Dei M. Elurbe 1,2 and Robert Castelo 3,4

December 5, 2017

- ¹CIBER de Enfermedades Raras (CIBERER), Barcelona, Spain
- ²Present address: CMBI, Radboud University Medical Centre, Nijmegen, The Netherlands.
- ²Department of Experimental and Health Sciences, Universitat Pompeu Fabra, Barcelona, Spain.
- 3 Research Program on Biomedical Informatics (GRIB), Hospital del Mar Medical Research Institute, Barcelona, Spain

1 Overview

The aim of this software package is to facilitate the filtering and annotation of coding and non-coding genetic variants from a group of unrelated individuals, or a family of related ones among which at least one of them is affected by a genetic disorder. When working with related individuals, *VariantFiltering* can search for variants from the affected individuals that segregate according to a particular inheritance model acting on autosomes (dominant, recessive homozygous or recessive heterozygous -also known as compound heterozygous) or on allosomes (X-linked), or that occur *de novo*. When working with unrelated individuals, no mode of inheritance is used for filtering but it can be used to search for variants shared among individuals affected by a common genetic disorder.

The main input is a multisample Variant Call Format (VCF) file which is parsed using the R/Bioconductor infrastructure, and particularly the functionality from the *VariantAnnotation* package for that purpose. This infrastructure also allows *VariantFiltering* to stream over large VCF files to reduce the memory footprint and to annotate the input variants with diverse functional information.

A core set of functional data are annotated by *VariantFiltering* but this set can be modified and extended using Bioconductor annotation packages such as *MafDb.1Kgenomes.phase3.hs37d5*, which stores and exposes to the user minor allele frequency (MAF) values frozen from the latest realease of the 1000 Genomes project.

The package contains a toy data set to illustrate how it works through this vignette, and it consists of a multisample VCF file with variants from chromosomes 20, 21, 22 and allosomes X, Y from a trio of CEU individuals of the 1000 Genomes project. To further reduce the execution time of this vignette, only the code for the first analysis is actually evaluated and its results reported.

2 Setting up the analysis

To start using *VariantFiltering* the user should consider installing the packages listed in the Suggests: field from its DESCRIPTION file. After loading *VariantFiltering* the first step is to build a parameter object, of class *VariantFilteringParam* which requires at least a character string with the input VCF filename, as follows:

```
> library(VariantFiltering)
> CEUvcf <- file.path(system.file("extdata", package="VariantFiltering"),</pre>
                      "CEUtrio.vcf.bgz")
> vfpar <- VariantFilteringParam(CEUvcf)</pre>
> class(vfpar)
[1] "VariantFilteringParam"
attr(,"package")
[1] "VariantFiltering"
> vfpar
VariantFiltering parameter object
  VCF file(s): CEUtrio.vcf.bgz
  Genome version(s): hg19(NCBI)
  Number of individuals: 3 (NA12878, NA12891, NA12892)
  Genome-centric annotation package: BSgenome.Hsapiens.1000genomes.hs37d5 (1000genomes hs37d5 hs37d5)
  Variant-centric annotation package: SNPlocs.Hsapiens.dbSNP144.GRCh37 (dbSNP dbSNP Human BUILD 144)
  Transcript-centric annotation package: TxDb.Hsapiens.UCSC.hg19.knownGene
  Gene-centric annotation package: org.Hs.eg.db
  Radical/Conservative AA changes: AA_chemical_properties_HanadaGojoboriLi2006.tsv
  Codon usage table: humanCodonUsage.txt
  Regions to annotate: CodingVariants, IntronVariants, FiveSpliceSiteVariants, ThreeSpliceSiteVariants, Prom
  Other annotation pkg/obj: MafDb.1Kgenomes.phase1.hs37d5,
                            PolyPhen.Hsapiens.dbSNP131,
                            SIFT.Hsapiens.dbSNP137,
                            phastCons100way.UCSC.hg19,
                            humanGenesPhylostrata
  All transcripts: FALSE
```

The display of the *VariantFilteringParam* object indicates a number of default values which can be overriden when calling the construction function. To quickly see all the available arguments and their default values we should type:

The manual page of VariantFilteringParam contains more information about these arguments and their default values.

