appreci8R – an R/Bioconductor package for filtering SNVs and short indels with high sensitivity and high PPV

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April 19, 2024

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1 Introduction

For the use of next-generation sequencing in clinical routine valid variant calling results are crucial. However, numerous variant calling tools are available. These tools usually differ in the variant calling algorithsms, the characteristics reported along with the variant calls, the recommended filtration strategies for the raw calls and thus, also in the output. Especially when calling variants with a low variant allele frequency (VAF), perfect results are hard to obtain. High sensitivity is usually accompanied by low positive predictive value (PPV).

appreci8R is a package for combining and filterating the output of differen variant calling tools according to the 'appreci8'-algorithm [3]. vcf as well as txt files containing variant calls can be evaluated. The number of variant calling tools to consider is unlimited (for the user interface version it is limited to 13). The final output contains a list of variant calls, classified as "probably true", "polymophism" or "artifact".

Important note: Currently, only hg19 is supported.

1.1 Loading the package

The package can be downloaded and installed with

> BiocManager::install("appreci8R")

After installation, the package can be loaded into R by typing

> library(appreci8R)

into the R console.

appreci8R requires the R-packages shiny, shinyjs, DT, VariantAnnotation, BSgenome, BSgenome. Hsapiens. UCSC.hg1 TxDb. Hsapiens. UCSC.hg19.knownGene, Homo.sapiens, SNPlocs. Hsapiens. dbSNP144. GRCh37, XtraSNPlocs. Hsapiens. drsnps, Biostrings, MafDb.1Kgenomes.phase3.hs37d5, MafDb.ExAC.r1.0.hs37d5, MafDb.gnomADex.r2.1.hs37d5, COSMIC.67, rentrez, PolyPhen. Hsapiens. dbSNP131, SIFT. Hsapiens. dbSNP137, seqinr, openxlsx, Rsamtools, stringr, utils, stats, GenomicRanges, S4 Vectors, GenomicFeatures, IRanges, GenomicScores and SummarizedExperiment. All of them are loaded automatically when using appreci8R.

2 appreci8R – classical

2.1 1st analysis step: Target filtration

In the 1st analysis step, all off-target calls are excluded from further analysis.

The function filterTarget covers two steps: reading input and target filtration. Available arguments are:

- output_folder The folder to write the output files into. If an empty string is provided, no files are written out.
- caller_name Name of the variant calling tool (only necessary if an output folder is provided).
- caller_folder Folder containing the variant calling results.
- caller_file_names_add Suffix for naming the variant calling files. If an empty string is provided, it is assumed that the files only contain the sample name, e.g. "Sample1.vcf".
- caller_file_type File type of the variant calling results (".vcf" or ".txt").
- caller_snv_indel SNVs and indels are reported in the same file (TRUE or FALSE).
- caller_snv_names_add Suffix for naming the variant calling files containing SNVs (only evaluated if caller_snv_indel==TRUE).
- caller_indel_names_add Suffix for naming the variant calling files containing indels (only evaluated if caller_snv_indel==TRUE).
- caller_chr Column of the variant calling input containing information on chr (default: 1).
- caller_pos Column of the variant calling input containing information on pos (default: 2).

- caller_ref Column of the variant calling input containing information on ref (default: 4).
- caller_alt Column of the variant calling input containing information on alt (default: 5).
- targetRegions Data frame object containing the target regions to be analyzed (bed-format: 1st column chr, 2nd column 0-based start pos, 3rd column 1-based end pos). Or: GRanges object containing the target regions to be analyzed.

First, all files in caller_folder of the file type caller_file_type with the suffix caller_file_names_add are read. Sample names are automatically derived from the file names (e.g. a sample name would be called "Sample1" if a file was called "Sample1.txt" and no suffix was defined; a sample would be called "Sample1.mutations" if a file was called "Sample1.mutations.vcf" and no suffix was defined, but "Sample1" if the suffix ".mutations" was defined).

If SNVs and indels are reported in separated files (in the same folder), caller_snv_indel==TRUE and caller_snv_names_add and caller_indel_names_add are defined, input from two files per sample is read and automatically combined (e.g. a sample would be called "Sample1" if files "Sample1.SNV.vcf" and "Sample1.indel.vcf" are read and caller_snv_indel==TRUE, caller_snv_names_add was defined as ".SNV" and caller_indel_names_add was defined as ".indel").

Subsequently, the read variant calling results are filtered according to the defined target region. All off-target calls are excluded from further analysis. A list of data.frames (one list per sample) with all on-target calls is returned. Every data.frame contains the columns: the SampleID (taken from the input file names), Chr. Pos. Ref and Alt.

Exemplary filtration of two vcf-files Sample1.rawMutations.vcf and Sample2.rawMutations.vcf:

```
> output_folder<-""</pre>
 target < -data.frame(chr = c("2", "4", "12", "17", "21", "X"),
                      start = c(25469500, 106196950, 12046280, 7579470, 36164400, 15838363),
                      end = c(25469510, 106196960, 12046350, 7579475, 36164410, 15838366))
 target
  chr
          start
                       end
1
    2
       25469500
                 25469510
2
    4 106196950 106196960
3
   12
       12046280
                  12046350
4
   17
        7579470
                   7579475
5
   21
       36164400
                 36164410
6
       15838363
                 15838366
> caller_folder <- system.file("extdata", package = "appreci8R")</pre>
> #Input data Sample1:
> read.table(paste(caller_folder, "/Sample1.rawMutations.vcf", sep=""),
             header = FALSE, sep = "\t", stringsAsFactors = FALSE)
  V1
           ٧2
                      V3 V4 V5
                                     V6 V7 V8
                                                           ۷9
                                                                                           V10
   2 25469502 rs2276598
                                         . NA GT:AD:DP:GQ:PL 0/1:111,113:224:99:3461,0,3634
                          С
                             T 3432.77
2 12 12044195 rs1058028
                          Τ
                             С
                               2953.77
                                         . NA GT:AD:DP:GQ:PL
                                                                0/1:91,97:188:99:2982,0,3095
3 17
     7579472 rs1042522
                          G
                             C 3074.77
                                         . NA GT:AD:DP:GQ:PL
                                                                    1/1:0,75:75:99:3103,224,0
 X 15838366 rs2301724 C T 3574.77
                                         . NA GT:AD:DP:GQ:PL 0/1:161,106:267:99:3603,0,5853
```

```
> #Input data Sample2:
> read.table(paste(caller_folder, "/Sample2.rawMutations.vcf", sep=""),
             header = FALSE, sep = "\t", stringsAsFactors = FALSE)
  V1
                        V3 V4
                                 ۷5
                                         ۷6
                                                  V7 V8
  4 106196951
                                  G 3135.77
                                                   . NA GT:AD:DP:GQ:PL
                                                                         0/1:84,86:170:99:3164,0,3076
                rs2454206
                           Α
2 12
     12046289 rs10649660
                           C CAAAG
                                      15.85 LowQual NA GT:AD:DP:GQ:PL
                                                                              0/1:52,5:57:53:53,0,2353
3 12
                                  G 2711.77
                                                   . NA GT:AD:DP:GQ:PL
                                                                         0/1:94,81:175:99:2740,0,3467
      12046341
                rs2855748
                            Α
4 17
       7579472
                                  C 1264.77
                                                   . NA GT:AD:DP:GQ:PL
                                                                              1/1:0,31:31:93:1293,93,0
                rs1042522
                            G
5 20
      31025689 rs41289852
                           G
                                  C 4098.77
                                                   . NA GT:AD:DP:GQ:PL 0/1:120,120:240:99:4127,0,4038
                                                                              0/1:4,10:14:99:299,0,119
6 21
      36164405 rs13051066
                                     270.77
                                                   . NA GT:AD:DP:GQ:PL
> filterTarget(output_folder, "GATK", caller_folder, ".rawMutations",
               ".vcf", TRUE, "", "", targetRegions = target)
[[1]]
  SampleID Chr
                    Pos Ref Alt
1 Sample1
             2 25469502
  Sample1
            17
                          G
                               С
                7579472
  Sample1
             X 15838366
                          С
                               Τ
[[2]]
  SampleID Chr
                                Alt
                     Pos Ref
  Sample2
             4 106196951
                            Α
                                  G
                            C CAAAG
2 Sample2
            12
                12046289
3
  Sample2
            12
                12046341
                            A
                                  G
```

2.2 2nd analysis step: Normalization

7579472

36164405

G

С

Τ

Sample2

Sample2

17

21

In the 2nd analysis step, all calls are normalized with respect to reporting indels, MNVs, reporting of several alternate alleles and reporting of complex indels. Available arguments are:

- output_folder The folder to write the output files into. If an empty string is provided, no files are written out.
- caller_name Name of the variant calling tool (only necessary if an output folder is provided).
- target_calls List of data.frames. One list element per sample. FilterTarget()-output can directly be taken as input.
- caller_indels_pm Deletions are reported with a "minus" (e.g. C > -A), insertions are reported with a "plus" (e.g. C > +A) (TRUE or FALSE).
- caller_mnvs MNVs are reported (e.g. CA > GT instead of C > G and A > T; or CGAG > TGAT instead of C > T and G > T) (TRUE or FALSE).

