JACUSA manual Version 1.0

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1 Introduction

JAVA framework for accurate SNV assessment (JACUSA) is a one-stop solution to detect single nucleotide variants (SNVs) from comparing matched sequencing samples. Robust identification has proven to be a daunting task due to artefacts specific for NGS-data and employed mapping strategies. We implement various feature filters that reduce the number of false positives. JACUSA employs a window-based approach to traverse provided BAM files featuring highly parallel processing. JACUSA has been extensively evaluated and optimized to identify RNA editing sites in RNA-DNA and RNA-RNA sequencing samples. JACUSA requires an operating JAVA environment and uses sorted and indexed BAM files as input.

2 Download

Download the latest version of JACUSA from . Check the source in the repository on GiTHub.

2.1 Installation and requirements

JACUSA does not need any configuration but needs a correctly configured Java environment. We developed and tested JACUSA with Java v1.7. If you encounter any Java related problems please consider to change to Java v1.7.

2.2 Sample in silico data

Download sample data that we used for the development of JACUSA. You can choose between different setups and species where the later greatly influences the data size and running time to detect variants. The gDNA VS cDNA represents the typical data setup that is encountered in detection of RNA editing sites via comparing genomic and transcriptomic sequencing reads. In this setup, variants have been only imputed to the cDNA BAM file. The cDNA VS cDNA data setup can be interpreted as representing allele specific expression of single variants or differential RNA editing. In this setup, variants with pairwise different base frequency have been imputed into both cDNA BAM files. Additionally, to make the identification of variants more challenging SNPs with pairwise similar base frequencies have been included to both BAM files. This sites should not be identified as true positive sites. gDNA data has been simulate with art¹ and cDNA reads have been simulated with flux². Read simulations have been restricted to the corresponding first chromosome of the respective species. Sample data is available for C. elegans ce10 and Homo sapien hg19. Each archive consists of:

gDNA.bam, cDNA.bam BAM files: gDNA.bam and cDNA OR cDNA_1.bam and cDNA 2.bam

snps.txt Only available for cDNA VS cDNA. Coordinates of imputed SNPs. In both BAM files matching SNPs have the same target frequency but different effective or sampled frequency. The shape parameter determines how much the sampled frequency will deviate from the target frequency in each BAM file. The suffixes: _cdna_1 and _cdna_2 correspond to the respective BAM file

variants.txt Coordinates of imputed variants and their target and sample frequencies

Available sample data organized by data type and species:

- hg19 chr1 gDNA VS cDNA.tar.gz
- hg19 chr1 cDNA VS cDNA.tar.gz

 $^{^{1}\}mathrm{art}$

²flux simulator

3 Input

3.1 Alignment files

JACUSA needs sorted and indexed BAM files. BAM is a standardized file format for efficient storage of alignments. Check the manuals for 3 and/or 4 for how to use the respective tool to convert your alignment files to valid JACUSA input BAM.

In the following, commands for SAMtools are presented:

 $\mathbf{SAM} o BAM$ samtools view -Sb mapping.sam > mapping.bam

sort BAM samtools sort mapping.bam mapping.sorted

index BAM samtools index mapping.sorted.bam

It is recommended pre-processing step to remove duplicate reads when identifying variants. Duplicated reads occur mostly due to PCR-artefacts. They are likely to harbour false variants and most statistical test require that reads are sampled independently. In the following, commands for picarf tools are presented:

```
java -jar MarkDuplicates.jar \
  I=mapping.sorted.bam O=dedup_mapping.sorted.bam \
  M=duplication.info
```

Invoke JACUSA with the additional command line option "-F 1024" to filter read that have been marked as duplicates.

3.1.1 Strand information

Depending on the employed sequencing library, JACUSA can use the strand orientation to build pileups. With the command line parameter "-P,-build-pileup <BUILD-PILEUP>" the user can choose from combinations of:

S stranded, and

U unstranded

to define if strand information will be utilized to build pileups. The format of <BUILD-PILEUP> is:

$$x, y: x, y \in \{S, U\}$$

where x corresponds corresponds to the first sample (BAM1_1-n files) and y to the second sample (BAM2_1-m files). The default is to ignore strand information for both samples ("-P U,U") JACUSA will process unstraded single-end and paired-end but strand orientation of base counts needs to be inferred from annotation (GFF). For stranded single-end RNA-Seq distiguish after which strand synthesis the read is sequenced:

- first-strand or
- second-strand

 $^{^3{\}rm SAM tools/BCF tools}$

⁴picard tools

and invert JACUSA output when first-strand library type is used. In order to utilize read orientation in paired end RNA-Seq both fragments must have same orientation f-f or r-r. Otherwise, use unstranded and invert correct orientation from annotation (see 1 for summary). In order to identify RNA editing sites by

Table 1: Summary of supported library types by JACUSA

	$\mathbf{single}\mathbf{-end}$	paired end		
unstranded	infer correct orientation from annotation			
$\operatorname{stranded}$				
first-strand	invert the orienation in	invert the orienation in		
	JACUSA output	JACUSA output (both		
		fragments $must$ have same		
		orientation f-f or r-r)		
second-strand	JACUSA output has cor-	JACUSA output has		
	rect orientation	correct orientation (both		
		fragments must have		
		same orientation f-f or r-r)		

comparing gDNA and stranded RNA-Seq use "-P U,S"

3.2 Traverse BED-like file

Variant detection can be limited to specific regions of the genome or transcriptome. Provide a minimalistic BED-like file to restrict the search to this region(s) or site(s). Remaining region(s) of the BAM files will not be considered.