3 Annotating variants

After setting up the parameters object, the next step is to annotate variants. This can be done using upfront an inheritance model that will substantially filter and reduce the number of variants and annotations or, as we illustrate here below, calling the function unrelatedIn
dividuals()
that just annotates the variants without filtering out any of them:

```
> uind <- unrelatedIndividuals(vfpar)</pre>
```

```
> class(uind)
[1] "VariantFilteringResults"
attr(,"package")
[1] "VariantFiltering"
> uind
VariantFiltering results object
  Genome version(s): "hq19"(NCBI) "hq19"(Ensembl)
  Number of individuals: 3 (NA12878, NA12891, NA12892)
  Variants segregate according to a(n) unrelated individuals inheritance model
  Quality filters
INDELQual
          LowQual
                      SNPQual
     TRUE
               TRUE
                         TRUE
  Functional annotation filters
     variantType
                                           location
                                                         consequence
           FALSE
                                              FALSE
                                                               FALSE
                            FALSE
    codonusageFC
                                                              maxMAF
                             MIMO
                                      aaChangeType
                            FALSE
           FALSE
                                              FALSE
                                                               FALSE
 phastCons100way genePhyloStratum
                                            S0terms
           FALSE
                            FALSE
                                              FALSE
```

The resulting object belongs to the *VariantFilteringResults* class of objects, defined within *VariantFiltering*, whose purpose is to ease the task of filtering and prioritizing the annotated variants. The display of the object just tells us the genome version of the input VCF file, the number of individuals, the inheritance model and what variant filters are activated.

To get a summary of the number of variants annotated to a particular feature we should use the function summary():

```
> summary(uind)
                        Description Nr. Variants % Variants
1 S0:0002012
                         start_lost
                                                      0.27
                                             1
2 S0:0001631 upstream_gene_variant
                                             52
                                                     14.13
3 S0:0001624
               3_prime_UTR_variant
                                                      4.35
                                             16
4
  S0:0001623
                5_prime_UTR_variant
                                             13
                                                      3.53
5 S0:0001589
                frameshift\_variant
                                             1
                                                      0.27
6 S0:0001587
                                             2
                                                      0.54
                        stop_gained
7 S0:0001583
                   missense_variant
                                             16
                                                      4.35
8 S0:0001575
              splice_donor_variant
                                              3
                                                      0.82
9 S0:0001574 splice_acceptor_variant
                                              6
                                                      1.63
10 S0:0001627
                     intron_variant
                                            309
                                                     83.97
11 S0:0001819
                  synonymous\_variant
                                                      4.35
                                             16
```

The default setting of the <code>summary()</code> function is to provide feature annotations in terms of Sequence Ontology (SO) terms. The reported number of variants refer to the number of different variants in the input VCF file annotated to the feature while the percentage of variants refers to the fraction of this number over the total number of different variants in the input VCF file.

We can also obtain a summary based on the Bioconductor feature annotations provided by the functions locateVariants() and predictCoding() from the VariantAnnotation package, as follows:

```
> summary(uind, method="bioc")
           BIOCID Nr. Variants % Variants
2
           intron
                       309
                                  83.97
3
          fiveUTR
                           13
                                   3.53
4
         threeUTR
                           16
                                   4.35
5
                           35
                                   9.51
           coding
7
                           52
                                  14.13
         promoter
   fiveSpliceSite
                           3
                                   0.82
8
9 threeSpliceSite
                           6
                                   1.63
10
       frameshift
                           1
                                   0.27
                            2
                                    0.54
11
         nonsense
12
    nonsynonymous
                           16
                                   4.35
                                    4.35
13
       synonymous
                           16
```