The function normalize covers two to four normalization steps:

- 1. Check alternative bases: Calls containing a "comma" are split up. A call like C > A, G is converted to C > A and an additional C > G call. This enables evaluation of the output of different callers while caller1 reports C > A, G, caller2 only reports C > A and caller3 only reports C > G. This normalization step is always performed.
- 2. Find string differences: Calls are checked for un-mutated bases. The smallest option of reporting a variant at the left-most position is chosen. For example, CAAAC > CAAC is converted to CA > C. This normalization step is always performed.
- 3. Convert indels: If deletions are reported with a "minus" and insertions are reported with a "plus", these are converted. An deletion like C > -G is converted to CG > C, while an insertion like C > +G is converted to C > CG. This normalization step is only performed if caller_indels_pm is TRUE.
- 4. Convert MNVs: If MNVs are reported, these are converted. This enables evaluation of the output of different callers if not all callers report all mutations being part of an MNV. A call like CA > GT is split up to a C > G and an A > T variant. But also a call like CGAG > TGAT is split up to C > T and G > T (G > G and A > A are not reported as they do not pass the normalization step "Find string differences"). This normalization step is only performed if caller_mnvs is TRUE.

A GRanges object with all normalized calls is returned (metadata columns: SampleID, Ref, Alt). If an output folder is provided, the output is saved as <caller_name>.normalized.txt. Exemplary normalization of a set of raw calls:

```
> output_folder <- ""</pre>
> caller_name <- ""
> sample1 <- data.frame(SampleID = c("Sample1", "Sample1", "Sample1"),</pre>
                          Chr = c("2", "17", "X"),
                          Pos = c(25469502, 7579472, 15838366),
                          Ref = c("CAG", "G", "C"),
                          Alt = c("TAT", "C", "T, A"),
                          stringsAsFactors = FALSE)
  sample2 <- data.frame(SampleID = c("Sample2", "Sample2", "Sample2", "sample2"),</pre>
                          Chr = c("4", "12", "12", "21"),
                          Pos = c(106196951, 12046289, 12046341, 36164405),
                          Ref = c("A", "C", "A", "GGG"),
                          Alt = c("G", "+AAAG", "G", "TGG"),
                          stringsAsFactors = FALSE)
> #Input:
> input <- list(sample1, sample2)</pre>
> input
[[1]]
  SampleID Chr
                      Pos Ref Alt
1 Sample1
              2 25469502 CAG TAT
2 Sample1
            17
                 7579472
                            G
                                 С
3 Sample1
              X 15838366
                            C T,A
```

```
[[2]]
  SampleID Chr
                     Pos Ref
                                Alt
             4 106196951
1 Sample2
                                 G
2 Sample2
                           C +AAAG
            12 12046289
  Sample2
            12
                12046341
                                 G
                           Α
4 sample2
            21
                36164405 GGG
                                TGG
> normalized <- appreci8R::normalize(output_folder, caller_name, input,
                                      TRUE, TRUE)
> normalized
```

GRanges object with 9 ranges and 3 metadata columns:

	-				
d	eqnames ranges	i	${\tt SampleID}$	Ref	Alt
>	<rle> <iranges></iranges></rle>	>	<character></character>	<character></character>	<character></character>
*	17 7579472	k	Sample1	G	C
*	X 15838366	¢	Sample1	С	T
*	4 106196951	k	Sample2	A	G
*	12 12046289	k	Sample2	C	CAAAG
*	12 12046341	k	Sample2	A	G
*	X 15838366	k	Sample1	С	A
*	2 25469502	k	Sample1	C	T
*	21 36164405	k	sample2	G	T
*	2 25469504	k	Sample1	G	T
			-		

seqinfo: 6 sequences from an unspecified genome; no seqlengths

2.3 3rd analysis step: Annotation

In the 3rd analysis step, all calls are annotated (using VariantAnnotation) and filtered according to the selected locations and consequences of interest.

The function annotate performs annotations of all variants in the input GRanges object (according to VariantAnnotation [2]) and filteres the annotated calls with respect to the selected locations and consequences of interest (based on the functions locateVariants and predictCoding of the package VariantAnnotation). At least one location of interest has to be chosen. If - among others - "coding" is selected, at least one consequence of interest has to be chosen as well. Avilable input arguments are:

- output_folder The folder to write the output files into. If an empty string is provided, no files are written out.
- caller_name Name of the variant calling tool (only necessary if an output folder is provided).
- normalized_calls A GRanges object. One normalized variant call per line. Necessary metadata columns are: SampleID, Ref, Alt. Normalize()-output can directly be taken as input.
- locations A vector containing the locations of interest. Accepted values are: coding, intron, threeUTR, fiveUTR, intergenic, spliceSite, promoter.

• consequences A vector containing the consequences of interest. Accepted values are: synonymous, nonsynonymous, frameshift, nonsense, not translated.

All possible transcripts are investigated. If "coding" is chosen and a variant is annotated as "coding" in only 1 out of many transcripts, the matching transcript is, together with the annotation information, reported. All the other non-matching transcripts are not reported.

A GRanges object is returned containing only the calls at the selected locations of interest and with the selected consequences of interest. Reported metadata columns are: SampleID, Ref, Alt, Location, c. (position of variant on cDNA level), p. (position of variant on protein level), AA_ref, AA_alt, Codon_ref, Codon_alt, Consequence, Gene, GeneID, TranscriptID. TxDb.Hsapiens.UCSC.hg19.knownGene is used for annotation. Whenever there is more than one transcriptID, all of them are reported, separated by commas. The first transcriptID matches the first entry under Location, c., p. etc.

If an output folder is provided, the output is saved as <caller_name>.annotated.txt. Exemplary annotation of a set of normalized calls:

```
> library(GenomicRanges)
> output_folder <- ""
> caller_name <- ""
> input <- GRanges(seqnames = c("2", "X", "4", "21"),
                   ranges = IRanges(start = c (25469504, 15838366, 106196951, 36164405),
                                     end = c (25469504,15838366,106196951,36164405)),
                   SampleID = c("Sample1", "Sample1", "Sample2", "Sample2"),
                   Ref = c("G", "C", "A", "G"),
                   Alt = c("T", "A", "G", "T"))
> #Input:
> input
GRanges object with 4 ranges and 3 metadata columns:
      segnames
                  ranges strand
                                      SampleID
                                                        Ref
         <Rle> <IRanges> <Rle> |
                                   <character> <character> <character>
  [1]
             2 25469504
                                       Sample1
                               *
                                                         G
  Γ21
             X 15838366
                               *
                                       Sample1
                                                         С
                                                                      Α
  [3]
             4 106196951
                               *
                                       Sample2
                                                         Α
                                                                      G
  [4]
            21 36164405
                                       Sample2
  seqinfo: 4 sequences from an unspecified genome; no seqlengths
> library(GO.db)
> library(org.Hs.eg.db)
> library(TxDb.Hsapiens.UCSC.hg19.knownGene)
> library(Biostrings)
> annotated <- annotate(output_folder, caller_name, input,
               locations = c("coding", "spliceSite"),
+
               consequences = c("nonsynonymous", "frameshift", "nonsense"))
```

> annotated

GRanges object with 3 ranges and 14 metadata columns:											
	seqnames	ra	nges	strand	Sample	ID	Re	ef	Alt	;	Location
	<rle></rle>	<irar< td=""><td>iges></td><td><rle></rle></td><td><pre><character< pre=""></character<></pre></td><td>c> <cha< td=""><td>aracter</td><td>c> <charac< td=""><td>cter></td><td>•</td><td><character></character></td></charac<></td></cha<></td></irar<>	iges>	<rle></rle>	<pre><character< pre=""></character<></pre>	c> <cha< td=""><td>aracter</td><td>c> <charac< td=""><td>cter></td><td>•</td><td><character></character></td></charac<></td></cha<>	aracter	c> <charac< td=""><td>cter></td><td>•</td><td><character></character></td></charac<>	cter>	•	<character></character>
[1]	2	2546	9504	*	Sample	e 1		G	T	coding,co	ding,coding
[2]	Х	1583	88366	*	Sample	e1		C	А	L	coding
[3]	4	10619	6951	*	Sample	e2		A	G	со	ding,coding
		с.		р.	AA_ref	I	AA_alt	Codon_1	ref	Codon_alt	
	<charac< td=""><td>ter></td><td><char< td=""><td>cacter></td><td><character></character></td><td><chara< td=""><td>acter></td><td><characte< td=""><td>er> <</td><td>character></td><td></td></characte<></td></chara<></td></char<></td></charac<>	ter>	<char< td=""><td>cacter></td><td><character></character></td><td><chara< td=""><td>acter></td><td><characte< td=""><td>er> <</td><td>character></td><td></td></characte<></td></chara<></td></char<>	cacter>	<character></character>	<chara< td=""><td>acter></td><td><characte< td=""><td>er> <</td><td>character></td><td></td></characte<></td></chara<>	acter>	<characte< td=""><td>er> <</td><td>character></td><td></td></characte<>	er> <	character>	
[1]	697,1264,	1264	233,4	122,422	L, L, L		M,M,M	CTG,CTG,	CTG A	TG, ATG, ATG	
[2]		864		288	N		K	I	AAC	AAA	
[3]	5284,	5347	176	52,1783	I,I		V,V	ATA,	ATA	GTA,GTA	
		Cor	seque	ence		${\tt Gene}$		${\tt GeneID}$		Transcrip	tID
		<cl< td=""><td>aract</td><td>er></td><td><chara< td=""><td>cter></td><td><ch< td=""><td>naracter></td><td></td><td><charact< td=""><td>er></td></charact<></td></ch<></td></chara<></td></cl<>	aract	er>	<chara< td=""><td>cter></td><td><ch< td=""><td>naracter></td><td></td><td><charact< td=""><td>er></td></charact<></td></ch<></td></chara<>	cter>	<ch< td=""><td>naracter></td><td></td><td><charact< td=""><td>er></td></charact<></td></ch<>	naracter>		<charact< td=""><td>er></td></charact<>	er>
[1]	nonsynony	mous,	nonsy	yn DNN	MT3A,DNMT3A,I	ASTMNC	1788,1	1788,1788	1080	8,10809,10	810
[2]		nonsy	nonyn	nous		ZRSR2		8233		75	467
[3]	nonsynony	mous,	nonsy	/n	TET2	2,TET2	547	790,54790		18308,18	309

