# SomaticSeq Documentation

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## 1 Introduction

SomaticSeq is a flexible post-somatic-mutation-calling workflow for improved accuracy. We have incorporated multiple somatic mutation caller(s) to obtain a combined call set, and then it uses machine learning to distinguish true mutations from false positives from that call set. We have incorporated the following somatic mutation caller: MuTect/Indelocator, MuTect2, VarScan2, JointSNVMix, SomaticSniper, VarDict, MuSE, LoFreq, and Scalpel. You may incorporate some or all of those callers into your own pipeline with SomaticSeq.

The manuscript, An ensemble approach to accurately detect somatic mutations using SomaticSeq, is published in Genome Biology 2015, 16:197. The SomaticSeq project is located at <a href="http://bioinform.github.io/somaticseq/">http://bioinform.github.io/somaticseq/</a>. The data described in the manuscript is also described at <a href="http://bioinform.github.io/somaticseq/data.html">http://bioinform.github.io/somaticseq/data.html</a>. There have been some major improvements since the publication.

SomaticSeq.Wrapper.sh is a bash script that calls a series of scripts to combine the output of the somatic mutation caller(s), after the somatic mutation callers are run. Then, depending on what R scripts are fed to SomaticSeq.Wrapper.sh, it will either 1) train the call set into a classifier, 2) predict high-confidence somatic mutations from the call set based on a pre-defined classifier, or 3) simply label the calls (i.e., PASS, LowQual, or REJECT) based on majority vote of the tools.

## 1.1 Dependencies

- Python 3, plus regex, pysam, numpy, and scipy libraries. All the .py scripts are written in Python 3 (Yes, Python 3. I deserve brownie points from Python developers).
- R, plus the ada package in R.
- BEDTools (if there is an inclusion and/or an exclusion region file)
- Optional: Free GATK version 3. You can use GATK CombineVariants to merge VCF files if the jar file is provided to –gatk. Otherwise, the sorting is done by the vcfsort.pl script written by German Gaston Leparc.
- Optional: dbSNP and COSMIC files in VCF format (if you want to use these features as a
  part of the training).
- At least one of MuTect/Indelocator, MuTect2, VarScan2, JointSNVMix, SomaticSniper, VarDict, MuSE, LoFreq, Scalpel, and Strelka2. Those are the tools we have incorporated in SomaticSeq. If there are other somatic tools that may be good addition to our list, please make the suggestion to us.

## 1.2 Docker repos

- SomaticSeq is dockerized at https://hub.docker.com/r/lethalfang/somaticseq/.
- MuTect (untested): https://hub.docker.com/r/aarjunrao/mutect/
- MuTect2: https://hub.docker.com/r/broadinstitute/gatk
- VarScan2 (untested): https://hub.docker.com/r/jeltje/varscan2/
- JointSNVMix2: https://hub.docker.com/r/lethalfang/jointsnvmix2/
- SomaticSniper: https://hub.docker.com/r/lethalfang/somaticsniper/
- VarDict: https://hub.docker.com/r/lethalfang/vardictjava/
- MuSE: https://hub.docker.com/r/marghoob/muse/
- LoFreq: https://hub.docker.com/r/marghoob/lofreq/
- Scalpel: https://hub.docker.com/r/lethalfang/scalpel/
- Strelka2: https://hub.docker.com/r/lethalfang/strelka/

## 2 To use SomaticSeq.Wrapper.sh

The SomaticSeq.Wrapper.sh is a wrapper script that calls a series of programs and procedures. It can be easily run if all the dependencies are met. Some optional parameters are hard-coded as defaults, but can be easily edited by advanced users. In the next section, we also described the workflow in more detail, so you are not dependent on this wrapper script. You can either modify this wrapper script or create your own workflow.

#### 2.1 To train data set into a classifier

To create a trained classifier, ground truth files are required for the data sets. There is also an option to include a list of regions to include and/or exclude. The exclusion or inclusion regions can be VCF or BED files. An inclusion region may be subset of the call sets where you have validated their true/false mutation status, so that only those regions will be used for training. An exclusion region can be regions where the "truth" is ambigious.