In the following traverse file, the search is confined to a 100nt region on contig 1 staring at 1,000 and a single site on contig 2 at coordinates 10,000: HINT: Many individual sites will slow down JACUSA. If possible, try to merge nearby sites into contiguous regions and extract specific sites from JACUSA output with bedtools⁵ intersect:

merge sites

bedtools merge -d 500 singular_sites.bed > \
 contigous_regions.bed

run JACUSA

java -jar JACUSA.jar call-2 -b contigous_regions.bed -o
JACUSA.out mapping_1.sorted.bam mapping_2.sorted.bam

extract sites

bedtools intersect -wa -a JACUSA.out -b singular_sites.bed

Table 2: Example of BED-like traverse file

contig	start	end
1	1000	1100
2	10000	10000

⁵bedtools

4 Output

JACUSA writes its output to a user defined file or a pipe. When using multiple threads, JACUSA will create a gzipped temporary file for each allocated thread. Chosen command line parameters and current genomic position are printed to the command prompt. Furthermore, depending on the provided command line parameters, JACUSA will generate a file with sites that have been identified as potential artefacts when "-s" is provided. Currently, JACUSA supports the following output formats, controlled by "-f":

- Default (JACUSA output)
- Variant Call Format (VCF)⁶

The default output format is based on BED6⁷ with additional JACUSA specific columns. The actual number of columns depends on number of provided BAM files.

Table 3: JACUSA default output format										
Column:	1	2	3	4	5	6	7	8	9	10
	1	100	101	variant	8.07	-	0,0,0,6	0,6,0,0	*	*

- (1, 2, 3) contig + start + end 0-based, genomic coordinates of potential variant site
- (4) name Currently, constant string: "variant". This dummy field is to ensure BED6 compatibility
- (5) score Test-statistic $z \in \mathbb{R}$ that indicates the likelihood that this is a true variant. Higher number indicates a higher likelihood for a variant
- (6) strand Possible values are: ".", "+", and "-" which correspond to "unstranded", "positive strand", and "negative strand" respectively. If strand is != ".", then the following base columns will be indicating base counts according to the strand inverted base count if on the "negative strand"
- (7,8) basesIJ The number of base columns depends on the number of BAM files. In basesIJ: I corresponds to sample and J to the respective replicate. Numbers indicate the base count of the following base vector: (A, C, G, T)
- (9) info Additional info for this specific site. Currently, details about the parameter estimation of the Dirichlet-Multinomial can be shown. If nothing provided, the empty field is equal to "*"
- (10) filter_info Relevant, if feature filter(s) X have been provided with "-a X" on the command line. The column will contain a comma-separated list of feature filters that predict this site to be a potential artefact. Possible

⁶VCF file format

 $^{^7\}mathrm{BED}$ file format

values are	e:
Value	Description of potential artefact
D	Variant call in the vicinity of Read Start/End, Intron, and/or
	INDEL position
В	Variant call in the vicinity of Read Start/End
I	Variant call in the vicinity of INDEL position
\mathbf{S}	Variant call in the vicinity of Splice Site
Y	Variant call in the vicinity of homopolymer

M Max allowed alleles exceeded

H "Control" sample contains non-homozygous pileup

d Some pileup exceeds max depth

5 Usage

Calling JACUSA without any arguments will print the available tools which currently are:

```
java -jarJACUSA.jar
call-2 Call variants - two samples
  pileup SAMtools like mpileup for two samples
usage: jacusa.jar [OPTIONS] BAM1_1[,BAM1_2,BAM1_3,...] BAM2_1[,BAM2_2,BAM2_3,...]
 -a,-pileup-filter
                      <PILEUP-
                                   chain of PILEUP-FILTER to apply to pileups:
 FILTER>
                                   D | Filter distance to Read Start/End, In-
                                   tron, and INDEL position.
                                                                   Default:
                                   (D:distance:min_ratio)
                                   S | Filter distance to Splice Site. Default: 6:0.5
                                   (S:distance:min_ratio)
                                   B | Filter distance to Read Start/End. Default:
                                   6:0.5 (F:distance:min ratio)
                                   M | Max allowed alleles per parallel pileup. Default:
                                   Y | Filter wrong variant calls in the vicinity of ho-
                                   mopolymers. Default: 7 (Y:length)
                                   H | Filter non-homozygous pileup/BAM (1 or 2).
                                   Default: none
                                   I | Filter distance to INDEL position. Default: 6:0.5
                                   (I:distance:min ratio)
                                   Separate multiple PILEUP-FILTER with ',' (e.g.:
                                   D,I)
 -b,-bed < BED >
                                   BED file to scan for variants
 -C,-base-config
                        <BASE-
                                   Choose what bases should be considered for variant
 CONFIG>
                                   calling: TC or AG or ACGT or AT.... Default:
                                   ACGT
 -c,-min-coverage
                          <MIN-
                                   filter positions with coverage < MIN-COVERAGE.
 COVERAGE >
                                   Default: 5
                          <MIN-
                                   filter 1 positions with
 -c1,-min-coverage1
                                                              coverage
                                                                             MIN-
 COVERAGE1>
                                   COVERAGE1. Default: 5
                          <MIN-
                                   filter 2 positions with
 \hbox{-}c2, \hbox{-}min\hbox{-}coverage2
                                                                             MIN-
                                                              coverage
 COVERAGE2>
                                   COVERAGE2. Default: 5
```