seqinfo: 4 sequences from an unspecified genome; no seqlengths

2.4 4th analysis step: Output combination

In the 4th analysis step, all normalized and annotated calls of the different variant calling tools are combined.

The function combineOutput performs a combination of the normalized and annotated variant calls from different variant calling tools. The results are sorted according to Chr, Pos, Ref, Alt and SampleID. If two callers - caller1 and caller2 - report the same variant for the same sample, it is only reported once in the output file with a "1" in the column "caller1" and another "1" in the column "caller2". Available arguments are:

- output_folder The folder to write the output files into. If an empty string is provided, no files are written out.
- caller_names Vector containing the name of the variant calling tools.
- annotated_calls A GRangesList object. Every list element contains the variant calls of one variant calling tool. Annotate()-output can directly be taken as input. Providing a list with only one list element, i.e. evaluating only one variant calling tool, is possible, but not assumed to be useful. The point of the appreci8-algorithm lies in the combined evaluation and filtration of the output of several variant calling tools.

A GRanges object is returned containing all combined calls. Reported metadata columns are: SampleID, Ref, Alt, Location, c. (position of variant on cDNA level), p. (position of variant on protein level), AA_ref, AA_alt, Codon_ref, Codon_alt, Consequence, Gene, GeneID, TranscriptID. In addition, one column for every variant calling tool is reported, containing a "1" for every call detected by that tool and "NA" for every call not detected by that tool. If an output folder is provided, the output is saved as Results_Raw.txt.

Exemplary combination of a set of annotated calls:

```
> output_folder <- ""
> caller_name <- ""</pre>
> #Input:
> gatk <- annotated[c(1,3),]</pre>
> varscan <- annotated[c(2,3),]</pre>
> annotated<-GRangesList()</pre>
> annotated[[1]]<-gatk</pre>
> annotated[[2]]<-varscan</pre>
> annotated
GRangesList object of length 2:
[[1]]
GRanges object with 2 ranges and 14 metadata columns:
      seqnames
                  ranges strand
                                       SampleID
                                                                     Alt
                                                                                      Location
                                                         Ref
         <Rle> <IRanges> <Rle> | <character> <character> <character>
                                                                                   <character>
                                                                       T coding, coding, coding
  [1]
             2 25469504
                               *
                                        Sample1
                                                           G
  [2]
             4 106196951
                                       Sample2
                                                           Α
                                                                                 coding, coding
                  с.
                              р.
                                       AA_ref
                                                   AA_alt
                                                             Codon_ref
                                                                         Codon_alt
        <character> <character> <character> <character> <character> <character>
  [1] 697,1264,1264 233,422,422
                                                    M,M,M CTG,CTG,CTG ATG,ATG,ATG
                                       L,L,L
  [2]
          5284,5347
                       1762,1783
                                          I.I
                                                      V.V
                                                               ATA, ATA
                                                                            GTA, GTA
                 Consequence
                                                             GeneID
                                                                          TranscriptID
                                               Gene
                  <character>
                                        <character>
                                                        <character>
                                                                           <character>
  [1] nonsynonymous,nonsyn.. DNMT3A,DNMT3A,DNMT3A 1788,1788,1788 10808,10809,10810
  [2] nonsynonymous, nonsyn...
                                          TET2, TET2
                                                        54790,54790
                                                                           18308,18309
  seqinfo: 4 sequences from an unspecified genome; no seqlengths
[[2]]
GRanges object with 2 ranges and 14 metadata columns:
                                                                     Alt
      seqnames
                  ranges strand
                                       SampleID
                                                         Ref
                                                                               Location
                                                                                                  с.
         <Rle> <IRanges> <Rle> | <character> <character> <character>
                                                                            <character> <character>
  [1]
             X 15838366
                               *
                                        Sample1
                                                           С
                                                                       Α
                                                                                 coding
  [2]
             4 106196951
                                        Sample2
                                                                       G coding, coding
                                                           Α
                                                                                          5284,5347
                                    \mathtt{AA\_alt}
                                              Codon_ref
                                                           Codon_alt
                                                                                 Consequence
               р.
                        AA_ref
      <character> <character> <character> <character> <character>
                                                                                 <character>
  [1]
              288
                                          K
                                                    AAC
                                                                              nonsynonymous
                             N
                                                                 AAA
  Γ21
        1762,1783
                           I,I
                                        V.V
                                                ATA, ATA
                                                             GTA, GTA nonsynonymous, nonsyn...
                        GeneID TranscriptID
             Gene
      <character> <character> <character>
  [1]
            ZRSR2
                          8233
                                       75467
  [2] TET2,TET2 54790,54790 18308,18309
  seqinfo: 4 sequences from an unspecified genome; no seqlengths
> combined<-combineOutput("", c("GATK", "VarScan"), annotated)</pre>
```

> combined

```
GRanges object with 3 ranges and 16 metadata columns:
    segnames
                 ranges strand
                                      SampleID
                                                         Ref
                                                                      Alt
                                                                                       Location
                                   <character> <character> <character>
       <Rle> <IRanges>
                          <Rle>
                                                                                    <character>
  2
           2 25469504
                                       Sample1
                                                           G
                                                                        T coding, coding, coding
  3
           4 106196951
                                                                        G
                                       Sample2
                                                           A
                                                                                  coding, coding
              15838366
                                       Sample1
                                                           C
                                                                        A
                                                                                         coding
                с.
                             р.
                                      AA_ref
                                                   AA_alt
                                                             Codon_ref
                                                                          Codon alt
      <character> <character> <character> <character> <character> <character>
  2 697,1264,1264 233,422,422
                                       L, L, L
                                                    M, M, M CTG, CTG, CTG ATG, ATG, ATG
  3
        5284,5347
                     1762,1783
                                         I,I
                                                      V, V
                                                               ATA, ATA
                                                                            GTA, GTA
  4
               864
                            288
                                           N
                                                        K
                                                                    AAC
                                                                                 AAA
                                               Gene
                                                             GeneID
                                                                          TranscriptID
                                                                                              GATK
                Consequence
                <character>
                                       <character>
                                                        <character>
                                                                           <character> <numeric>
  2 nonsynonymous, nonsyn.. DNMT3A, DNMT3A, DNMT3A, 1788, 1788, 1788, 10808, 10809, 10810
                                                                                                 1
  3 nonsynonymous, nonsyn..
                                         TET2, TET2
                                                       54790,54790
                                                                           18308,18309
                                                                                                 1
  4
              nonsynonymous
                                              ZRSR2
                                                               8233
                                                                                  75467
                                                                                                NA
      VarScan
    <numeric>
  2
           NA
  3
             1
  4
             1
```

2.5 5th analysis step: Coverage and BQ determination

seqinfo: 3 sequences from an unspecified genome; no seqlengths

In the 5th analysis step, all calls are evaluated with respect to coverage and basequality (using Rsamtools [1]). Calls with insufficient coverage and/or basequality are filtered from further analysis. The function evaluateCovAndBQ evaluates coverage and base quality of all calls and filteres them according to user-definable thresholds. Only those calls that feature sufficient coverage and base quality are reported. Filtration is performed in a 2-step procedure. Available arguments are:

- output_folder The folder to write the output files into. If an empty string is provided, no files are written out.
- combined_calls A GRanges object; necessary input object. The GRanges object contains one call per line. CombineOutput()-output can directly be taken as input.
- bam_folder The folder containing the bam- and bai-files for every sample. The file names have to match the sample names, i.e. Sample1.bam and Sample1.bai if Sample1.vcf (without any defined suffixes) was analyzed for caller1.
- dp Minimum depth that is required for a call to pass the filter (default: 50).
- nr_alt Minimum number of reads carrying the alternate allele that is required for a call to pass the filter (default: 20).
- vaf Minimum VAF that is required for a call to pass the filter (default: 0.01).