All the output VCF files from individual callers are optional. Those VCF files can be bgzipped if they have .vcf.gz extensions. It is imperative that the parameters used for the training and prediction are identical.

```
# For training, truth file and the correct R script are required.
SomaticSeq.Wrapper.sh \
                     MuTect/variants.snp.vcf \setminus
-mutect
 -mutect2
                     MuTect2/variants.vcf \
                     Indelocator/variants.indel.vcf \
 -indelocator
                     VarScan2/variants.snp.vcf
 -varscan-snv
 -varscan-indel
                     VarScan2/variants.indel.vcf
                     JointSNVMix2/variants.snp.vcf
-jsm
 -sniper
                     SomaticSniper/variants.snp.vcf \
                     VarDict/variants.vcf \
 -vardict
 -muse
                     MuSE/variants.snp.vcf
 -lofreq-snv
                     LoFreq/variants.snp.vcf
 -lofreq-indel
                     LoFreq/variants.indel.vcf \
```

```
--scalpel
                     Scalpel/variants.indel.vcf \
 -strelka -snv
                     Strelka/variants.snv.vcf
 strelka -indel
                     Strelka/variants.indel.vcf \
---normal-bam
                     matched_normal.bam \
-tumor-bam
                     tumor.bam
-ada-r-script
                     ada_model_builder.R \
                    human_b37.fasta \
-genome-reference
 -cosmic
                     cosmic.b37.v71.vcf
--dbsnp
                    dbSNP.b37.v141.vcf \
                    $PATH/TO/GenomeAnalysisTK.jar \
 -gatk
 -exclusion-region
                    ignore.bed \
                     validated.bed
--inclusion-region
--truth-snv
                     truth.snp.vcf
--truth-indel
                     truth.indel.vcf \
 -output-dir
                    $OUTPUT_DIR
```

SomaticSeq.Wrapper.sh supports any combination of the somatic mutation callers we have incorporated into the workflow. SomaticSeq will run based on the output VCFs you have provided. It will train for SNV and/or INDEL if you provide the truth.snp.vcf and/or truth.indel.vcf file(s) as well as the proper R script (ada\_model\_builder.R). Otherwise, it will fall back to the simple caller consensus mode.

## 2.2 To predict somatic mutation based on trained classifiers

Make sure the classifiers and the proper R script (ada\_model\_predictor.R) are supplied, Without either of them, it will fall back to the simple caller consensus mode.

```
# The *. RData files are trained classifier from the training mode.
SomaticSeq.Wrapper.sh
                     MuTect/variants.snp.vcf \
-mutect
                     MuTect2/variants.vcf \
 -mutect2
                     Indelocator/variants.indel.vcf \
--indelocator
 -varscan-snv
                     VarScan2/variants.snp.vcf
---varscan-indel
                     VarScan2/variants.indel.vcf
                     JointSNVMix2/variants.snp.vcf \
--ism
                     SomaticSniper/variants.snp.vcf \
 -sniper
                     VarDict/variants.vcf \
--vardict
 -muse
                     MuSE/variants.snp.vcf
--lofreq-snv
                     LoFreq/variants.snp.vcf \
-lofreq-indel
                     LoFreq/variants.indel.vcf
 -scalpel
                     Scalpel/variants.indel.vcf \
 -strelka-snv
                     Strelka/variants.snv.vcf
 -strelka-indel
                     Strelka/variants.indel.vcf \
---normal-bam
                     matched_normal.bam \
-tumor-bam
                     tumor.bam \
--ada-r-script
                     ada_model_predictor.R \
                     human_b37.fasta \
-genome-reference
                     cosmic.b37.v71.vcf
 -cosmic
--dbsnp
                     dbSNP.b37.v141.vcf
 -snpeff-dir
                     $PATH/TO/DIR/snpSift \
--gatk
                     $PATH/TO/GenomeAnalysisTK.jar \
--exclusion-region
                     ignore.bed \
--inclusion-region
                     validated.bed
---classifier --snv
                     sSNV. Classifier . RData \
--classifier -indel
                     sINDEL. Classifier . RData \
--pass-threshold
                     0.5 \ \
 -lowqual-threshold
                    0.1 \
 -output-dir
                     $OUTPUT_DIR
```

If both MuTect2 and MuTect/Indelocator VCF files are provided, the script is written such that it will use MuTect2 and ignore MuTect.

