

Overview of ensemblVEP

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Contents

1	Introduction	1
2	Results as R objects	1
3	Write results to a file	5
4	Configuring runtime options	5
5	sessionInfo()	6

1 Introduction

Ensembl provides the facility to predict functional consequences of known and unknown variants using the Variant Effect Predictor (VEP). The `ensemblVEP` package wraps Ensembl VEP and returns the results as R objects or a file on disk. To use this package the Ensembl VEP perl script must be installed in your path. See the package README for details.

Downloads: <http://uswest.ensembl.org/info/docs/tools/vep/index.html>

Complete documentation for runtime options: http://uswest.ensembl.org/info/docs/tools/vep/script/vep_options.html

To test that Ensembl VEP is properly installed, enter the name of the script from the command line:

```
variant_effect_predictor.pl
```

2 Results as R objects

```
> library(ensemblVEP)
```

The `ensemblVEP` function can return variant consequences from Ensembl VEP as R objects (`GRanges` or `VCF`) or write them to a file. The default behavior returns a `GRanges`. Runtime options are stored in a `VEPParam` object and allow a great deal of control over the content and format of the results. See the man pages for more details.

```
> ?ensemblVEP
```

```
> ?VEPParam
```

The default runtime options can be inspected by creating a `VEPParam`.

```
> param <- VEPParam()
```

```
> param
```

```
class: VEPParam82
```

```
identifier(0):
```

```
colocatedVariants(0):
```

```
dataformat(0):
```

```
basic(0):
```

```
input(1): species
```

```
cache(3): dir, dir_cache, dir_plugins
```

```
output(1): terms
```

```

filterqc(0):
database(2): host, database
advanced(1): buffer_size
version: 82
scriptPath:

```

```
> basic(param)
```

```
$verbose
[1] FALSE
```

```
$quiet
[1] FALSE
```

```
$no_progress
[1] FALSE
```

```
$config
character(0)
```

```
$everything
[1] FALSE
```

```
$fork
numeric(0)
```

Using a vcf file from VariantAnnotation as input, we query Ensembl VEP with the default runtime parameters.

```

> fl <- system.file("extdata", "gl_chr1.vcf", package="VariantAnnotation")
> gr <- ensemblVEP(fl)

```

Consequence data are parsed into the metadata columns of the GRanges. To control the type and amount of data returned see the options in output(VEPParam()).

```
> head(gr, 3)
```

GRanges object with 3 ranges and 23 metadata columns:

	seqnames	ranges	strand	Allele					
	<Rle>	<IRanges>	<Rle>	<factor>					
	rs6054257	20 [14370, 14370]	*	A					
	20:17330_T/A	20 [17330, 17330]	*	A					
	rs6040355	20 [1110696, 1110696]	*	G					
		Consequence	IMPACT	SYMBOL	Gene				
		<factor>	<factor>	<factor>	<factor>				
	rs6054257	intergenic_variant	MODIFIER	<NA>	<NA>				
	20:17330_T/A	intergenic_variant	MODIFIER	<NA>	<NA>				
	rs6040355	upstream_gene_variant	MODIFIER	PSMF1	ENSG00000125818				
		Feature_type	Feature	BIOTYPE	EXON				
		<factor>	<factor>	<factor>	<factor>				
	rs6054257	<NA>	<NA>	<NA>	<NA>				
	20:17330_T/A	<NA>	<NA>	<NA>	<NA>				
	rs6040355	Transcript	ENST00000479715	processed_transcript	<NA>				
		INTRON	HGVSc	HGVSp	cDNA_position	CDS_position			
		<factor>	<factor>	<factor>	<factor>	<factor>			
	rs6054257	<NA>	<NA>	<NA>	<NA>	<NA>			
	20:17330_T/A	<NA>	<NA>	<NA>	<NA>	<NA>			
	rs6040355	<NA>	<NA>	<NA>	<NA>	<NA>			
		Protein_position	Amino_acids	Codons	Existing_variation				
		<factor>	<factor>	<factor>	<factor>				
	rs6054257	<NA>	<NA>	<NA>	<NA>	<NA>			
	20:17330_T/A	<NA>	<NA>	<NA>	<NA>	<NA>			

```

rs6040355      <NA>      <NA>      <NA>      <NA>
      DISTANCE  STRAND   FLAGS SYMBOL_SOURCE  HGNC_ID
      <factor> <factor> <factor>   <factor> <factor>
rs6054257      <NA>      <NA>      <NA>      <NA>      <NA>
20:17330_T/A   <NA>      <NA>      <NA>      <NA>      <NA>
rs6040355      2610      1      <NA>      HGNC HGNC:9571
-----