- bq Minimum base quality that is required for a call to pass the filter (default: 15; values above 44 not recommended).
- bq_diff Maximum difference in basequality between reference and alternative that is allowed for a call to pass the filter (default: 7).

First, coverage is evaluated using Rsamtools pileup and a threshold of min_base_quality=0. If depth is below the defined threshold dp and/or the number of reads carrying the alternate allele is below the defined threshold nr_alt and/or VAF is below the defined threshold VAF, a call is excluded. However, if coverage is sufficient, base quality - which is more time-consuming - is evaluated.

To evaluate base quality, the threshold for min_base_quality is successively increased from 0 to 44 (thus, no values above 44 are recommended for bq). The average base quality for all reads carrying the reference and the alternate allele is calculated. For indels, which are always summed up as "+" by Rsamtools, no base quality can be determined. If the average base quality of the alternate allele is lower than bq, the call is filtered. If "average base quality reference allele" - "average base quality alternate allele" is higher than bq_diff, the call is filtered.

Coverage is reported with respect to the forward- and reverse reads separately. It is not yet evaluated. This is done in the 7th analysis step (finalFiltration).

A GRanges object is returned containing all calls with sufficient coverage and base quality. Reported metadata columns are: SampleID, Ref, Alt, Location, c. (position of variant on cDNA level), p. (position of variant on protein level), AA_ref, AA_alt, Codon_ref, Codon_alt, Consequence, Gene, GeneID, TranscriptID, Caller1 to CallerX (dependent on the number of callers that is evaluated), Nr_Ref, Nr_Alt, DP, VAF, BQ_REF, BQ_ALT, Nr_Ref_fwd, Nr_Alt_fwd, DP_fwd, VAF_rev, Nr_Ref_rev, Nr_Alt_rev, DP_rev, VAF_rev. If an output folder is provided, the output is saved as Results Frequency.txt.

Exemplary filtration of a set of combined calls:

```
> output_folder <- ""
> bam_folder <- system.file("extdata", package = "appreci8R")
> bam_folder <- paste(bam_folder, "/", sep="")
> #Input:
> combined
```

GRanges object with 3 ranges and 16 metadata columns:

```
seqnames
               ranges strand
                                    SampleID
                                                      Ref
                                                                   Alt
                                                                                     Location
     <Rle> <IRanges>
                        <Rle>
                                <character> <character> <character>
                                                                                  <character>
2
           25469504
                                     Sample1
                                                        G
                                                                      T coding, coding, coding
                                                                               coding, coding
3
         4 106196951
                                     Sample2
                                                        A
                                                                      G
4
            15838366
                                                        C
                                                                      A
                                     Sample1
                                                                                       coding
              с.
                                    AA_ref
                                                 AA_alt
                                                           Codon_ref
                                                                        Codon alt
                           р.
    <character> <character> <character> <character> <character> <character>
2 697,1264,1264 233,422,422
                                                  M, M, M CTG, CTG, CTG ATG, ATG, ATG
                                     L,L,L
3
      5284,5347
                   1762,1783
                                       I,I
                                                    V, V
                                                                          GTA, GTA
                                                             ATA, ATA
4
             864
                          288
                                         N
                                                      K
                                                                 AAC
                                                                              AAA
                                                                        TranscriptID
                                                                                            GATK
              Consequence
                                            Gene
                                                           GeneID
              <character>
                                     <character>
                                                     <character>
                                                                         <character> <numeric>
```

```
2 nonsynonymous, nonsyn.. DNMT3A, DNMT3A, DNMT3A, 1788, 1788, 1788, 10808, 10809, 10810
                                                                                               1
  3 nonsynonymous, nonsyn...
                                        TET2, TET2
                                                      54790,54790
                                                                          18308,18309
                                                                                               1
             nonsynonymous
                                             ZRSR2
                                                              8233
                                                                                75467
                                                                                              NA
      VarScan
    <numeric>
  2
           NA
  3
            1
  4
            1
  seqinfo: 3 sequences from an unspecified genome; no seqlengths
> filtered <- evaluate Cov And BQ (output_folder, combined, bam_folder, bq_diff = 20,
                               vaf = 0.001)
> filtered
GRanges object with 2 ranges and 30 metadata columns:
                ranges strand
                                                                              Location
    seqnames
                                     SampleID
                                                       Ref
                                                                    Alt
                                                                                                 С.
       <Rle> <IRanges>
                         <Rle>
                                  <character> <character> <character>
                                                                           <character> <character>
  3
           4 106196951
                                      Sample2
                                                          A
                                                                      G coding, coding
                                                                                          5284,5347
  4
           Х
             15838366
                              *
                                      Sample1
                                                          C
                                                                                coding
                                                                                                864
                                   AA_alt
                                                                                                     Gene
                      AA_ref
                                             Codon_ref
                                                         Codon_alt
                                                                                Consequence
             р.
    <character> <character> <character> <character> <character>
                                                                                <character> <character>
  3
      1762,1783
                                      V, V
                                                            GTA, GTA nonsynonymous, nonsyn...
                                                                                               TET2, TET2
                         I,I
                                               ATA, ATA
            288
                           N
                                        K
                                                   AAC
                                                                AAA
                                                                              nonsynonymous
                                                                                                    ZRSR2
         GeneID TranscriptID
                                    GATK
                                            VarScan
                                                       Nr_Ref
                                                                  Nr_Alt
                                                                                 DP
                                                                                            VAF
                                                                                                    BQ_REF
    <character>
                  <character> <numeric> <numeric> <numeric> <numeric> <numeric> <numeric>
                                                                                      <numeric> <numeric>
  3 54790,54790
                  18308,18309
                                                  1
                                                          1268
                                                                    1283
                                                                               2551 0.50294002
                                                                                                   38.6680
                                       1
           8233
                                                          1991
                        75467
                                      NA
                                                  1
                                                                       31
                                                                               3630 0.00853994
                                                                                                   37.4591
                                                   VAF_fwd Nr_Ref_rev Nr_Alt_rev
                                                                                                VAF_rev
       BQ_ALT Nr_Ref_fwd Nr_Alt_fwd
                                          DP_fwd
                                                                                       DP_rev
    <numeric>
                <numeric>
                           <numeric> <numeric> <numeric>
                                                             <numeric>
                                                                         <numeric> <numeric> <numeric>
      38.8145
                      428
                                  469
                                                  0.522854
                                                                   840
  3
                                             897
                                                                               814
                                                                                         1654
                                                                                               0.492140
                      839
                                            1507
                                                  0.000000
  4
      18,0000
                                                                  1152
                                                                                31
                                                                                         2123
                                                                                               0.014602
```

seqinfo: 3 sequences from an unspecified genome; no seqlengths

2.6 6th analysis step: Characteristics determination

In the 6th analysis step, an extended set of characteristics for all calls is determined. This includes database check-ups and impact prediction on protein level.

The function evaluateCharacteristics determines an extended set of characteristics for the normalized, annotated, coverage and base quality filtered calls. This includes database check-ups and impact prediction on protein level. Available arguments are:

- output_folder The folder to write the output files into. If an empty string is provided, no files are written out.
- frequency_calls A GRanges object. The GRanges object contains one call per line. EvaluateCovAndBQ()-output can directly be taken as input.

- predict String defining the prediction tool to use (accepted strings are: SIFT, Provean, PolyPhen2)
- dbSNP Query dbSNP (144.GRCh37) for the presence of your variants (TRUE or FALSE; default TRUE).
- '1kgenomes' Query 1000 Genomes (phase3) for the presence of your variants (TRUE or FALSE; default TRUE).
- exacDB Query ExAC (r1.0) for the presence of your variants (TRUE or FALSE; default TRUE).
- gadDB Query Genome Aggregation Database (r2.0.1) for the presence of your variants (TRUE or FALSE; default TRUE).
- cosmicDB Query COSMIC (67) for the presence of your variants (TRUE or FALSE; default TRUE).
- clinvarDB Query ClinVar (via rentrez) for the presence of your variants (TRUE or FALSE; default TRUE).

First, the database check-up is performed. At a maximum, dbSNP, 1000Genomes, ExAC, Genome Aggregation Database, COSMIC and ClinVar can be checked. If all the variables are set to FALSE, no database is checked and none will be reported. Thus, the check-up can be disabled.

Second, the impact perdiction on protein level is performed. Impact can be predicted using either SIFT, Provean or PolyPhen2. In any case, the prediction is performed on the basis of rs-IDs. For every call, the predicted effect as well as the score is reported.

