Bamsurgeon: Methods for spike-in mutations on BAM files

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1 Adding SNVs to existing BAM alignments (addsnv)

1.1 Usage

```
usage: addsnv.py [-h] -v VARFILENAME -f BAMFILENAME -r REFFASTA -o OUTBAMFILE
[-s SNVFRAC] [-m MUTFRAC] [-n NUMSNVS] [--nomut] [--det]
[--force]
```

adds SNVs to reads, outputs modified reads as .bam along with mates

optional arguments:

-h, --help show this help message and exit

-v VARFILENAME, --varfile VARFILENAME

Target regions to try and add a SNV, as BED

-f BAMFILENAME, --sambamfile BAMFILENAME

sam/bam file from which to obtain reads

-r REFFASTA, --reference REFFASTA

reference genome, fasta indexed with bwa index -a

stdsw _and_ samtools faidx

-o OUTBAMFILE, --outbam OUTBAMFILE

.bam file name for output

-s SNVFRAC, --snvfrac SNVFRAC

maximum allowable linked SNP MAF (for avoiding

haplotypes) (default = 1)

-m MUTFRAC, --mutfrac MUTFRAC

allelic fraction at which to make SNVs (default = 0.5)

-n NUMSNVS, --numsnvs NUMSNVS

maximum number of mutations to make (default: entire

input)

--nomut dry run

--det deterministic base changes: make transitions only force mutation to happen regardless of nearby SNP or

low coverage

1.2 Description

Single nucleotide changes are introduced to an existing BAM alignment using addsnv.py as described in Figure 4. Input consists of a list of locations where SNVs will be made, a target BAM alignment, and a reference genome indexed using bwa (preferably the same genome used to align the reads in the BAM file). Additionally there are a number of optional arguments, including the option to make deterministic mutations (default is random, deterministic always makes transitions), an option to skip locations if a SNP above a specified MAF is located nearby (avoid interfering with haplotypes), and the MAF of the spike-in SNVs (default 0.5/heterozygous). Output consists of a BAM file containing the spike-in mutations and a log file describing which bases were changed in which reads.

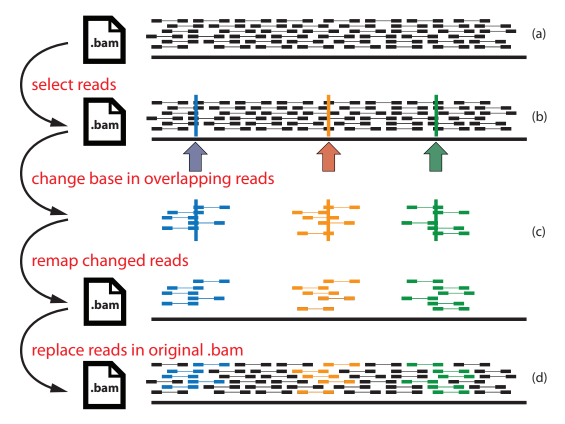


Figure 1: Method for adding SNVs to existing BAM alignments. Starting with the original BAM alignment (a) and a list of positions (b), reads overlapping the chosen positions are selected the target base change is made (c). Altered reads are re-mapped the the reference genome using bwa, and the modified, realigned reads replace the unmodified versions in the original BAM (d).

2 Adding indels and SVs to existing BAM alignments (addsv.py)

2.1 Usage

```
adds SNVs to reads, outputs modified reads as .bam along with mates
optional arguments:
  -h, --help
                        show this help message and exit
  -v VARFILENAME, --varfile VARFILENAME
                        whitespace-delimited target regions to try and add a
                        SNV: chr, start, stop, action, seqfile if
                        insertion, TSDlength if insertion
  -f BAMFILENAME, --sambamfile BAMFILENAME
                        sam/bam file from which to obtain reads
  -r REFFASTA, --reference REFFASTA
                        reference genome, fasta indexed with bwa index -a
                        stdsw _and_ samtools faidx
  -o OUTBAMFILE, --outbam OUTBAMFILE
                        .bam file name for output
  -1 MAXLIBSIZE, --maxlibsize MAXLIBSIZE
                        maximum fragment length of seq. library
  -k KMERSIZE, --kmer KMERSIZE
                        kmer size for assembly (default = 31)
  -s SVFRAC, --svfrac SVFRAC
                        allele fraction of variant (default = 1.0)
  -x EXCLFILE, --excluded EXCLFILE
                        output excluded (e.g. from a deletion) read names to
                        file (default=excluded.txt)
  --maxctglen MAXCTGLEN
                        maximum contig length for assembly - can increase if
                        velvet is compiled with LONGSEQUENCES
                        maximum number of mutations to make
  -n MAXMUTS
  --nomut
                        dry run
  --noremap
                        dry run
  --noref
                        do not perform reference based assembly
```