```

seqinfo: 1 sequence from genome

Next we use a vcf of structural variants as input

```
> fl <- system.file("extdata", "structural.vcf", package="VariantAnnotation")
```

and request that a VCF object be returned by setting the *vcf* option in the *dataformat* slot to TRUE.

```
> param <- VEPParam(dataformat=c(vcf=TRUE))
```

An call to *ensemblVEP* results in an error.

```
> vcf <- ensemblVEP(fl, param)
2012-12-03 16:40:55 - Starting...
ERROR: Could not detect input file format
```

In most situations Ensembl VEP can auto-detect the input format. In this case, however, it cannot so we explicitly set the *format* option to 'vcf'.

```
> input(param)$format <- "vcf"
```

Try again.

```
> vep <- ensemblVEP(fl, param)
```

Success! When a VCF is returned, consequence data are included as an unparsed INFO column labeled *CSQ*.

```
> info(vep)$CSQ
```

```

CharacterList of length 5
[[1]] deletion|intron_variant&non_coding_transcript_variant&feature_truncatio...
[[2]] -|intergenic_variant|
[[3]] insertion|upstream_gene_variant|MODIFIER|SETD5|ENSG00000168137|Transcri...
[[4]] duplication|transcript_amplification|HIGH|CRIP1P1|ENSG00000233252|Trans...
[[5]] -|intergenic_variant|

```

The *parseCSQToGRanges* function parses these data into a *GRanges*. When the rownames of the original VCF are provided as *VCFRowID* a metadata column of the same name is included in the output.

```
> vcf <- readVcf(fl, "hg19")
> csq <- parseCSQToGRanges(vep, VCFRowID=rownames(vcf))
> head(csq, 3)
```

GRanges object with 3 ranges and 24 metadata columns:

```

      seqnames      ranges strand | VCFRowID
      <Rle>      <IRanges> <Rle> | <integer>
2:321682_T/<DEL>    2 [ 321682, 321682] * | 3
2:321682_T/<DEL>    2 [ 321682, 321682] * | 3
2:14477084_C/<DEL:ME:ALU> 2 [14477084, 14477084] * | 4
      Allele
      <factor>
2:321682_T/<DEL> deletion
2:321682_T/<DEL> deletion
2:14477084_C/<DEL:ME:ALU> -
      Consequence
      <factor>
2:321682_T/<DEL> intron_variant&non_coding_transcript_variant&feature_truncation