A GRanges Object is returned containing all calls, information on their presence in the selected databases and information on the predicted effect. Reported metadata columns are (if all databases are selected): SampleID, Ref, Alt, dbSNP (containing the rs-ID), dbSNP_MAF, G1000_AF, ExAC_AF, GAD_AF, CosmicID, Cosmic_Counts (number of Cosmic entries), Clin-Var, Prediction (damaging or neutral), Score (on the basis of the selected prediction tool), c. (variant on cDNA level containing position and variant), c.complement (complement of the variant; if c. is c.5284A>G, then c.complement is c.5284T>C), p. (variant on protein level containing position and amino acids).

Exemplary characteristics of a set of filtered calls:

- > output_folder <- ""</pre>
- > #Input:
- > filtered

GRanges object with 2 ranges and 30 metadata columns:

	seqnames	ranges	strar	ıd	Samp]	eID		Ref		Alt	Location		с.	
	<rle></rle>	Ranges</td <td><rle< td=""><td>> </td><td><charact< td=""><td>er></td><td><charact< td=""><td>ter></td><td><charac< td=""><td>ter></td><td><character></character></td><td><char< td=""><td>cacter></td><td></td></char<></td></charac<></td></charact<></td></charact<></td></rle<></td>	<rle< td=""><td>> </td><td><charact< td=""><td>er></td><td><charact< td=""><td>ter></td><td><charac< td=""><td>ter></td><td><character></character></td><td><char< td=""><td>cacter></td><td></td></char<></td></charac<></td></charact<></td></charact<></td></rle<>	>	<charact< td=""><td>er></td><td><charact< td=""><td>ter></td><td><charac< td=""><td>ter></td><td><character></character></td><td><char< td=""><td>cacter></td><td></td></char<></td></charac<></td></charact<></td></charact<>	er>	<charact< td=""><td>ter></td><td><charac< td=""><td>ter></td><td><character></character></td><td><char< td=""><td>cacter></td><td></td></char<></td></charac<></td></charact<>	ter>	<charac< td=""><td>ter></td><td><character></character></td><td><char< td=""><td>cacter></td><td></td></char<></td></charac<>	ter>	<character></character>	<char< td=""><td>cacter></td><td></td></char<>	cacter>	
3	4	10619695	-	*	Samp	le2		Α		G	coding, coding	528	34,5347	
4	Х	15838366	3	*	Samp	le1		С		Α	coding		864	
		p	AA_ref		AA_alt	Со	don_ref	Co	odon_alt		Consequ	ience		Gene
	<characte< td=""><td>er> <char< td=""><td>cter></td><td><ch< td=""><td>aracter></td><td><cha< td=""><td>racter></td><td><cha< td=""><td>aracter></td><td></td><td><charac< td=""><td>cter></td><td><charac< td=""><td>ter></td></charac<></td></charac<></td></cha<></td></cha<></td></ch<></td></char<></td></characte<>	er> <char< td=""><td>cter></td><td><ch< td=""><td>aracter></td><td><cha< td=""><td>racter></td><td><cha< td=""><td>aracter></td><td></td><td><charac< td=""><td>cter></td><td><charac< td=""><td>ter></td></charac<></td></charac<></td></cha<></td></cha<></td></ch<></td></char<>	cter>	<ch< td=""><td>aracter></td><td><cha< td=""><td>racter></td><td><cha< td=""><td>aracter></td><td></td><td><charac< td=""><td>cter></td><td><charac< td=""><td>ter></td></charac<></td></charac<></td></cha<></td></cha<></td></ch<>	aracter>	<cha< td=""><td>racter></td><td><cha< td=""><td>aracter></td><td></td><td><charac< td=""><td>cter></td><td><charac< td=""><td>ter></td></charac<></td></charac<></td></cha<></td></cha<>	racter>	<cha< td=""><td>aracter></td><td></td><td><charac< td=""><td>cter></td><td><charac< td=""><td>ter></td></charac<></td></charac<></td></cha<>	aracter>		<charac< td=""><td>cter></td><td><charac< td=""><td>ter></td></charac<></td></charac<>	cter>	<charac< td=""><td>ter></td></charac<>	ter>
3	1762,17	783	I,I		V,V		ATA,ATA		GTA,GTA	nons	synonymous, nons	syn	TET2	TET2
4	2	288	N		K		AAC		AAA		nonsynony	mous	2	RSR2

```
DΡ
                                                                                             VAF
                                                                                                    BQ_REF
         GeneID TranscriptID
                                    GATK
                                            VarScan
                                                        Nr_Ref
                                                                   Nr_Alt
    <character>
                  <character> <numeric> <numeric> <numeric> <numeric> <numeric> <numeric>
                                                                                      <numeric> <numeric>
  3 54790,54790
                  18308,18309
                                                          1268
                                                                     1283
                                                                                2551 0.50294002
                                                                                                   38.6680
                                        1
                                                  1
  4
           8233
                         75467
                                      NA
                                                  1
                                                          1991
                                                                       31
                                                                                3630 0.00853994
                                                                                                   37.4591
       BQ_ALT Nr_Ref_fwd Nr_Alt_fwd
                                          DP_fwd
                                                   VAF_fwd Nr_Ref_rev Nr_Alt_rev
                                                                                       DP_rev
                                                                                                 VAF_rev
                <numeric>
                           <numeric> <numeric> <numeric>
                                                             <numeric>
                                                                         <numeric> <numeric> <numeric>
    <numeric>
                                  469
  3
      38.8145
                      428
                                             897
                                                  0.522854
                                                                    840
                                                                                814
                                                                                          1654
                                                                                                0.492140
      18,0000
                                                  0.000000
  4
                      839
                                    0
                                            1507
                                                                   1152
                                                                                 31
                                                                                         2123
                                                                                                0.014602
  seqinfo: 3 sequences from an unspecified genome; no seqlengths
 characteristics <- determineCharacteristics(output_folder, filtered,</pre>
                                                 predict = "Provean")
> characteristics
GRanges object with 2 ranges and 15 metadata columns:
    seqnames
                 ranges strand
                                     SampleID
                                                        Ref
                                                                     Alt
                                                                                dbSNP dbSNP_MAF
                                                                                                  G1000 AF
       <Rle> <IRanges>
                         <Rle>
                                  <character> <character> <character>
                                                                         <character> <logical>
                                                                                                 <numeric>
           4 106196951
                                      Sample2
                                                                           rs2454206
  3
                                                                       G
                                                                                            < NA >
                                                                                                      0.23
                                                          A
  4
              15838366
                                                          C
                                                                                 <NA>
                                                                                                      0.44
           Х
                                      Sample1
                                                                       A
                                                                                            <NA>
      ExAC AF
                          CosmicID Cosmic_Counts Prediction
                  GAD AF
                                                                     Score
                                                                                              с.
    <numeric> <numeric> <logical>
                                         <logical> <character> <numeric>
                                                                                    <character>
  3
         0.27
                    0.30
                               <NA>
                                              < NA >
                                                        Neutral
                                                                    -0.061 \text{ c.} 5284A>G,c.5347A>G
  4
         0.47
                    0.47
                               <NA>
                                              <NA>
                                                           <NA>
                                                                        NΑ
                                                                                       c.864C>A
           c.complement
                                          р.
             <character>
                                <character>
  3 c.5284T>C,c.5347T>C p.I1762V,p.I1783V
  4
                c.864G>T
                                    p.N288K
```

seqinfo: 3 sequences from an unspecified genome; no seqlengths

2.7 7th analysis step: Final filtration

In the 7th analysis step, the final filtration according to the appreci8-algorithm is performed.

The function finalFiltration performs the final filtration according to the appreci8-algorithm. The previously determined characteristics of the calls are evaluated and an automatic categorization of the calls is performed. Possible categories are: Probably true or Hotspot, Polymorphism and Artifact. Available arguments are:

- output_folder The folder to write the output files into. If an empty string is provided, no files are written out.
- frequency_calls A GRanges object. The GRanges object contains one call per line. EvaluateCovAndBQ()-output can directly be taken as input.
- database_calls A GRanges object. The GRanges object contains one call per line. EvaluateCharacteristics()-output can directly be taken as input.