## 2.3 To classify based on simple majority vote

Same as the command previously, but not including the R script or the ground truth files. Without those information, SomaticSeq will fall back into a simple majority vote.

## 3 The step-by-step SomaticSeq Workflow

We'll describe the workflow here, so you may modify the workflow and/or create your own workflow instead of using the wrapper script described previously.

#### 3.1 Combine the call sets

We use GATK CombineVariants to combine the VCF files fkeep-intermediates:rom different callers, although it does not matter what tools are used to merge VCF files. We GATK CombineVariants because it's quite fast. To make them compatible with GATK, the VCF files are modified.

A simple alternative method is to use GNU sort and uniq in Linux to list all the unique first-5-column (i.e., CHROM, POS, ID, REF, and ALT) from all the VCF files, and then fill the remaining required VCF columns with whatever string and sort it according to the reference. That VCF file will do the job just fine.

For our GATK CombineVariants procedure:

1. Modify MuTect and/or Indelocator output VCF files. Since MuTect's output VCF do not always put the tumor and normal samples in the same columns, the script will determine that information based on either the BAM files (the header has sample name information), or based on the sample information that you tell it, and then determine which column belongs to the normal, and which column belongs to the tumor.

```
# Modify MuTect and Indelocator's output VCF based on BAM files
modify_MuTect.py -infile input.vcf -outfile output.vcf -nbam normal.bam -tbam
tumor.bam

# Based on the sample name you supply:
modify_MuTect.py -infile input.vcf -outfile output.vcf -nsm NormalSampleName -
tsm TumorSampleName
```

2. Modify VarScan's output VCF files to be rigorously concordant to VCF format standard, and to attach the tag 'VarScan2' to somatic calls.

```
# Do it for both the SNV and indel modify_VJSD.py -method VarScan2 -infile input.vcf -outfile output.vcf
```

3. JointSNVMix2 does not output VCF files. In our own workflow, we have already converted its text file into a basic VCF file with an 2 awk one-liners, which you may see in the Run\_5\_callers directory, which are:

```
# To avoid text files in the order of terabytes, this awk one-liner keeps
     entries where the reference is not "N", and the somatic probabilities are
    at least 0.95.
awk -F "\t" 'NR!=1 && $4!="N" && $10+$11>=0.95'
# This awk one-liner converts the text file into a basic VCF file
awk -F "\t" '{ print $1 "\t" $2 "\t.\t" $3 "\t" $4 "\t.\t.\tAAAB=" $10 "; AABB=" $11 "\tRD:AD\t" $5 ":" $6 "\t" $7 ":" $8}'
## The actual commands we've used in our workflow:
echo -e '##fileformat=VCFv4.1' > unsorted.vcf
echo -e '##NFO=<ID=AAAB, Number=1, Type=Float, Description="Probability of Joint Genotype AA in Normal and AB in Tumor">'>> unsorted.vcf
echo -e '##NFO=<ID=AABB, Number=1, Type=Float, Description="Probability of Joint
      Genotype AA in Normal and BB in Tumor">'>> unsorted.vcf
echo -e '##FORMAT=<ID=RD, Number=1, Type=Integer, Description="Depth of reference
    -supporting bases (reads1)">'>> unsorted.vcf
echo -e '##FORMAT=<ID=AD, Number=1, Type=Integer, Description="Depth of variant-
    supporting bases (reads2)">'>' >> unsorted.vcf
echo -e '#CHROM\tPOS\tID\tREF\tALT\tQUAL\tFILTER\tINFO\tFORMAT\tNORMAL\tTUMOR'
     >> unsorted.vcf
python $PATH/TO/jsm.py classify joint_snv_mix_two genome.GRCh37.fa normal.bam
    tumor.bam\ trained.parameter.cfg\ /dev/stdout\ |\backslash
awk -F "\t" 'NR!=1 && $4!="N" && $10+$11>=0.95' |\
awk -F "\t" '{print $1 "\t" $2 "\t.\t" $3 "\t" $4 "\t.\t.\tAAAB=" $10 ";AABB="
$11 "\tRD:AD\t" $5 ":" $6 "\t" $7 ":" $8}' >> unsorted.vcf
```