```

```

2:321682_T/<DEL> intron_variant&non_coding_transcript_variant&feature_truncation
2:14477084_C/<DEL:ME:ALU> intergenic_variant

IMPACT SYMBOL Gene Feature_type
<factor> <factor> <factor> <factor>
2:321682_T/<DEL> MODIFIER AC079779.6 ENSG00000233684 Transcript
2:321682_T/<DEL> MODIFIER AC079779.6 ENSG00000233684 Transcript
2:14477084_C/<DEL:ME:ALU> <NA> <NA> <NA> <NA>

Feature BIOTYPE EXON INTRON HGVSc
<factor> <factor> <factor> <factor> <factor>
2:321682_T/<DEL> ENST00000436808 lincRNA <NA> 1/3 <NA>
2:321682_T/<DEL> ENST00000430529 lincRNA <NA> 1/1 <NA>
2:14477084_C/<DEL:ME:ALU> <NA> <NA> <NA> <NA> <NA>

HGVSposition cDNA_position CDS_position
<factor> <factor> <factor>
2:321682_T/<DEL> <NA> <NA> <NA>
2:321682_T/<DEL> <NA> <NA> <NA>
2:14477084_C/<DEL:ME:ALU> <NA> <NA> <NA>

Protein_position Amino_acids Codons
<factor> <factor> <factor>
2:321682_T/<DEL> <NA> <NA> <NA>
2:321682_T/<DEL> <NA> <NA> <NA>
2:14477084_C/<DEL:ME:ALU> <NA> <NA> <NA>

Existing_variation DISTANCE STRAND FLAGS
<factor> <factor> <factor> <factor>
2:321682_T/<DEL> <NA> <NA> 1 <NA>
2:321682_T/<DEL> <NA> <NA> 1 <NA>
2:14477084_C/<DEL:ME:ALU> <NA> <NA> <NA> <NA>

SYMBOL_SOURCE HGNC_ID
<factor> <factor>
2:321682_T/<DEL> Clone_based_vega_gene <NA>
2:321682_T/<DEL> Clone_based_vega_gene <NA>
2:14477084_C/<DEL:ME:ALU> <NA> <NA>

```

```
-----
seqinfo: 3 sequences from genome; no seqlengths
```

The `VCFRowID` column maps the expanded `CSQ` data back to the rows in the `VCF` object. This index can be used to subset the original VCF.

```
> vcf[csq$"VCFRowID"]
```

```
class: CollapsedVCF
```

```
dim: 22 1
```

```
rowRanges(vcf):
```

```
GRanges with 5 metadata columns: paramRangeID, REF, ALT, QUAL, FILTER
```

```
info(vcf):
```

```
DataFrame with 10 columns: BKPTID, CIEND, CIPOS, END, HOMLEN, HOMSEQ, IMPR...
```

```
info(header(vcf)):
```

	Number	Type	Description
BKPTID	.	String	ID of the assembled alternate allele in the asse...
CIEND	2	Integer	Confidence interval around END for imprecise var...
CIPOS	2	Integer	Confidence interval around POS for imprecise var...
END	1	Integer	End position of the variant described in this re...
HOMLEN	.	Integer	Length of base pair identical micro-homology at ...
HOMSEQ	.	String	Sequence of base pair identical micro-homology a...
IMPRECISE	0	Flag	Imprecise structural variation
MEINFO	4	String	Mobile element info of the form NAME,START,END,P...
SVLEN	.	Integer	Difference in length between REF and ALT alleles
SVTYPE	1	String	Type of structural variant

```
geno(vcf):
```

```
SimpleList of length 4: GT, GQ, CN, CNQ
geno(header(vcf)):
  Number Type      Description
GT 1      String    Genotype
GQ 1      Float     Genotype quality
CN 1      Integer   Copy number genotype for imprecise events
CNQ 1     Float     Copy number genotype quality for imprecise events
```

3 Write results to a file

In the previous section we saw Ensembl VEP results returned as R objects in the workspace. Alternatively, these results can be written directly to a file. The flag that controls how the data are returned is the *output_file* flag in the *input* options.

When *output_file* is an empty character (default), the results are returned as either a *GRanges* or *VCF* object.

```
> input(param)$output_file
```

```
character(0)
```

To write results directly to a file, specify a file name for the *output_file* flag.

```
> input(param)$output_file <- "/mypath/myfile"
```

The file can be written as a *vcf* or *gvf* by setting the options in the *dataformat* slot to TRUE. If neither of *vcf* or *gvf* are TRUE the file is written out as tab delimited.

```
> ## Write a vcf file to myfile.vcf:
> myparam <- VEPParam(dataformat=c(vcf=TRUE),
+                       input=c(output_file="/path/myfile.vcf"))
> ## Write a gvf file to myfile.gvf:
> myparam <- VEPParam(dataformat=c(gvf=TRUE),
+                       input=c(output_file="/path/myfile.gvf"))
> ## Write a tab delimited file to myfile.txt:
> myparam <- VEPParam(input=c(output_file="/path/myfile.txt"))
```

