RBcf: An VCF API for R.

Pierre Lindenbaum / yokofakun/ Institut du Thorax . Nantes. ${\it April~23,~2020}$

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.

2 Examples

2.1 Htslib and Rbcf versions

Code:

```
# load the library
library(rbcf)
#print the version of the associated htslib
htslib.version()
#print the version of rbcf
rcbf.version()
```

Output:

```
> # load the library
> library(rbcf)
> #print the version of the associated htslib
> htslib.version()
[1] "1.10.2-dirty"
> #print the version of rbcf
> rcbf.version()
[1] "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("../tests/data/rotavirus_rf.01.vcf",FALSE)
# dispose the vcf reader
bcf.close(fp)</pre>
```

Output:

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

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("../tests/data/rotavirus_rf.01.vcf",FALSE)
info <- bcf.infos(fp)
# dispose the vcf reader
bcf.close(fp)
# print the table
info</pre>
```

Output:

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

```
[1] TRUE
 > # print the table
 > info
                                                  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
                                                                                                                    Float
                                                                                               1
BQB
                                           BQB
                                                                                               1
                                                                                                                  Float
MQSB
                                       MQSB
                                                                                               1
                                                                                                                  Float
SGB
                                            SGB
                                                                                               1
                                                                                                                  Float
MQOF
                                       MQOF
                                                                                               1
                                                                                                                   Float
 ICB
                                            ICB
                                                                                               1
                                                                                                                    Float
HOB
                                           HOB
                                                                                                                    Float
                                                                                               1
 AC
                                                 AC
                                                                                               A Integer
AN
                                                 AN
                                                                                               1 Integer
DP4
                                            DP4
                                                                                               4 Integer
MQ
                                                 MQ
                                                                                               1 Integer
 INDEL
                                                                                                                                                                                                                                                                                                                                        "Indicates_that_the_v
IDV
                                                                                                                                                                                                                                                                                                                  "Maximum_number_of_reads_
IMF
                                                                                                                                                                                                                                                                                                      "Maximum<sub>□</sub>fraction<sub>□</sub>of<sub>□</sub>reads<sub>□</sub>
DP
 VDB
                                  "Variant_Distance_Bias_for_filtering_splice-site_artefacts_in_RNA-seq_data
RPB
                                                                                                                                                                                                                    "Mann-Whitney Uutest of Read Position Bias
MQB
                                                                                                                                                                                                         "Mann-Whitney Uutest of Mapping Quality Bias
BQB
                                                                                                                                                                                                                          "Mann-Whitney Uutest of Base Quality Bias
MQSB
                                                                                                                                                 \verb|"Mann-Whitney|| U_{\sqcup} test_{\sqcup} of_{\sqcup} \verb|Mapping|| Quality_{\sqcup} vs_{\sqcup} Strand_{\sqcup} Bias
 SGB
                                                                                                                                                                                                                                                                                                                                                                                                                     "Segreg
MQOF
                                                                                                                                                                                                                                                                                                                             "Fraction of MQ0 reads to make the reads of the large term of the
 ICB
                                                                                                                                                                                                                                               "Inbreeding Coefficient Binomial test
HOB
                                                                                                                                                                                                                                                           "Bias_in_the_number_of_HOMs_number_
AC
                                                                                                                                      "Allele\_count\_in\_genotypes\_for\_each\_ALT\_allele,\_in\_the\_s
                                                                                                                                                                                                                                                                                                                  "Total_{\sqcup}number_{\sqcup}of_{\sqcup}alleles_{\sqcup}
 AN
DP4
                                                                                    \verb"Number_lof_lhigh-quality_ref-forward_l,_lref-reverse,_lalt-forward_land or the property of the property of
MQ
                                                                                                                                                                                                                                                                                                                                                                                                                                "Aver
```

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("../tests/data/rotavirus_rf.01.vcf",FALSE)
fmts <- bcf.formats(fp)
# dispose the vcf reader
bcf.close(fp)
# print the table
fmts</pre>
```