After that, you'll also want to sort the VCF file. Now, to modify that basic VCF into something that will be compatible with other VCF files under GATK CombineVariants:

```
modify_VJSD.py -method JointSNVMix2 -infile input.vcf -outfile output.vcf
```

4. Modify SomaticSniper's output:

```
modify_VJSD.py -method SomaticSniper -infile input.vcf -outfile output.vcf
```

5. VarDict has both SNV and indel, plus some other variants in the same VCF file. Our script will create two files, one for SNV and one for indel, while everything else is ignored for now. By default, LikelySomatic/StrongSomatic and PASS calls will be labeled VarDict. However, in our SomaticSeq paper, based on our experience in DREAM Challenge, we implemented two custom filters to relax the VarDict tagging criteria.

```
# Default VarDict tagging criteria, only PASS (and Likely or Strong Somatic):
modify_VJSD.py -method VarDict -infile intput.vcf -outfile output.vcf

# When running VarDict, if var2vcf_paired.pl is used to generate the VCF file,
you may relax the tagging criteria with -filter paired
modify_VJSD.py -method VarDict -infile intput.vcf -outfile output.vcf -filter
paired

# When running VarDict, if var2vcf_somatic.pl is used to generate the VCF file
, you may relax the tagging criteria with -filter somatic
```

```
| modify_VJSD.py -method VarDict -infile intput.vcf -outfile output.vcf -filter somatic
```

In the SomaticSeq paper, -filter somatic was used because var2vcf\_somatic.pl was used to generate VarDict's VCF files. In the SomaticSeq.Wrapper.sh script, however, -filter paired is used because VarDict authors have since recommended var2vcf\_paired.pl script to create the VCF files. While there are some differences (different stringencies in some filters) in what VarDict labels as PASS between the somatic.pl and paired.pl scripts, the difference is miniscule after applying our custom filter (which relaxes the filter, resulting in a difference about 5 calls out of 15,000).

The output files will be snp.output.vcf and indel.output.vcf.

6. MuSE was not a part of our analysis in the SomaticSeq paper. We have implemented it later.

```
modify_VJSD.py -method MuSE -infile input.vcf -outfile output.vcf
```

- 7. LoFreq and Scalpel do not require modification. LoFreq has no sample columns anyway.
- 8. Strelka's INDEL VCF clashes with Scalple's VCF during GATK CombineVariants, for reasons we haven't bothered to figure out. We just removed the sample columns for Strelka's INDEL to solve the problem during variant merging.

```
gunzip -c somatic.indel.vcf.gz | awk -F "\t" '{ if ($0 !~ /^##/) print $1 "\t" $2 "\t" $3 "\t" $4 "\t" $5 "\t" $6 "\t" $7 "\t" $8; else print $0}' OFS ="\t" > strelka.indel.truncated.vcf
```

9. Finally, with the VCF files modified, you may combine them with GATK CombineVariants: one for SNV and one for indel separately. There is no particular reason to use GATK CombineVariants. Other combiners should also work. The only useful thing here is to combine the calls and GATK CombineVariants does it well and pretty fast.

```
# Combine the VCF files for SNV. Any or all of the VCF files may be present.

# -nt 6 means to use 6 threads in parallel
java -jar $PATH/TO/GenomeAnalysisTK.jar -T CombineVariants -R genome.GRCh37.fa
-nt 6 --setKey null --genotypemergeoption UNSORTED -V mutect.vcf -V
varscan.snp.vcf -V jointsnvmix.vcf -V snp.vardict.vcf -V muse.vcf --out
CombineVariants.snp.vcf
java -jar $PATH/TO/GenomeAnalysisTK.jar -T CombineVariants -R genome.GRCh37.fa
-nt 6 --setKey null --genotypemergeoption UNSORTED -V indelocator.vcf -V
varscan.snp.vcf -V indel.vardict.vcf --out CombineVariants.indel.vcf
```