4 Configuring runtime options

The Ensembl VEP web page has complete descriptions of all runtime options. http://uswest.ensembl.org/info/docs/tools/vep/script/vep_options.html Below are examples of how to configure the runtime options in the *VEP-Param* for specific situations. Investigate the differences in results using a sample file from *VariantAnnotation*.

```
> fl <- system.file("extdata", "ex2.vcf", package="VariantAnnotation")
```

- Add regulatory region consequences:

```
> param <- VEPParam(output=c(regulatory=TRUE))
> gr <- ensemblVEP(fl, param)
```

- Specify input file format as VCF, add HGNC gene identifiers, output SO consequence terms:

```
> param <- VEPParam(input=c(format="vcf"),
+                   output=c(terms="so"),
+                   identifiers=c(symbol=TRUE))
> gr <- ensemblVEP(fl, param)
```

- Check for co-located variants, output only coding sequence consequences, output HGVS names:

```
> param <- VEPParam(filterqc=c(coding_only=TRUE),
+                   colocatedVariants=c(check_existing=TRUE),
+                   identifiers=c(symbol=TRUE))
> gr <- ensemblVEP(fl, param)
```

- Add SIFT score and prediction, PolyPhen prediction only, output results as VCF:

```

fl <- system.file("extdata", "chr22.vcf.gz", package="VariantAnnotation")
param <- VEPParam(output=c(sift="b", polyphen="p"),
                 dataformat=c(vcf=TRUE))
vcf <- ensemblVEP(fl, param)
csq <- parseCSQToGRanges(vcf)

> head(levels(mcols(csq)$SIFT))
[1] "deleterious(0.01)" "deleterious(0.02)" "deleterious(0.03)"
[4] "deleterious(0.04)" "deleterious(0.05)" "deleterious(0)"

> levels(mcols(csq)$PolyPhen)
[1] "benign"           "possibly_damaging" "probably_damaging"
[4] "unknown"

```

5 sessionInfo()

```
> sessionInfo()
```

```

R version 3.2.4 Revised (2016-03-16 r70336)
Platform: x86_64-pc-linux-gnu (64-bit)
Running under: Ubuntu 14.04.4 LTS

```

locale:

```

[1] LC_CTYPE=en_US.UTF-8      LC_NUMERIC=C
[3] LC_TIME=en_US.UTF-8      LC_COLLATE=C
[5] LC_MONETARY=en_US.UTF-8  LC_MESSAGES=en_US.UTF-8
[7] LC_PAPER=en_US.UTF-8     LC_NAME=C
[9] LC_ADDRESS=C             LC_TELEPHONE=C
[11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C

```

attached base packages:

```

[1] stats4    parallel  stats      graphics  grDevices  utils      datasets
[8] methods   base

```

other attached packages:

```

[1] ensemblVEP_1.10.3          VariantAnnotation_1.16.4
[3] Rsamtools_1.22.0          Biostrings_2.38.4
[5] XVector_0.10.0            SummarizedExperiment_1.0.2
[7] Biobase_2.30.0            GenomicRanges_1.22.4
[9] GenomeInfoDb_1.6.3       IRanges_2.4.8
[11] S4Vectors_0.8.11         BiocGenerics_0.16.1

```

loaded via a namespace (and not attached):

```

[1] AnnotationDbi_1.32.3      GenomicAlignments_1.6.3  zlibbioc_1.16.0
[4] BiocParallel_1.4.3       BSgenome_1.38.0          tools_3.2.4
[7] DBI_0.3.1                lambda.r_1.1.7           futile.logger_1.4.1
[10] rtracklayer_1.30.3       futile.options_1.0.0     bitops_1.0-6
[13] RCurl_1.95-4.8           biomaRt_2.26.1          RSQLite_1.0.0
[16] GenomicFeatures_1.22.13  XML_3.98-1.4

```