; VDB=0.706942; SGB=7
                         Α
RF04
                                 Τ
                                                          DP=39; VDB=0.966939; SGB=0
        1920
                         Α
                                         42.014
```