2.2 Description

--recycle

Small insertions and deletions (indels), and larger structural variants (insertions, deletions, duplications, inversions, and compound variants) are added to existing BAM alignments using addsv.py as described in Figure 5. Input consists of a list of regions where SVs will be made along with a specification of each variant, a target BAM alignment, and a reference genome indexed using bwa. The input mutation list consists of four columns: chromosome, start of region, end of region, and a controlled-vocabulary description of the mutation. A mutation will not be made if the largest contig obtained from local assembly of the specified region is less than 3 times the maximum expected library size (specified by -1/--maxlibsize). The mutation description starts with either INS, DEL, DUP, or INV for insertion, deletion, duplication, and inversion, respectively and is followed by mutation-specific options.

For insertions, INS should be followed by either a FASTA file containing the sequence to be

inserted, or by the nucleotide sequence itself. For example, INS ATG would insert the sequence ''ATG'' in the middle of the largest contig obtained from the specified region, while INS LINE1.fa would insert the sequence in the FASTA-formatted file LINE1.fa into the largest contig. For deletions, DEL should be followed by the fraction of the largest contig to be deleted, where 1.0 indicates a deletion of the largest assembled region minus one library length on each end. e.g. DEL 0.5. Inversions (INV) have no further options - the region inside the largest contig will be inverted. Duplications (DUP) have one optional parameter, an integer specifying the number of times the sequence of the largest contig should be duplicated, e.g. DUP 2 specifies the region is duplicated twice.

Compound variants are also possible by chaining a number of mutations together in a commadelimited list, e.g. DUP 1, DEL 0.5, INS AAATCC, INV would duplicate the region inside the largest contig once, delete half the width of the region, insert the sequence AAATCC, and invert the region.

Where mutations add sequence (e.g. duplications), new reads are created that will be added to the original BAM, and where mutations remove sequence (e.g. deletions) those reads are maked as excluded (excluded read names are recorded in the file specified by -x/--excluded). Excluded reads are not carried over from the original BAM to the mutated BAM, creating the copy number effect associated with deletion.

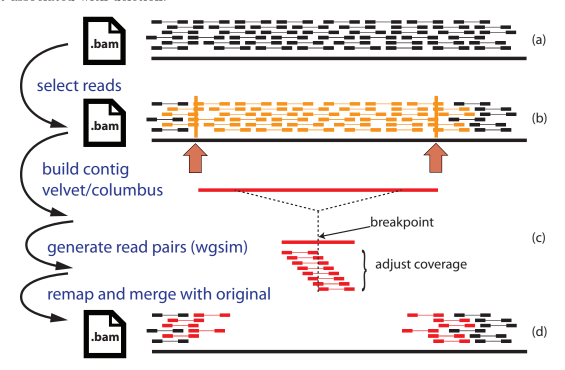


Figure 2: Method for adding multi-nucleotide variants (indels and SVs) to existing BAMs: deletion example. Starting with the original BAM alignment (a) regions, e.g. (b), are assembled using Velvet/Columbus (http://www.ebi.ac.uk/zerbino/velvet/) and the desired mutation(s) are created in the largest contig. Read coverage is simulated over the contig using wgsim (https://github.com/lh3/wgsim, also included with samtools) with parameters -e 0 -r 0 -R 0 to suppress additional mutations, other parameters are set based on the input BAM and desired coverage. Simulated coverage is scaled based on whether the mutation is adds sequence (duplication) or removes sequence (deletions), and replaced into the original BAM while excluded reads are removed from the original BAM (d).