Since SO terms are organized hierarchically, we can use this structure to aggregate feature annotations into more coarse-grained SO terms using the argument method="SOfull":

```
> uindS0 <- summary(uind, method="S0full")</pre>
> uindS0
        SOID Level
                                Description Nr. Variants % Variants
1 S0:0002012
                                 start_lost
                                                              0.27
                                                      1
2 S0:0001582
                 8
                    initiator_codon_variant
                                                      1
                                                              0.27
3 S0:0001819 8
                         synonymous_variant
                                                     16
                                                              4.35
4 S0:0001631 5
                     upstream_gene_variant
                                                      52
                                                             14.13
5 S0:0001628
                         intergenic_variant
                                                      52
                                                             14.13
                 4
6 S0:0001624
                 8
                         3_prime_UTR_variant
                                                      16
                                                              4.35
```

7	S0:0001623	8	5_prime_UTR_variant	13	3.53	
8	S0:0001622	7	UTR_variant	29	7.88	
9	S0:0001589	9	${\sf frameshift_variant}$	1	0.27	
10	S0:0001587	5	$stop_{-}gained$	2	0.54	
11	S0:0001906	4	$feature_truncation$	2	0.54	
12	S0:0001583	11	${ t missense_variant}$	16	4.35	
13	S0:0001992	10	${\sf nonsynonymous_variant}$	18	4.89	
14	S0:0001650	9	${\sf inframe_variant}$	18	4.89	
15	S0:0001818	8	protein_altering_variant	19	5.16	
16	S0:0001580	7	coding_sequence_variant	35	9.51	
17	S0:0001968	6	${\tt coding_transcript_variant}$	55	14.95	
18	S0:0001791	6	exon_variant	55	14.95	
19	S0:0001575	8	splice_donor_variant	3	0.82	
20	S0:0001574	8	splice_acceptor_variant	6	1.63	
21	S0:0001629	7	${\sf splice_site_variant}$	9	2.45	
22	S0:0001627	6	${\sf intron_variant}$	310	84.24	
23	S0:0001568	6	${\sf splicing_variant}$	9	2.45	
24	S0:0001576	5	${\sf transcript_variant}$	351	95.38	
25	S0:0001564	4	$gene_{-}variant$	351	95.38	
26	S0:0001878	3	$feature_variant$	368	100.00	
27	S0:0001537	2	$structural_{-}variant$	368	100.00	
28	S0:0001060	1	${\sf sequence_variant}$	368	100.00	
29	S0:0002072	0	$sequence_comparison$	368	100.00	

Here the Level column refers to the shortest-path distance to the most general SO term sequence_variant within the SO ayclic digraph. We can use this level value to interrogate the annotations on a specific granularity:

```
> uindS0[uindS0$Level == 6, ]
        SOID Level
                                Description Nr. Variants % Variants
17 S0:0001968 6 coding_transcript_variant
                                                     55
                                                             14.95
18 S0:0001791
                                                     55
                                                             14.95
                               exon_variant
22 S0:0001627
                                                             84.24
                 6
                             intron_variant
                                                    310
23 S0:0001568
                 6
                           splicing_variant
                                                      9
                                                              2.45
```

Variants are stored internally in a *VRanges* object. We can retrieve the variants as a *Vranges* object with the function allVariants():

```
> allVariants(uind)

VRangesList of length 3
names(3): NA12878 NA12891 NA12892
```

This function in fact returns a *VRangesList* object with one element per sample by default. We can change the grouping of variants with the argument groupBy specifying the annotation column we want to use to group variants. If the specified column does not exist, then it will return a single *VRanges object* with all annotated variants.

Using the following code we can obtain a graphical display of a variant, including the aligned reads and the running coverage, to have a visual representation of its support. For this purpose we need to have the BAM files used to perform the variant calling. In this case we are using toy BAM files stored along with the *VariantFiltering* package, which for practical reasons only include a tiny subset of the aligned reads.