- combined_calls A GRanges object. The GRanges object contains one call per line. CombineOutput ()-output can directly be taken as input.
- damaging_safe Threshold for the impact prediction score, identifying reliably damaging calls. For example, if Provean has been selected as the impact prediction tool, the internal threshold of the tool is -2.5. However, calls with a score of -2.51 are less likely to have a correct "Damaging" prediction compared to calls with a score of -12.0. A threshold of -5.0 for damaging_safe marks all calls with a score below -5.0 as reliably damaging.
- tolerated_safe Threshold for the impact prediction score, identifying reliably tolerated calls. For example, if Provean has been selected as the impact prediction tool, the internal threshold of the tool is -2.5. However, calls with a score of -2.49 are less likely to have a correct "Neutral" prediction compared to calls with a score of +7.0. A threshold of -1.0 for tolerated_safe marks all calls with a score above -1.0 as reliably tolerated.
- primer Optional: Data frame containing primer positions in bed-format, i.e. 1st column chromosome, 2nd column 0-based start position, 3rd column 1-based end position (default: NA).
- hotspots Optional: Data.frame containing expected/hotspot mutations: 1st column gene name (e.g. ASXL1), 2nd column mutation on AA level (e.g. G12S or E628fs or I836del or G12 if any AA change at this position is considered an expected/hotspot mutation), 3rd column minimum VAF or NA (default: NA).
- overlapTools Vector of strings containing the names of variant calling tools that, should a call be overlappingly reported by these tools, will be rewarded.
- nrsamples Threshold for the number of samples. Shold a variant be detected in more than the threshold defined by nrsamples, it will be interpreted as possible evidence for an artifact (default: 3).
- dp Minimum depth that is required for a call (default: 50).
- nr_alt Minimum number of reads carrying the alternate allele that is required for a call (default: 20).
- vaf Minimum VAF that is required for a call (default: 0.01).
- bq Minimum base quality that is required for a call (default: 15; values above 44 not recommended).
- bq_diff Maximum difference in basequality between reference and alternative that is allowed for a call (default: 7).
- detectedLow Value added to the artifact score if more than nrsamples number of samples feature the same call (default: 2).
- detectedHigh Value added to the artifact score if more than 50% of the samples analyzed (if more than 1 sample is analyzed) feature the same call, which is not an expected/hotspot mutation (default: 2).

- isIndel Value added to the artifact score if the call is an indel (default: 1).
- isIndelVAF Value added to the artifact score if the call is an indel and the VAF is below 0.05 (default: 1).
- detectedLowVAF Value added to the artifact score if more than nrsamples number of samples feature the same call and the VAF is above 0.85 (default: 2).
- noPrimerP Value added to the artifact score if no primer information is provided or no primer is located at this position and the p value (Fisher's Exact Test evaluating forward and reverse reads to determine strandbias) is below 0.001 (default: 1).
- primerPalt Value added to the artifact score if no primer information is provided or no primer is located at this position and the p value (Fisher's Exact Test evaluating forward and reverse reads to determine strandbias) is below 0.001 and Nr_Alt_fwd is at least Nr_Alt/2 and Nr_Alt_rev is at least Nr_Alt/1 (default: -1).
- noPrimerPFwd Value added to the artifact score if no primer information is provided or no primer is located at this position and the p value (Fisher's Exact Test evaluating forward and reverse reads to determine strandbias) is at least 0.001 and Nr_Ref_fwd is at least (DP-Nr_Alt)/2 and Nr Alt fwd is below 3 (default: 1).
- primerPFwd Value added to the artifact score if no primer information is provided or no primer is located at this position and the p value (Fisher's Exact Test evaluating forward and reverse reads to determine strandbias) is below 0.001 and Nr_Ref_fwd is below (DP-Nr_Alt)/2 and Nr_Alt_fwd is below 3 (default: -1).
- noPrimerPRev Value added to the artifact score if no primer information is provided or no primer is located at this position and the p value (Fisher's Exact Test evaluating forward and reverse reads to determine strandbias) is at least 0.001 and Nr_Ref_rev is at least (DP-Nr_Alt)/2 and Nr Alt rev is below 3 (default: 1).
- primerPRev Value added to the artifact score if no primer information is provided or no primer is located at this position and the p value (Fisher's Exact Test evaluating forward and reverse reads to determine strandbias) is below 0.001 and Nr_Ref_rev is below (DP-Nr_Alt)/2 and Nr_Alt_rev is below 3 (default: -1).
- primerLocation Value added to the artifact score if a call is located at a position where a primer is located (default: -1).
- vafLow Value added to the artifact score if the VAF is below 0.02 (default: 2).
- databaseVAF Value added to the artifact score if the VAF is below 0.10 and the variant was not found in any database (default: 1).
- databaseHigh Value added to the artifact score if the variant was detected in at least 50% of all samples, but not found in any database (default: 1).
- predictionSafe Value added to the artifact score if the variant has a reliable damaging prediction (default: -1).

- predictionVAF Value added to the artifact score if the variant has a reliable tolerated prediction and a VAF below 0.35 or between 0.65 and 0.85 (default: 1).
- nrcaller4 Intermediate number of callers that overlappingly reported a variant (default: 4).
- reward4 Value added to the artifact score if a variant is overlappingly reported by nrcaller4 callers (default: -1).
- nrcaller5 High number of callers that overlappingly reported a variant (default: 5).
- reward5 Value added to the artifact score if a variant is overlappingly reported by nrcaller5 callers (default: -1).
- nrcaller6 Very high number of callers that overlappingly reported a variant (default: 6).
- reward6 Value added to the artifact score if a variant is overlappingly reported by nrcaller6 callers (default: -1).
- oneCaller Value added to the artifact score if a variant is only reported by one caller (default: 1).
- BQ_AltMean Value added to the artifact score if the base quality of a variant is below mean(BQ_Alt)-3*(BQ_Alt) over all variants (default: 4).
- knownHotspot Value added to the artifact score if a variant is an expected/hotspot mutation (default: -3).
- overlapReward Value added to the artifact score if a variant is overlappingly reported by the tools defined in overlapTools (default: -3).
- artifactThreshold Threshold for the artifact score, i.e. variants with a score below this threshold will be categorized as "probably true" (default: 0).
- polyDetected Value added to the polymorphism score if more than nrsamples number of samples feature the same call (default: 1).
- polyDetectedOnce Value added to the polymorphism score if a variant is only detected in one sample (default: -1).
- polyDatabasesPolyLow Intermediate number of polymorphism databases that have information on a variant (default: 2).
- polyDatabasesPolyLowReward Value added to the polymorphism score if a variant is present in at least polyDatabasesPolyLow databases (default: 1).
- polyDatabasesPolyHigh High number of polymorphism databases that have information on a variant (default: 4).
- polyDatabasesPolyHighReward Value added to the polymorphism score if a variant is present in at least polyDatabasesPolyHigh databases (default: 1).
- polyDatabasesMut Critical number of mutation databases that have information on a variant (default: 2).

- polyDatabasesMutReward Value added to the polymorphism score if a variant is present in at least polyDatabasesMut databases (default: -1).
- polyNoDatabase Value added to the polymorphism score if a variant is not present in any polymorphism database (default: -1).
- polyDatabases High number of databases that have information on a variant (default: 6).
- polyDatabasesReward Value added to the polymorphism score if a variant is present in at least polyDatabases databases (default: 1).
- polyEffect Value added to the polymorphism score if a variant is not a frameshift variant, not a stop gain variant and not a stop lost variant (default: 1).
- polyVAF Value added to the polymorphism score if the VAF of a variant is between 0.65 and 0.35 or above 0.85 (default: 1).
- polyPrediction Value added to the polymorphism score if the variant has a reliable tolerated prediction (default: 1).
- polyPredictionEffect Value added to the polymorphism score if a variant has a reliable damaging prediction or is a stop gain variant or is a stop lost variant (default: -1).
- polyCosmic Critical number of COSMIC entries (default: 100).
- polyThresholdCritical Threshold for the polymorphism score if the number of COSMIC entries is not critical, i.e. variants with at least this score will be categorized as "polymorphism" (default: 2).
- polyThreshold Threshold for the polymorphism score if the number of COSMIC entries is critical, i.e. variants with at least this score will be categorized as "polymorphism" (default: 3).
- PolymorphismVAF10 Value added to the artifact score if a variant is a "polymorphism" based on the polymorphism score, but the VAF is below 0.10 (default: 5).
- PolymorphismVAF20 Value added to the artifact score if a variant is a "polymorphism" based on the polymorphism score, but the VAF is below 0.20 (default: 2).
- PolymorphismFrame Value added to the artifact score if a variant is a "polymorphism" based on the polymorphism score, but it is a frameshift variant (default: 2).

Final filtration consists of several steps:

- Frequency and base quality are re-considered. Stricter thresholds compared to evaluateCovAndBQ can be defined.
- 2. Samples with the same call are considered. Counting is based on the normalized and annotated calls, not on the coverage- and base quality filtered calls.
- 3. Samples with a call at the same position are considered (e.g. A>AG at pos 1 in sample 1 and A>AGG at pos 1 in sample 2 are reported as 2 calls at the same position). Counting is based on the normalized and annotated calls, not on the coverage- and base quality filtered calls.

- 4. Background information is considered. The number of samples with the same variant in the coverage- and base quality filtered data are considered.
- 5. Number of databases is considered. Dependent on the previously selected number of databases. The presence of a variant in mutation- and polymorphism databases is evaluated.
- 6. VAF in relation to a predicted effect is considered. Variants are marked if their VAF is typical of polymorphisms and their predicted effect is "tolerated".
- 7. VAF in relation to the number of samples is considered. Variants are marked if they are detected in more than nrsamples samples and the VAF is at least 0.85 in more than 90% of these samples.
- 8. Strandbias is considered. Fisher's Exact Test is performed to analyze the relation between forward-reverse and reference-alternate allele.
- 9. Hotspot list if any is provided is considered. Variants are marked if they are present on the hotspot list.
- 10. Final filtration is performed. The artifact- and the polymorphism score are calculated. On the basis of these two scores, a call is either classified as a probably true/hotspot call, polymorphism or artifact.