# 3.2 For model training: process and annotate the VCF files (union of call sets)

This step may be needed for model training. The workflow in SomaticSeq.Wrapper.sh allows for inclusion and exclusion region. An inclusion region means we will only use calls inside these regions. An exclusion region means we do not care about calls inside this region. DREAM Challenge had exclusion regions, e.g., blacklisted regions, etc.

```
# In the DREAM_Stage_3 directory, we have included an exclusion region BED file as an example
# This command uses BEDtools to rid of all calls in the exclusion region intersectBed -header -a BINA_somatic.snp.vcf -b ignore.bed -v > somatic.snp. processed.vcf
intersectBed -header -a BINA_somatic.indel.vcf -b ignore.bed -v > somatic.indel. processed.vcf

# Alternatively (or both), this command uses BEDtools to keep only calls in the inclusion region intersectBed -header -a BINA_somatic.snp.vcf -b inclusion.bed > somatic.snp. processed.vcf
intersectBed -header -a BINA_somatic.indel.vcf -b inclusion.bed > somatic.indel. processed.vcf
```

# 3.3 Convert the VCF file, annotated or otherwise, into a tab separated file

This script works for all VCF files. It extracts information from BAM files as well as some VCF files created by the individual callers. If the ground truth VCF file is included, a called variant will be annotated as a true positive, and everything will be annotated as a false positive.

```
# SNV
SSeq_merged.vcf2tsv.py -ref genome.GRCh37.fa -myvcf somatic.snp.processed.vcf -
truth Ground.truth.snp.vcf -mutect MuTect/variants.snp.vcf.gz -varscan VarScan2
/variants.snp.vcf -jsm JSM2/variants.vcf -sniper SomaticSniper/variants.vcf -
vardict VarDict/snp.variants.vcf -muse MuSE/variants.vcf -lofreq LoFreq/
variants.snp.vcf -strelka Strelka/variants.snp.vcf -dedup -tbam tumor.bam -nbam
normal.bam -outfile Ensemble.sSNV.tsv
```

That was for SNV, and indel is almost the same thing. After version 2.1, we have replaced all information from SAMtools and HaplotypeCaller with information directly from the BAM files. The accuracy differences are negligible with significant improvement in usability and resource requirement.

At the end of this, Ensemble.sSNV.tsv and Ensemble.sINDEL.tsv are created.

All the options for SSeq\_merged.vcf2tsv.py are listed here. They can also be displayed by running SSeq\_merged.vcf2tsv.py –help.

```
-myvcf Input VCF file of the merged calls [REQUIRED]
-ref Genome reference fa/fasta file [REQUIRED]
-nbam BAM file of the matched normal sample [REQUIRED]
-tbam BAM file of the tumor sample [REQUIRED]
-ref Genome reference fa/fastq file [REQUIRED]
-truth Ground truth VCF file. Every other position is a False Positive.
```

```
dbSNP VCF file
-dbsnp
             COSMIC VCF file
-cosmic
             VCF file from either MuTect2, MuTect, or Indelocator
-mutect
             VCF file from SomaticSniper
-\operatorname{sniper}
             VCF file from VarScan2
-varscan
-jsm
             VCF file from Bina's workflow that contains JointSNVMix2
             VCF file that contains only SNV or only INDEL from VarDict
-vardict
-muse
             VCF
                 file from MuSE
             VCF file from LoFreq
-lofreq
-scalpel
             VCF file from Scalpel
-\operatorname{strelk} a
             VCF file from Strelka
-dedup
             A flag to consider only primary reads
-minMQ
             Minimum mapping quality for reads to be considered (Default = 1)
             Minimum base quality for reads to be considered (Default = 5)
-minBQ
-mincaller
             Minimum number of caller classification for a call to be considered
 (Use 0.5 to consider some LowQual calls. Default = 0).
-scale
             The options are phred, fraction, or None, to convert numbers to
 Phred scale or fractional scale. (default = None, i.e., no conversion)
-outfile
             Output TSV file name
```

Note: Do not worry if Python throws a warning like this.

```
RuntimeWarning: invalid value encountered in double_scalars z = (s - expected) / np.sqrt(n1*n2*(n1+n2+1)/12.0)
```

This is to tell you that scipy was attempting some statistical test with empty data. That's usually due to the fact that normal BAM file has no variant reads at that given position. That is why lots of values are NaN for the normal.