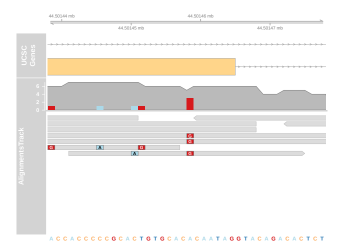


Figure 1: Browser-like display of a variant

4 Filters and cutoffs

In the case of having run the <u>unrelatedIndividuals()</u> annotation function we can filter variants by restricting the samples involved in the analysis, as follows:

```
> samples(uind)
[1] "NA12878" "NA12891" "NA12892"
> samples(uind) <- c("NA12891", "NA12892")
> uind
VariantFiltering results object
```

```
Genome version(s): "hg19"(NCBI) "hg19"(Ensembl)
 Number of individuals: 2 (NA12891, NA12892)
 Variants segregate according to a(n) unrelated individuals inheritance model
 Quality filters
INDELQual
            LowQual
                      SNPQual
     TRUE
               TRUE
                         TRUE
 Functional annotation filters
     variantType
                            dbSNP
                                           location
                                                         consequence
           FALSE
                            FALSE
                                              FALSE
                                                               FALSE
    codonusageFC
                             MIMO
                                       aaChangeType
                                                              maxMAF
           FALSE
                                              FALSE
                                                               FALSE
                            FALSE
phastCons100way genePhyloStratum
                                            S0terms
           FALSE
                            FALSE
                                              FALSE
> uindS02sam <- summary(uind, method="S0full")</pre>
> uindS02sam[uindS02sam$Level == 6, ]
                                  Description Nr. Variants % Variants
         SOID Level
17 S0:0001968
                  6 coding_transcript_variant
                                                         50
                                                                 15.34
18 S0:0001791
                                 exon_variant
                                                         50
                                                                 15.34
22 S0:0001627
                  6
                               intron_variant
                                                        272
                                                                 83.44
23 S0:0001568
                  6
                                                          7
                                                                  2.15
                             splicing_variant
```

As we can see, restricting the samples for filtering variants results in fewer variants. We can set the original samples back with the function resetSamples():

```
> uind <- resetSamples(uind)</pre>
> uind
VariantFiltering results object
  Genome version(s): "hg19"(NCBI) "hg19"(Ensembl)
  Number of individuals: 3 (NA12878, NA12891, NA12892)
  Variants segregate according to a(n) unrelated individuals inheritance model
  Quality filters
INDELQual
          LowQual
                     SNP0ual
     TRUE
               TRUE
                         TRUE
  Functional annotation filters
     variantType
                            dbSNP
                                           location
                                                         consequence
           FALSE
                            FALSE
                                              FALSE
                                                               FALSE
    codonusageFC
                             MIMO
                                       aaChangeType
                                                              maxMAF
           FALSE
                            FALSE
                                              FALSE
                                                               FALSE
 phastCons100way genePhyloStratum
                                            S0terms
           FALSE
                            FALSE
                                              FALSE
```

The rest of the filtering operations we can perform on a *VariantFilteringResults* objects are implemented through the *FilterRules* class which implements a general mechanism for generating logical masks to filter vector-like objects; consult its manual page at the *IRanges* package for full technical details.

The *Variantfiltering* package provides a number of default filters, which can be extended by the user. To see which are these filters we just have to use the **filters()** function:

```
> filters(uind)
FilterRules of length 14
names(14): INDELQual LowQual SNPQual ... genePhyloStratum S0terms
```

Filters may be active or inactive. Only active filters will participate in the filtering process when we interrogate for variants. To know what filters are active we should use the active() function as follows:

```
> active(filters(uind))
      INDELQual
                         LowQual
                                          SNPQual
                                                       variantType
           TRUE
                            TRUE
                                            TRUE
                                                             FALSE
          dbSNP
                        location
                                      consequence
                                                      codonusageFC
          FALSE
                           FALSE
                                           FALSE
                                                             FALSE
           MIMO
                    aaChangeType
                                          maxMAF phastCons100way
          FALSE
                           FALSE
                                            FALSE
                                                             FALSE
genePhyloStratum
                         S0terms
          FALSE
                           FALSE
```