A GRanges object is returned containing all calls with a predicted category. Categories can be: probably true, hotspot, polymorphism or artifact. Reported metadata columns are: SampleID, Ref, Alt, Gene, GeneID, TranscriptID, Location, Consequence, c. (variant on cDNA level containing position and variant), c.complement (complement of the variant; if c. is c.5284A>G, then c.complement is c.5284T>C), p. (variant on protein level containing position and amino acids), Codon_ref, Codon_alt, Nr_Ref, Nr_Alt, DP, VAF, Caller1 to CallerX (dependent on the number of callers that is evaluated), dbSNP (containing the rs-ID), dbSNP_MAF, G1000_AF, ExAC_AF, GAD_AF, CosmicID, Cosmic_Counts (number of Cosmic entries), ClinVar, Prediction (damaging or neutral), Score (on the basis of the selected prediction tool), BQ_REF, BQ_ALT, Nr_Ref_fwd, Nr_Alt_fwd, DP_fwd, VAF_rev, Nr_Ref_rev, Nr_Alt_rev, DP_rev, VAF_rev, strandbias, nr_samples (number of samples with the same variant), nr_samples_similar (number of samples with a variant at the same position), Category.

If an output folder is provided, the output is saved as Results_Final.txt. Additionally, Results_Final.xlsx is saved, containing three sheets: Mutations, Polymorphisms and Artifacts.

```
> output_folder <- ""
> #Input:
```

> filtered

GRanges object with 2 ranges and 30 metadata columns:

```
segnames
              ranges strand
                                   SampleID
                                                     Ref
                                                                  Alt
                                                                           Location
                                                                                               с.
     <Rle> <IRanges>
                       <Rle>
                               <character> <character> <character>
                                                                        <character> <character>
3
         4 106196951
                                                                                       5284,5347
                                    Sample2
                                                       A
                                                                    G coding, coding
4
           15838366
                                    Sample1
                                                       С
                                                                                             864
                                                                             coding
                                 AA alt
                                          Codon ref
                                                       Codon_alt
                                                                             Consequence
                    AA ref
                                                                                                  Gene
           р.
  <character> <character> <character> <character> <character>
                                                                             <character> <character>
```

```
V,V
    1762,1783
                      I,I
                                           ATA, ATA
                                                       GTA, GTA nonsynonymous, nonsyn..
3
                                                                                          TET2, TET2
          288
                        N
                                     K
                                               AAC
                                                           AAA
                                                                         nonsynonymous
                                                                                              ZRSR2
                                                                            DP
       GeneID TranscriptID
                                 GATK
                                                              Nr_Alt
                                                                                      VAF
                                                                                              BQ_REF
                                        VarScan
                                                   Nr_Ref
  <character> <character> <numeric> <numeric> <numeric> <numeric> <numeric> <numeric>
                                                                                <numeric> <numeric>
3 54790,54790
               18308,18309
                                                     1268
                                                                1283
                                                                          2551 0.50294002
                                                                                             38.6680
                                   1
                                              1
                                                                          3630 0.00853994
         8233
                     75467
                                   NA
                                              1
                                                     1991
                                                                  31
                                                                                             37.4591
     BQ_ALT Nr_Ref_fwd Nr_Alt_fwd
                                      DP_fwd
                                              VAF_fwd Nr_Ref_rev Nr_Alt_rev
                                                                                 DP_rev
                                                                                           VAF_rev
  <numeric>
             <numeric> <numeric> <numeric> <numeric>
                                                        <numeric> <numeric> <numeric> <numeric>
    38.8145
                               469
                                              0.522854
                                                              840
                                                                                   1654 0.492140
                   428
                                         897
                                                                          814
                   839
   18.0000
                                 0
                                        1507 0.000000
                                                              1152
                                                                           31
                                                                                   2123 0.014602
```

seqinfo: 3 sequences from an unspecified genome; no seqlengths

> characteristics

GRanges object with 2 ranges and 15 metadata columns: ranges strand SampleID dbSNP dbSNP_MAF G1000_AF Ref Alt <Rle> <IRanges> <Rle> | <character> <character> <character> <character> <logical> <numeric> 4 106196951 * Sample2 rs2454206 0.23 3 A G < NA >X 15838366 Sample1 С Α <NA> <NA> 0.44 ExAC AF GAD_AF CosmicID Cosmic_Counts Prediction Score с. <logical> <character> <numeric> <numeric> <numeric> <logical> <character> 3 0.27 0.30 < NA >< NA >Neutral -0.061 c. 5284A > G, c. 5347A > G4 0.47 0.47 <NA> <NA> <NA> NAc.864C>A c.complement р. <character> <character> 3 c.5284T>C,c.5347T>C p.I1762V,p.I1783V

seqinfo: 3 sequences from an unspecified genome; no seqlengths

p.N288K

> combined

GRanges object with 3 ranges and 16 metadata columns:

c.864G>T

41041	1600 obje.	00 111	011 0 1	. 411600	ina io modada	. oa oo 1	· ammi					
	seqnames	r	anges	strand	SampleI	D	Re	f	Alt	;	1	Location
	<rle></rle>	<ira:< td=""><td>nges></td><td><rle></rle></td><td> <character< td=""><td>> <cha< td=""><td>aracter</td><td>> <charac< td=""><td>cter></td><td>,</td><td><ch:< td=""><td>aracter></td></ch:<></td></charac<></td></cha<></td></character<></td></ira:<>	nges>	<rle></rle>	<character< td=""><td>> <cha< td=""><td>aracter</td><td>> <charac< td=""><td>cter></td><td>,</td><td><ch:< td=""><td>aracter></td></ch:<></td></charac<></td></cha<></td></character<>	> <cha< td=""><td>aracter</td><td>> <charac< td=""><td>cter></td><td>,</td><td><ch:< td=""><td>aracter></td></ch:<></td></charac<></td></cha<>	aracter	> <charac< td=""><td>cter></td><td>,</td><td><ch:< td=""><td>aracter></td></ch:<></td></charac<>	cter>	,	<ch:< td=""><td>aracter></td></ch:<>	aracter>
2	2	2546	69504	*	Sample	1		G	T	coding	,codin	g,coding
3	4	10619	96951	*	Sample	2		A	G	ł	coding	g,coding
4	X	158	38366	*	Sample	1		С	A	L		coding
		с.		р.	AA_ref	I	AA_alt	Codon_1	ref	Codon_	alt	
	<charac< td=""><td>cter></td><td><char< td=""><td>cacter></td><td><character></character></td><td><chara< td=""><td>acter></td><td><characte< td=""><td>er> <</td><td>charact</td><td>er></td><td></td></characte<></td></chara<></td></char<></td></charac<>	cter>	<char< td=""><td>cacter></td><td><character></character></td><td><chara< td=""><td>acter></td><td><characte< td=""><td>er> <</td><td>charact</td><td>er></td><td></td></characte<></td></chara<></td></char<>	cacter>	<character></character>	<chara< td=""><td>acter></td><td><characte< td=""><td>er> <</td><td>charact</td><td>er></td><td></td></characte<></td></chara<>	acter>	<characte< td=""><td>er> <</td><td>charact</td><td>er></td><td></td></characte<>	er> <	charact	er>	
2	697,1264	,1264	233,4	122,422	L, L, L		M,M,M	CTG,CTG,	CTG A	TG, ATG,	ATG	
3	5284	,5347	176	32,1783	I,I		V,V	ATA,	ATA	GTA,	GTA	
4		864		288	N		K	I	AAC		AAA	
		Co	nseque	ence		Gene		${\tt GeneID}$		Transc	riptID	GATK
		<c1< td=""><td>haract</td><td>er></td><td><chara< td=""><td>cter></td><td><ch< td=""><td>aracter></td><td></td><td><char< td=""><td>acter></td><td><numeric></numeric></td></char<></td></ch<></td></chara<></td></c1<>	haract	er>	<chara< td=""><td>cter></td><td><ch< td=""><td>aracter></td><td></td><td><char< td=""><td>acter></td><td><numeric></numeric></td></char<></td></ch<></td></chara<>	cter>	<ch< td=""><td>aracter></td><td></td><td><char< td=""><td>acter></td><td><numeric></numeric></td></char<></td></ch<>	aracter>		<char< td=""><td>acter></td><td><numeric></numeric></td></char<>	acter>	<numeric></numeric>
2	nonsynony	ymous	,nonsy	n DNN	MT3A,DNMT3A,D	NMT3A	1788,1	788,1788	1080	8,10809	,10810	1

```
TET2, TET2
                                                    54790,54790
                                                                      18308,18309
  3 nonsynonymous, nonsyn...
                                                                                           1
     nonsynonymous
                                          ZRSR2
                                                           8233
                                                                            75467
                                                                                          NA
      VarScan
    <numeric>
  2
          NΑ
  3
           1
  4
            1
  seqinfo: 3 sequences from an unspecified genome; no seqlengths
> final<-finalFiltration(output_folder, frequency_calls = filtered,
                         database_calls = characteristics,
                         combined_calls = combined, damaging_safe = -3,
                         tolerated_safe = -1.5, overlapTools = c("VarScan"),
                         bq_diff = 20, vaf = 0.001)
> final
GRanges object with 2 ranges and 43 metadata columns:
   seqnames
                ranges strand
                                   SampleID
                                                     Ref
                                                                 Alt
                                                                            Gene
                                                                                       GeneID
                       <Rle> | <character> <character> <character> <character> <character>
       <Rle> <IRanges>
  3
           4 106196951
                            *
                                    Sample2
                                                                  G
                                                                       TET2, TET2 54790, 54790
                                                       Α
           X 15838366
                            *
                                    Sample1
                                                       C
                                                                           ZRSR2
                                                                                        8233
   TranscriptID
                      Location
                                          Consequence
                                                                        с.
                                                                                   c.complement
                                          <character>
     <character>
                   <character>
                                                                                   <character>
                                                               <character>
    18308,18309 coding,coding nonsynonymous,nonsyn.. c.5284A>G,c.5347A>G c.5284T>C,c.5347T>C
           75467
                        coding
                                        nonsynonymous
                                                                  c.864C>A
                                                                                       c.864G>T
                        Codon_ref
                                    Codon_alt
                                                  Nr_Ref
                                                            Nr_Alt
                                                                          DP
                                                                                    VAF
                                                                                              GATK
                   р.
          <character> <character> <numeric> <numeric> <numeric> <numeric> <numeric> <numeric>
                                                              1283
                                                                        2551 0.50294002
  3 p.I1762V,p.I1783V
                          ATA, ATA
                                      GTA, GTA
                                                    1268
                                                                        3630 0.00853994
                                                                                                NA
             p.N288K
                              AAC
                                          AAA
                                                    1991
                                                                31
      VarScan
                 Called
                              dbSNP dbSNP_MAF G1000_AF
                                                           ExAC_AF
                                                                      GAD_AF CosmicID Cosmic_Counts
    <numeric> <numeric> <character> <logical> <numeric> <numeric> <numeric> <logical>
                                                                                            <logical>
  3
                          rs2454206
                                          <NA>
                                                    0.23
                                                              0.27
                                                                        0.30
                                                                                   <NA>
            1
                      2
                                                                                                 <NA>
  4
                      1
                                          <NA>
                                                    0.44
                                                                        0.47
                                                                                   <NA>
            1
                               < NA >
                                                              0.47
                                                                                                 <NA>
     Prediction
                    Score
                             BQ_REF
                                       BQ_ALT Nr_Ref_fwd Nr_Alt_fwd
                                                                        DP_fwd
                                                                                VAF_fwd Nr_Ref_rev
                                                                                           <numeric>
    <character> <numeric> <numeric> <numeric> <numeric>
                                                          <numeric> <numeric> <numeric>
  3
        Neutral
                   -0.061
                            38.6680
                                      38.8145
                                                      428
                                                                 469
                                                                           897 0.522854
                                                                                                 840
                                      18.0000
                                                      839
                                                                          1507
                                                                                                1152
           < NA >
                       NA
                            37.4591
                                                                   0
                                                                                0.000000
   Nr Alt rev
                  DP_rev
                           VAF_rev strandbias nr_samples nr_samples_similar
                                                                                   Category
     <numeric> <numeric> <numeric>
                                     <numeric> <integer>
                                                                    <integer> <character>
  3
           814
                    1654 0.492140 1.46693e-01
                                                                            1 Polymorphism
                                                         1
                    2123 0.014602 6.69655e-08
  4
            31
                                                         1
                                                                            1 Artifact (4)
```