## 3.4 Model Training or Mutation Prediction

You can use Ensemble.sSNV.tsv and Ensemble.sINDEL.tsv files either for model training (provided that their mutation status is annotated with 0 or 1) or mutation prediction. This is done with stochastic boosting algorithm we have implemented in R.

Model training:

```
# Training:
r_script/ada_model_builder.R Ensemble.sSNV.tsv
r_script/ada_model_builder.R Ensemble.sINDEL.tsv
```

Ensemble.sSNV.tsv.Classifier.RData and Ensemble.sINDEL.tsv.Classifier.RData will be created from model training.

Mutation prediction:

```
# Mutation prediction:
r_script/ada_model_predictor.R Ensemble.sSNV.tsv.Classifier.RData Ensemble.sSNV.
tsv Trained.sSNV.tsv
r_script/ada_model_predictor.R Ensemble.sINDEL.tsv.Classifier.RData Ensemble.sINDEL
.tsv Trained.sINDEL.tsv
```

After mutation prediction, if you feel like it, you may convert Trained.sSNV.tsv and Trained.sINDEL.tsv into VCF files. Use -tools to list ONLY the individual tools used to have appropriately annotated VCF files. Accepted tools are CGA (for MuTect/Indelocator), VarScan2,

JointSNVMix2, SomaticSniper, VarDict, MuSE, LoFreq, and/or Scalpel. To list a tool without having run it, the VCF will be annotated as if the tool was run but did not identify that position as a somatic variant.

```
# Probability above 0.7 labeled PASS (-pass 0.7), and between 0.1 and 0.7 labeled
LowQual (-low 0.1):

# Use -all to include REJECT calls in the VCF file

# Use -phred to convert probability values (between 0 to 1) into Phred scale in the
QUAL column in the VCF file

SSeq_tsv2vcf.py -tsv Trained.sSNV.tsv -vcf Trained.sSNV.vcf -pass 0.7 -low 0.1
-tools CGA VarScan2 JointSNVMix2 SomaticSniper VarDict MuSE LoFreq Strelka -all
-phred

SSeq_tsv2vcf.py -tsv Trained.sINDEL.tsv -vcf Trained.sINDEL.vcf -pass 0.7 -low 0.1
-tools CGA VarScan2 VarDict LoFreq Scalpel Strelka -all -phred
```

## 4 Release Notes

Make sure training and prediction use the same version. Otherwise the prediction is not valid.

#### 4.1 Version 1.0

Version used to generate data in the manuscript and Stage 5 of the ICGC-TCGA DREAM Somatic Mutation Challenge, where SomaticSeq's results were #1 for INDEL and #2 for SNV.

In the original manuscript, VarDict's var2vcf\_somatic.pl script was used to generate VarDict VCFs, and subsequently "-filter somatic" was used for SSeq\_merged.vcf2tsv.py. Since then (including DREAM Challenge Stage 5), VarDict recommends var2vcf\_paired.pl over var2vcf\_somatic.pl, and subsequently "-filter paired" was used for SSeq\_merged.vcf2tsv.py. The difference in SomaticSeq results, however, is pretty much negligible.

## 4.2 Version 1.1

Automated the SomaticSeq.Wrapper.sh script for both training and prediction mode. No change to any algorithm.

## 4.3 Version 1.2

Have implemented the following improvement, mostly for indels:

- SSeq\_merged.vcf2tsv.py can now accept pileup files to extract read depth and DP4 (reference
  forward, reference reverse, alternate forward, and alternate reverse) information (mainly for
  indels). Previously, that information can only be extracted from SAMtools VCF. Since the
  SAMtools or HaplotypeCaller generated VCFs hardly contain any indel information, this
  option improves the indel model. The SomaticSeq.Wrapper.sh script is modified accordingly.
- Extract mapping quality (MQ) from VarDict output if this information cannot be found in SAMtools VCF (also mostly benefits the indel model).
- Indel length now positive for insertions and negative for deletions, instead of using the absolute value previously.