By default, all filters are always inactive. To activate all of them, we can simply type:

```
> active(filters(uind)) <- TRUE</pre>
> active(filters(uind))
       INDELQual
                         LowQual
                                           SNPQual
                                                        variantType
           TRUE
                             TRUE
                                              TRUE
                                                               TRUE
           dbSNP
                         location
                                       consequence
                                                       codonusageFC
            TRUE
                             TRUE
                                             TRUE
                                                               TRUE
            MIMO
                     aaChangeType
                                            maxMAF
                                                    phastCons100way
            TRUE
                             TRUE
                                              TRUE
                                                               TRUE
genePhyloStratum
                          S0terms
            TRUE
                             TRUE
> summary(uind)
        SOID
                       Description Nr. Variants % Variants
1 S0:0001631 upstream_gene_variant
                                              1
                                                        11
2 S0:0001624
              3_prime_UTR_variant
                                              1
                                                        11
                                                        33
3 S0:0001623
               5_prime_UTR_variant
                                              3
4 S0:0001819
                synonymous_variant
                                              9
                                                       100
```

To deactivate all of them back and selectively activate one of them, we should use the bracket [] notation, as follows:

```
> active(filters(uind)) <- FALSE</pre>
> active(filters(uind))["dbSNP"] <- TRUE</pre>
> summary(uind)
         SOID
                          Description Nr. Variants % Variants
1 S0:0002012
                           start_lost
                                                         0.24
                                                1
2 S0:0001631 upstream_gene_variant
                                                55
                                                        13.16
                3_prime_UTR_variant
3 S0:0001624
                                                16
                                                         3.83
4 S0:0001623
                 5_prime_UTR_variant
                                                11
                                                         2.63
5 S0:0001587
                          stop_gained
                                                 2
                                                         0.48
```

```
6 S0:0001583
                                                 18
                                                          4.31
                     missense_variant
7 S0:0001575
                 splice_donor_variant
                                                  3
                                                          0.72
8 S0:0001574 splice_acceptor_variant
                                                  6
                                                          1.44
9 S0:0001627
                       intron_variant
                                                356
                                                         85.17
10 S0:0001819
                   synonymous\_variant
                                                          3.83
                                                 16
```

The previous filter just selects variants with an annotated dbSNP identifier. However, other filters may require cutoff values to decide what variants pass the filter. To set those values we can use the function cutoffs(). For instance, in the case of the SOterms filter, we should use set the cutoff values to select variants annotated to specific SO terms. Here we select, for instance, those annotated in UTR regions:

```
> change(cutoffs(uind), "S0terms") <- "UTR_variant"</pre>
> active(filters(uind))["S0terms"] <- TRUE</pre>
> summary(uind)
                 Description Nr. Variants % Variants
[1] SOID
<0 rows> (or 0-length row.names)
> summary(uind, method="S0full")
         SOID Level
                                  Description Nr. Variants % Variants
1 S0:0001624
                  8
                          3_prime_UTR_variant
                                                                    59
                                                         16
2 S0:0001623
                          5_prime_UTR_variant
                                                         11
                                                                    41
3 S0:0001622
                  7
                                  UTR_variant
                                                         27
                                                                   100
4 S0:0001968
                  6 coding_transcript_variant
                                                         27
                                                                   100
5 S0:0001791
                                                         27
                                                                   100
                  6
                                 exon_variant
6 S0:0001576
                  5
                           transcript_variant
                                                         27
                                                                   100
7 S0:0001564
                  4
                                 gene_variant
                                                         27
                                                                   100
8 S0:0001878
                  3
                                                         27
                              feature_variant
                                                                   100
9 S0:0001537
                  2
                           structural_variant
                                                         27
                                                                   100
10 SO:0001060
                  1
                             sequence_variant
                                                         27
                                                                   100
11 S0:0002072
                          sequence_comparison
                                                         27
                                                                   100
```

Note that the first call to summary() did not report any variant since there are no variants annotated to the SO term UTR_variant. However, when using the argument method="SOfull", all variants annotated to more specific SO terms in the hierarchy will be reported.