seqinfo: 3 sequences from an unspecified genome; no seqlengths

3 appreci8R - GUI

appreci8R combines and filters the output of different variant calling tools according to the 'appreci8'-algorithm. The whole analysis-pipeline consists of seven steps. Individual functions for executing these steps are available. However, an additional user-interface is available that enables easy execution of the analysis-pipeline, restarting parts of the pipeline from checkpoints, changing parameters, exporting and importing the configuration and evaluation of the results. The user-interface is opend by executing the function appreci8Rshiny (no arguments have to be provided).

The user-interface is structured by tabs: Analysis, Overview results and Detailed results.

- Analysis: The tab is essential for performing the whole analysis. It is further separated into 10 additional tabs:
 - Important notes: General information on prerequisites for performing the analysis and current limitations (only hg19).
 - 0. Preparations: Defining the general output folder and the varaint caller input. GATK, Platypus, VarScan, FreeBayes, LoFreq, SNVer, SamTools and VarDict are available as these make up the classical appreci8-pipeline. The tools can be added or removed. For every tool the following parameters have to be defined: the folder containing the variant calling results, the output file type (vcf or txt; as vcf is a standardized format, the columns containing chr, pos, ref and alt are assumed to be fixed; when selecting txt, the columns containing chr, pos, ref and alt have to be defined), a possible suffix for naming the output, if MNVs are reported (AG>CT instead of A>C and G>T), if SNVs and indels are reported in one file (if not, what is the suffix for the SNV files and what is the suffix for the indel files?), and how are indels reported. In addition to the eight tools, up to 5 additional tools can be added (select the number and click Go). In addition to the previous parameters, a name can be defined for each additionally added tool.
 - 1. Target filtration: Upload your target region (bed file; chromosome definition without 'chr').
 - 2. Normalization
 - 3. Annotation: Annotates the variants and filters them according to the selected locations (coding, intron, threeUTR, fiveUTR, intergenic, spliceSite, promoter) and consequences (synonymous, nonsynonymous, frameshift, nonsense, not translated) of interest.
 - 4. Combine output
 - 5. Evaluate Coverage and BQ: Evaluate coverage and base quality by analyzing the bam files. Calls with insufficient coverage and or base quality are filtered. The following parameters have to be defined: the folder containing the bam- and bai files, minimum coverage, minimum number of reads carrying the alternate allele, minimum VAF, minimum base quality and the maximum difference in base quality between reference and alternative.
 - 6. Extended Set of Characteristics: Select the databases you would like to query (dbSNP, 1000Genomes, ExAC, Genome Aggregation Database, COSMIC, ClinVar) and the source you would like to use for impact prediction (SIFT, Provean, PolyPhen2).

- 7. Final Filtration: Adjust parameters and define input for the final filtration. A bed file containing the primer positions can be uploaded (optional). A txt file containing expected/hotspot mutations (defined via gene, mutation on AA level, and an optional minimum VAF) can be uploaded (optional). Stricter thresholds for coverage and base quality can be defined (see parameters for 5. Evaluate Coverage and BQ). The number of samples making a call striking can be defined. Dependent on the previously selected source for impact prediction (SIFT, Provean, PolyPhen2), a threshold can be defined to identify reliable damaging predictions and reliable tolerated prediction. Furthermore, the default scoring for the artifact- and the polymorphism score can be changed (just recommended for experts).
- Action: The complete analysis can be executed (Start complete analysis). The availability of checkpoints can be checked (Check for possible checkpoints). If checkpoints are available, one can be selected and the analysis can be started from the selected checkpoint. The defined configuration can be exported (Export current configuration) as well as imported (Import configuration). Press Done if you are done with your analysis.
- Overview results: The tab provides an overview of the results. It is further separated into 5 additional tabs:
 - Log: The log of the analysis is displayed here. In case of errors, e.g. input not being available, these messages are also reported in the log.
 - Raw Calls: The raw number of calls per sample and per caller are reported.
 - Calls On Target: The number of calls on target per sample and per caller are reported.
 - Annotaed Calls: The number of annotated (and filtered according to annotation) calls per sample and per caller are reported.
 - Filtered Calls: The number of coverage and base quality filtered calls per sample is reported.
- Detailed results: The tab provides detailed information on the results. It is further separated into 3 additional tabs:
 - Mutations: Variant calls classified as likely true mutations.
 - Polymorphisms: Variant calls classified as likely polymorphisms.
 - Artifacts: Variant calls classified as likely artifacts.

No value is returned. Output files for the different analysis steps are automatically saved.

> appreci8Rshiny()

References

- Morgan, Η Pagès, V Obenchain, andΝ Hayden. Rsamtools: Bicall(bcf) $_{\mathrm{file}}$ alignment (bam), fasta, variant and tabix import. http://bioconductor.org/packages/release/bioc/html/Rsamtools.html.
- [2] V Obenchain, M Lawrence, V Carey, S Gogarten, P Shannon, and M Morgan. Variantannotation: a bioconductor package for exploration and annotation of genetic variants. *Bioinformatics*, 30(14), 2014.
- [3] S Sandmann, M Karimi, A de Graaf, C Rohde, S Göllner, J Varghese, J Ernsting, G Walldin, B van der Reijden, C Müller-Tidow, L Malcovati, E Hellström-Lindberg, J Jansen, and M Dugas. appreci8: A pipeline for precise variant calling integrating 8 tools, 2018. https://doi.org/10.1093/bioinformatics/bty518.