

Abstract

Motivation: Visualizing BAM and VCF files is a common task for biologists, but they're missing a way to filter and to explore the details of each short-read or variation.

Results: In that context, we wrote an interactive java-based interface named *JfxNgs* that uses javascript snippets to filter and reformat BAM and VCF files.

Availability: <https://github.com/lindenb/jvarkit/blob/master/docs/JfxNgs.md>

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JfxNgs : A BAM/VCF viewer with javascript-based filtering/reformatting functionalities.

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1 Description

Powerful tools like the *Integrative Genomics Viewer (IGV)* (Thorvalds *et al.*, 2013) are very efficient at visualizing the features laying in a genomic region but they're not really suitable for exploring the details of each item (quality, functional annotations, ...). Biologists in our lab currently explore the VCF files using knime4bio (Lindenbaum *et al.*, 2011), a plugin we developped for the knime plateforme. But by breaking the VCF structure into a table, we lose important informations like the VCF header, and we cannot use the existing programming interfaces like the '*java API for high-throughput sequencing data formats*' (htsjdk) (htsjdk, 2016) to analyze the data. Hence, we have written '*JfxNgs*' a java heavy client displaying the items of some VCF and BAM files. Nevertheless we do not see *JfxNgs* as a competitor of *IGV* but rather like a companion that will be used to get the details of each record in the file.

We have taken advantage that the standard edition of 'java' distributed by Oracle contains '*nashorn*', an embedded javascript engine. This engine gives the ability to manipulate java objects using simple javascript statements. When those objects are created by htsjdk (htsjdk, 2016), it gives the users a very simple way to filter or reformat a VCF or a BAM file. We had previously implemented this idea in the tools *FilterVcf* and *FilterSamReads* from the *picard* (Picard, 2017) suite and in *jvarkit* (Lindenbaum, 2015), and we have now implemented it in the *JfxNgs* interface (Fig. 1 and 2).

```
function myfilter(vc) {
  var countDP = 0;
  for (var i = 0; i < vc.getNSamples() ; ++i) {
    var genotype = vc.getGenotype(i);
    if ( ! genotype.hasDP() ) continue;
    var dp = genotype.getDP();
    if ( dp >= 200 ) continue;
    if ( ++countDP >= 2 ) return true;
  }
  return false;
}
myfilter( variant );
```

Figure 1: A simple javascript-based filter retaining the variants having at least two genotypes with a depth lower than 200 reads: an instance of the htsjdk java class `htsjdk.variant.variantcontext.VariantContext` named '*variant*' is injected in the javascript context, the function '*myfilter*' is invoked. It runs a loop over all the samples and count the number of genotypes having a depth ('DP') lower than 200.

```
while( iter.hasNext() ) {
  var samread = iter.next();
  if ( samread.getReadUnmappedFlag() ) continue;
  out.println( samread.getContig() + "\t" +
    (samread.getStart() - 1) + "\t" +
    samread.getEnd() );
}
```

Figure 2: Reformatting a BAM file to a BED file. An iterator named '*iter*', and scanning some instances of the htsjdk java class `htsjdk.samtools.SAMRecord`, is injected in the javascript context. As long as we can read a SAM record, mapped on the reference, its' genomic location is printed to '*out*', the current output stream.

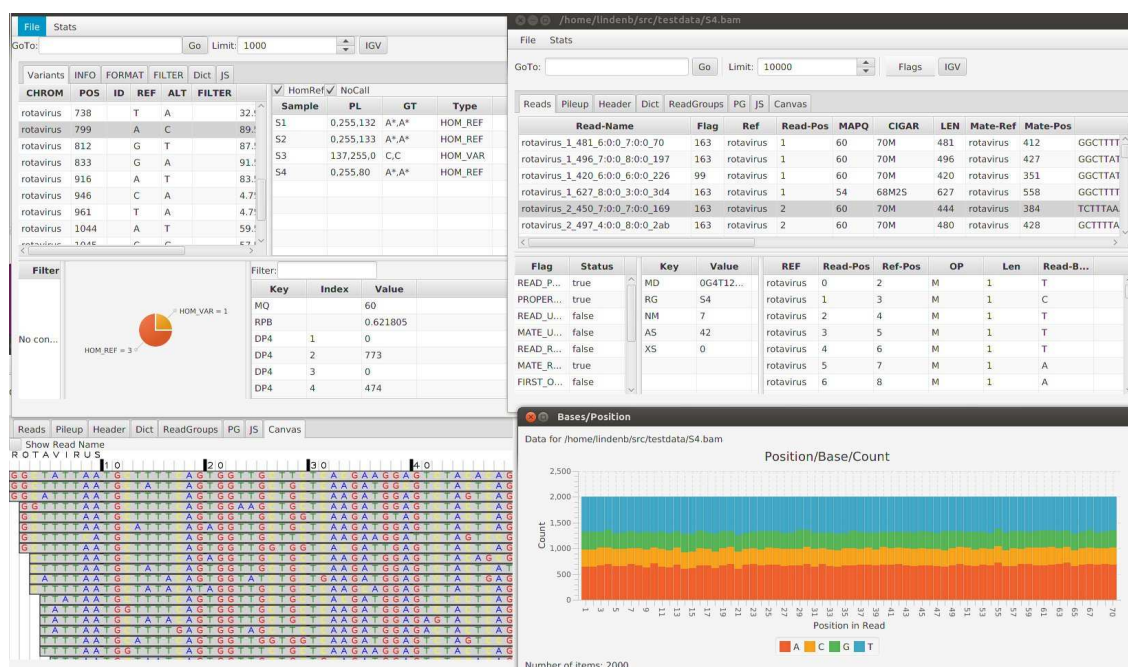


Figure 3: A Screen of jfxngs with a VCF and a BAM window. The 'JS' tabs provide a text area to write a javascript code to filter the data.

JfxNgs can read local files as long as they've been indexed with tabix or tribble. It can also access the remote files if the hosting server supports 'Byte-Range' requests. The main window provides tools for indexing BAM and VCF files. The VCF and the BAM windows have common functionalities: filtering the data with javascript, displaying the data in a very simple genome browser, viewing the selected item in a web browser for a common database like Exac (Song *et al.*, 2015), displaying simple statistics (Fig. 3), exporting the data with javascript, viewing the current selected genomic interval in *IGV*, viewing the components of the file header, saving the filtered data in a new file. The javascript areas contain a growing library of code snippets to help the user.

The BAM window shows the standard columns of the BAM specification (Read-Name, Sam-Flags, etc...), it provides some tables to view the details of the cigar string, the sam flags, the supplementary alignments. A pileup-like table displays the bases covering each position.

The VCF window displays the standard columns of a VCF file (CHROM, POS, etc...) , it provides some tables to clearly view the INFO, FILTER, ALTS data. If the annotations of a prediction algorithm like VEP (McLaren *et al.*, 2010) or SNPEff (Cingolani *et al.*, 2012) are detected, the predictions for each transcripts are displayed in a new table. A table of each genotypes displays the calls for each variant with the ability to filter out the Hom-Ref and No-Call genotypes. A pedigree file can be associated to a VCF and is then used to detect the Mendelian incompatibilities.

2 Availability

The software is available in the jvarkit package (Lindenbaum, 2015). It has been tested under Linux, Windows and MacOS and is freely available at <https://github.com/lindenb/jvarkit/blob/master>. At the time of writing, our tool is also available, packaged as a java webstart application, at http://redonlab.univ-nantes.fr/public_html/jnlp/jfxngs, meaning that it doesn't require any installation but java and an always up-to-date application is downloaded each time the user invokes it.

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