The methods <code>filters()</code> and <code>cutoffs()</code> can be employed to extend the filtering functionality. Here we show a simple example in which we add a filter to detect the loss of the codon that initiates translation. This constitutes already a feature annotated by <code>VariantFiltering</code> so that we can verify that it works:

```
INDELQual
                          LowQual
                                           SNPQual
                                                        variantType
           FALSE
                            FALSE
                                             FALSE
                                                               FALSE
           dbSNP
                         location
                                       consequence
                                                       codonusageFC
           FALSE
                            FALSE
                                             FALSE
                                                               FALSE
           MIMO
                     aaChangeType
                                            maxMAF
                                                    phastCons100way
           FALSE
                            FALSE
                                             FALSE
                                                               FALSE
genePhyloStratum
                          S0terms
                                         startLost
           FALSE
                            FALSE
                                              TRUE
> summary(uind)
        SOID
                       Description Nr. Variants % Variants
1 S0:0002012
                       start_lost
                                              1
                                              1
2 S0:0001631 upstream_gene_variant
                                                       100
3 S0:0001623 5_prime_UTR_variant
                                              1
                                                       100
4 S0:0001583
                 missense_variant
                                              1
                                                       100
5 S0:0001575 splice_donor_variant
                                              1
                                                        100
6 S0:0001627
                    intron_variant
                                                        100
```

As we can see, our filter works as expected and selects the only variant which was annotated with the SO term start_lost. Note that there is also an additional annotation indicating this variant belongs to an UTR region resulting from an alternative CDS.

Properly updating cutoff values may be problematic if we do not know how exactly are they employed by the corresponding filters. To facilitate setting the right cutoff values we use the change()) method and we illustrate it on the filter of minor allele frequency (MAF) values:

```
> active(filters(uind)) <- FALSE</pre>
> active(filters(uind))["maxMAF"] <- TRUE</pre>
> cutoffs(uind)$maxMAF
CutoffsList of length 2
names(2): popmask maxvalue
> cutoffs(uind)$maxMAF$popmask
    AFKGp1 AFR_AFKGp1 AMR_AFKGp1 ASN_AFKGp1 EUR_AFKGp1
                                        TRUE
      TRUE
                 TRUE
                             TRUE
                                                    TRUE
> cutoffs(uind)$maxMAF$maxvalue
[1] 0.5
> change(cutoffs(uind)$maxMAF, "popmask") <- FALSE</pre>
> cutoffs(uind)$maxMAF$popmask
    AFKGp1 AFR_AFKGp1 AMR_AFKGp1 ASN_AFKGp1 EUR_AFKGp1
                FALSE
                           FALSE
                                       FALSE
                                                   FALSE
     FALSE
> change(cutoffs(uind)$maxMAF, "popmask") <- c(AFKGp1=TRUE)</pre>
> cutoffs(uind)$maxMAF$popmask
    AFKGp1 AFR_AFKGp1 AMR_AFKGp1 ASN_AFKGp1 EUR_AFKGp1
      TRUE
                FALSE
                            FALSE
                                        FALSE
                                                   FALSE
> change(cutoffs(uind)$maxMAF, "maxvalue") <- 0.01</pre>
> summary(uind)
```

SOID	Description N	Ur Variants	% Variants
1 SO:0001631	•	23	15.86
	upstream_gene_variant		
2 S0:0001624	$3_{ extsf{prime}}_{ extsf{UTR}}$ variant	3	2.07
3 S0:0001623	5_prime_UTR_variant	5	3.45
4 S0:0001587	stop_gained	1	0.69
5 S0:0001583	${\tt missense_variant}$	6	4.14
6 S0:0001575	splice_donor_variant	1	0.69
7 S0:0001574	splice_acceptor_variant	3	2.07
8 S0:0001627	intron_variant	125	86.21
9 S0:0001819	$synonymous_variant$	4	2.76

In this case we selected variants with MAF< 0.01 in the global population of the 1000 Genomes project. If we are interested in retrieving the actual set of filtered variants, we can do it using the function filteredVariants():

```
> filteredVariants(uind)

VRangesList of length 3
names(3): NA12878 NA12891 NA12892
```

To further understand how to manipulate *Vranges* and *VRangesList* objects, please consult the package *VariantAnnotation*.