Output:

```
> # load rbcf
> library(rbcf)
> # we don't need the index for this file
> fp <- bcf.open("../tests/data/rotavirus_rf.01.vcf",FALSE)</pre>
> fmts <- bcf.formats(fp)</pre>
> # dispose the vcf reader
> bcf.close(fp)
[1] TRUE
> # print the table
> fmts
   ID Number
                Type
                                                        Description
PL PL
           G Integer "List⊔of⊔Phred-scaled⊔genotype⊔likelihoods"
           1 String
GT GT
                                                         "Genotype"
```

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("../tests/data/gnomad.exomes.r2.0.1.sites.vcf",FALSE)
flt <- bcf.filters(fp)
# dispose the vcf reader
bcf.close(fp)
# print the table
flt</pre>
```

Output:

```
> # load rbcf
> library(rbcf)
> # we don't need the index for this file
> fp <- bcf.open("../tests/data/gnomad.exomes.r2.0.1.sites.vcf",FALSE)</pre>
> flt <- bcf.filters(fp)</pre>
> # dispose the vcf reader
> bcf.close(fp)
[1] TRUE
> # print the table
> flt
                                 ID
PASS
                              PASS
AC<sub>0</sub>
                                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_{\sqcup}random_{\sqcup}forests_{\sqcup}filte
SEGDUP
```

2.6 Print the Samples in the VCF header

Code:

```
# load rbcf
library(rbcf)
# we don't need the index for this file
fp <- bcf.open("../tests/data/rotavirus_rf.01.vcf",FALSE)
# print the number of samples
cat(paste("Num._Samples=",bcf.nsamples(fp),".\n"))
# get the name for the 1st sample
cat(paste("First_sample_lis_",bcf.sample1(fp,1),".\n"))
# get the samples</pre>
```

```
samples <- bcf.samples(fp)
# dispose the vcf reader
bcf.close(fp)
# print the list
samples</pre>
```

Output:

```
> # load rbcf
> library(rbcf)
> # we don't need the index for this file
> fp <- bcf.open("../tests/data/rotavirus_rf.01.vcf",FALSE)</pre>
> # print the number of samples
> cat(paste("Num. Lamples=", bcf.nsamples(fp), ".\n"))
Num. Samples= 5 .
> # get the name for the 1st sample
> cat(paste("First_sample_is_",bcf.sample1(fp,1),".\n"))
First sample is S1 .
> # get the samples
> samples <- bcf.samples(fp)</pre>
> # dispose the vcf reader
> bcf.close(fp)
[1] TRUE
> # print the list
> samples
[1] "S1" "S2" "S3" "S4" "S5"
```

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("../tests/data/rotavirus_rf.01.vcf",FALSE)
dict <- bcf.dictionary(fp)
# dispose the vcf reader
bcf.close(fp)
# print the table
dict</pre>
```

Output:

```
> # load rbcf
> library(rbcf)
> # we don't need the index for this file
> fp <- bcf.open("../tests/data/rotavirus_rf.01.vcf",FALSE)</pre>
> dict <- bcf.dictionary(fp)</pre>
> # dispose the vcf reader
> bcf.close(fp)
[1] TRUE
> # print the table
> dict
     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
```

2.8 Print the Indexed Chromosomes

Code:

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

Output:

```
> # load rbcf
> library(rbcf)
> # Open the indexed VCF
> fp <- bcf.open("../tests/data/rotavirus_rf.02.vcf.gz")
> # get the indexed contigs
> contigs <- bcf.contigs(fp)
> # dispose the vcf reader
> bcf.close(fp)
[1] TRUE
> # print the table
> contigs
  [1] "RF01" "RF02" "RF03" "RF04" "RF05" "RF06" "RF07" "RF08" "RF09" "RF10"
[11] "RF11"
>
```