d man danth (MAY DEDTII)	man non DAM donth Default. 1
-d,-max-depth <max-depth> -d1,-max-depth1 <max-< td=""><td>max per-BAM depth. Default: -1 max per-sample 1 depth. Default: -1</td></max-<></max-depth>	max per-BAM depth. Default: -1 max per-sample 1 depth. Default: -1
DEPTH1>	max per sumple 1 depth. Delautt. 1
-d2,-max-depth2 <max- DEPTH2></max- 	max per-sample 2 depth. Default: -1
-F,-filter-flags <filter-< td=""><td>filter reads with flags FILTER-FLAGS. Default: 0</td></filter-<>	filter reads with flags FILTER-FLAGS. Default: 0
FLAGS>	intel leads with hags libibit-i bitoto. Delaute. 0
-f,-output-format <output-format></output-format>	Choose output format:
	<*> D: Default JACUSA
	< $>$ V: VCF
$-filter NH_1 < \! NH\text{-VALUE} \! >$	Max NH-VALUE for SAM tag NH
$- filter NH_2 < \! NH\text{-VALUE} \! >$	Max NH-VALUE for SAM tag NH
$- filter NM_1 < \! NM\text{-VALUE} \! >$	Max NM-VALUE for SAM tag NM
$- filter NM_2 < \! NM\text{-VALUE} \! >$	Max NM-VALUE for SAM tag NM
-h,-help	Print usage information
-m,-min-mapq $<$ MIN-MAPQ $>$	filter positions with MAPQ < MIN-MAPQ. Default: 20
-m1,-min-mapq1 <min-< td=""><td>filter 1 positions with MAPQ < MIN-MAPQ1. De-</td></min-<>	filter 1 positions with MAPQ < MIN-MAPQ1. De-
MAPQ1>	fault: 20
-m2,-min-mapq2 <min-< td=""><td>filter 2 positions with MAPQ < MIN-MAPQ2. De-</td></min-<>	filter 2 positions with MAPQ < MIN-MAPQ2. De-
m MAPQ2>	fault: 20
-P,-build-pileup <build-< td=""><td>Choose how parallel pileups are build: strand spe-</td></build-<>	Choose how parallel pileups are build: strand spe-
PILEUP>	cific (S) or strand unspecific (U) default: U,U
$\hbox{-p,-threads} < \hbox{THREADS} >$	use $\#$ THREADS. Default: 1
-q,-min-basq $<$ MIN-BASQ $>$	filter positions with base quality $<$ MIN-BASQ. Default: 20
-q1,-min-basq1 < MIN-BASQ1>	filter 1 positions with base quality $<$ MIN-BASQ1. Default: 20
-q2,-min-basq2 $<$ MIN-BASQ2 $>$	filter 2 positions with base quality < MIN-BASQ2. Default: 20
-r,-result-file $<$ RESULT-FILE $>$	results are written to RESULT-FILE or STDOUT
,	if empty
-s, $-separate$	Put feature-filtered results in to a separate file (=
	RESULT-FILE.filtered)
$\hbox{-T,-threshold} < \hbox{THRESHOLD}>$	Filter positions depending on test-statistic
	THRESHOLD default: DO NOT FILTER
-v,-version	Print version information.
-W,-thread-window-size	size of the window used per thread. Default: 100000
<thread-window-size></thread-window-size>	size of the window used for eaching. Mal
-w,-window-size <window- SIZE></window- 	size of the window used for caching. Make sure
DILL	this is greater than the read size and smaller than
	this is greater than the read size and smaller than THREAD-WINDOW-SIZE>. Default: 10000

5.1 SAMtools like mpileup for two samples

See "Call variant - two samples" for details.

6 Identification of RNA editing sites

Use the following command line to identify RNA-DNA differences in BAM files that might give rise to RNA editing sites:

```
java -jar call-2 -o JACUSA.out -s -a H:1 gDNA.bam cDNA.bam
```

Option "-a H:1" ensures that potential polymorphisms in gDNA will be eliminated as artefacts. The number $x \in \{1,2\}$ determines which sample has to be homomorph - in this case: gDNA.bam.

Use the following command line to identify RNA-DNA differences:

```
java -jar call-2 -o JACUSA.out -s cDNA1.bam cDNA2.bam
```

WARNING: If you want to identify RNA-RNA differences make sure NOT to use the filter "-a H:x"! Otherwise, potential valid variants will be filtered out.

7 Used libraries

picard tools