5 Inheritance models

We can filter upfront variants that do not segregate according to a given inheritance model. In such a case, we also need to provide a PED file at the time we build the parameter object, as follows:

Here we are using a PED file included in the *VariantFiltering* package and specifying information about the CEU trio employed int his vignette.

To use an inherintance model we need to replace the previous call to unrelatedIndividuals()
by one specific to the inheritance model. The VariantFiltering package offers 5 possible ones:

Autosomal recessive inheritance analysis: Homozygous.

Homozygous variants responsible for a recessive trait in the affected individuals can be identified calling the autosomalRecessiveHomozygous() function. This function selects homozygous variants that are present in all the affected individuals and occur in heterozygosity in the unaffected ones.

Autosomal recessive inheritance analysis: Heterozygous.

To filter by this mode of inheritance, also known as compound heterozygous, we need two unaffected parents/ancestors and at least one affected descendant. Variants are filtered in five steps: 1. select heterozygous variants in one of the parents and homozygous in the other; 2. discard previously selected variants that are common between the two parents; 3. group variants by gene; 4. select those genes, and the variants that

occur within them, which have two or more variants and there is at least one from each parent; 5. from the previously selected variants, discard those that do not occur in the affected descendants. This is implemented in the function autosomalRecessive Heterozygous().

Autosomal dominant inheritance analysis.

The function autosomalDominant() identifies variants present in all the affected individual(s) discarding the ones that also occur in at least one of the unaffected subjects.

X-Linked inheritance analysis.

The function <code>xLinked()</code> identifies variants that appear only in the X chromosome of the unaffected females as heterozygous, don't appear in the unaffected males analyzed and finally are present (as homozygous) in the affected male(s). This function is currently restricted to affected males, and therefore, it cannot search for X-linked segregating variants affecting daughters.

De Novo variants analysis

The function deNovo() searches for *de novo* variants which are present in one descendant and present in both parents/ancestors. It is currently restricted to a trio of individuals.

6 Create a report from the filtered variants

The function reportVariants() allows us to easily create a report from the filtered variants into a CSV or a TSV file as follows:

```
> reportVariants(uind, type="csv", file="uind.csv")
```

The default value on the type argument ("shiny") starts a shiny web app which allows one to interactively filter the variants, obtaining an udpated *VariantFilteringResults* object and downloading the filtered variants and the corresponding full reproducible R code, if necessary.

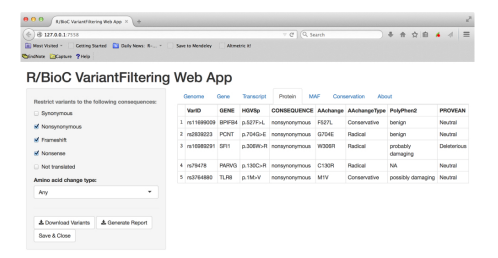


Figure 2: Snapshot of the shiny web app run from VariantFiltering with the function reportVariants() Some of the parameters has been filled for illustrative purposes.

7 Using the package with parallel execution

Functions in *VariantFiltering* to annotate and filter variants leverage the functionality of the Bioconductor package *BiocParallel* to perform in parallel some of the tasks and calculations and reduce the overall execution time. These functions have an argument called BPPARAM that allows the user to control how this parallelism is exploited. In particular the user must give as value to this argument the result from a call to the function bpparam(), which actually is its default behavior. Here below we modify that behavior to force a call being executed without parallelism. The interested reader should consult the help page of bpparam() and the vignette of the *BiocParallel* for further information.

8 Session information

> toLatex(sessionInfo())

- R Under development (unstable) (2017-10-30 r73642), x86_64-pc-linux-gnu
- Locale: LC_CTYPE=en_US.UTF-8, LC_NUMERIC=C, LC_TIME=en_US.UTF-8, LC_COLLATE=C, LC_MONETARY=en_US.UTF-8, LC_MESSAGES=en_US.UTF-8, LC_PAPER=en_US.UTF-8, LC_NAME=C, LC_ADDRESS=C, LC_TELEPHONE=C, LC_MEASUREMENT=en_US.UTF-8, LC_IDENTIFICATION=C
- Running under: CentOS Linux 7 (Core)
- Matrix products: default
- BLAS: /opt/R/R-devel/lib64/R/lib/libRblas.so
- LAPACK: /opt/R/R-devel/lib64/R/lib/libRlapack.so

- Base packages: base, datasets, grDevices, graphics, methods, parallel, stats, stats4, utils
- Other packages: AnnotationDbi 1.41.3, BSgenome 1.47.0, BSgenome.Hsapiens.1000genomes.hs37d5 0.99.1, Biobase 2.39.0, BiocGenerics 0.25.0, Biostrings 2.47.0, DelayedArray 0.5.5, GenomeInfoDb 1.15.1, GenomicFeatures 1.31.0, GenomicRanges 1.31.1, GenomicScores 1.3.0, IRanges 2.13.4, MafDb.1Kgenomes.phase1.hs37d5 3.6.0, PolyPhen.Hsapiens.dbSNP131 1.0.2, RSQLite 2.0, Rsamtools 1.31.0, S4Vectors 0.17.10, SIFT.Hsapiens.dbSNP137 1.0.0, SNPlocs.Hsapiens.dbSNP144.GRCh37 0.99.20, SummarizedExperiment 1.9.2, TxDb.Hsapiens.UCSC.hg19.knownGene 3.2.2, VariantAnnotation 1.25.1, VariantFiltering 1.15.13, XVector 0.19.1, matrixStats 0.52.2, org.Hs.eg.db 3.5.0, phastCons100way.UCSC.hg19 3.6.0, rtracklayer 1.39.0
- Loaded via a namespace (and not attached): AnnotationFilter 1.3.0, AnnotationHub 2.11.2, BiocInstaller 1.29.2, BiocParallel 1.13.0, BiocStyle 2.7.0, DBI 0.7, DT 0.2, Formula 1.2-2, GenomeInfoDbData 0.99.2, GenomicAlignments 1.15.1, Gviz 1.23.0, Hmisc 4.0-3, Matrix 1.2-12, ProtGenerics 1.11.0, R6 2.2.2, RBGL 1.55.0, RColorBrewer 1.1-2, RCurl 1.95-4.8, RMySQL 0.10.13, Rcpp 0.12.14, XML 3.98-1.9, acepack 1.4.1, assertthat 0.2.0, backports 1.1.1, base64enc 0.1-3, biomaRt 2.35.0, biovizBase 1.27.0, bit 1.1-12, bit64 0.9-7, bitops 1.0-6, blob 1.1.0, checkmate 1.8.5, cluster 2.0.6, colorspace 1.3-2, compiler 3.5.0, curl 3.0, data.table 1.10.4-3, dichromat 2.0-0, digest 0.6.12, ensembldb 2.3.1, evaluate 0.10.1, foreign 0.8-69, ggplot2 2.2.1, graph 1.57.0, grid 3.5.0, gridExtra 2.3, gtable 0.2.0, htmlTable 1.9, htmltools 0.3.6, htmlwidgets 0.9, httpuv 1.3.5, httr 1.3.1, interactiveDisplayBase 1.17.0, knitr 1.17, lattice 0.20-35, latticeExtra 0.6-28, lazyeval 0.2.1, magrittr 1.5, memoise 1.1.0, mime 0.5, miniUI 0.1.1, munsell 0.4.3, nnet 7.3-12, pkgconfig 2.0.1, plyr 1.8.4, prettyunits 1.0.2, progress 1.1.2, rlang 0.1.4, rmarkdown 1.8, rpart 4.1-11, rprojroot 1.2, scales 0.5.0, shiny 1.0.5, shinyjs 0.9.1, shinythemes 1.1.1, splines 3.5.0, stringi 1.1.6, stringr 1.2.0, survival 2.41-3, tibble 1.3.4, tools 3.5.0, xtable 1.8-2, yaml 2.1.14, zlibbioc 1.25.0