#### 4.4 Version 2.0

- Removed dependencies for SAMtools and HaplotypeCaller during feature extraction. SSeq\_merged.vcf2tsv.py extracts those information (plus more) directly from BAM files.
- Allow not only VCF file, but also BED file or a list of chromosome coordinate as input format for SSeq\_merged.vcf2tsv.py, i.e., use -mybed or -mypos instead of -myvcf.
- Instead of a separate step to annotate ground truth, that can be done directly by SSeq\_merged.vcf2tsv.py by supplying the ground truth VCF via -truth.
- SSeq\_merged.vcf2tsv.py can annotate dbSNP and COSMIC information directly if BED file or a list of chromosome coordinates are used as input in lieu of an annotated VCF file.
- Consolidated feature sets, e.g., removed some redunda Fixed a bug: if JointSNVMix2 is not included, the values should be "NaN" instead of 0's. This is to keep consistency with how we handle all other callersnt feature sets coming from different resources.

#### 4.5 Version 2.0.2

- Incorporated LoFreq.
- Used getopt to replace getopts in the SomaticSeq.Wrapper.sh script to allow long options.

#### 4.6 Version 2.1.2

- Properly handle cases when multiple ALT's are calls in the same position. The VCF files can either contain multiple calls in the ALT column (i.e., A,G), or have multiple lines corresponding to the same position (one line for each variant call). Some functions were significantly re-written to allow this.
- Incorporated Scalpel.
- Deprecated HaplotypeCaller and SAMTools dependencies completely as far as feature generation is concerned.
- The Wrapper script removed SnpSift/SnpEff dependencies. Those information can be directly obtained during the SSeq\_merged.vcf2tsv.py step. Also removed some additional legacy steps that has become useless since v2 (i.e., score\_Somatic.Variants.py). Added a step to check the correctness of the input. The v2.1 and 2.1.1 had some typos in the wrapper script, so only describing v2.1.2 here.

#### 4.7 Version 2.2

• Added MuTect2 support.

#### 4.8 Version 2.2.1

- InDel\_3bp now stands for indel counts within 3 bps of the variant site, instead of exactly 3 bps from the variant site as it was previously (likewise for InDel\_2bp).
- Collapse MQ0 (mapping quality of 0) reads supporting reference/variant reads into a single metric of MQ0 reads (i.e., tBAM\_MQ0 and nBAM\_MQ0). From experience, the number of MQ0 reads is at least equally predictive of false positive calls, rather than distinguishing if those MQ0 reads support reference or variant.

- Obtain SOR (Somatic Odds Ratio) from BAM files instead of VarDict's VCF file.
- Fixed a typo in the SomaticSeq.Wrapper.sh script that did not handle inclusion region correctly.

## 4.9 Version 2.2.2

- Got around an occasional unexplained issue in then ada package were the SOR is sometimes categorized as type, by forcing it to be numeric.
- Defaults PASS score from 0.7 to 0.5, and make them tunable in the SomaticSeq.Wrapper.sh script (-pass-threshold and -lowqual-threshold).

#### 4.10 Version 2.2.3

- Incorporated Strelka2 since it's now GPLv3.
- Added another R script (ada\_model\_builder\_ntChange.R) that uses nucleotide substitution pattern as a feature. Limited experiences have shown us that it improves the accuracy, but it's not heavily tested yet.
- If a COSMIC site is labeled SNP in the COSMIC VCF file, if\_cosmic and CNT will be labeled as 0. The COSMIC ID will still appear in the ID column. This will not change any results because both of those features are turned off in the training R script.
- Fixed a bug: if JointSNVMix2 is not included, the values should be "NaN" instead of 0's. This is to keep consistency with how we handle all other callers.

## 4.11 Version 2.2.4

- Resolved a bug in v2.2.3 where the VCF files of Strelka INDEL and Scalpel clash on GATK CombineVariants, by outputting a temporary VCF file for Strelka INDEL without the sample columns.
- Caller classification: consider if\_Scalpel = 1 only if there is a SOMATIC flag in its INFO.

#### 4.12 Version 2.2.5

- Added a dockerfile. Docker repo at https://hub.docker.com/r/lethalfang/somaticseq/.
- Ability to use vcfsort.pl instead of GATK CombineVariants to merge VCF files.

#### 4.13 Version 2.3.0

- Moved some scripts to the utilities directory to clean up the clutter.
- Added the split\_Bed\_into\_equal\_regions.py to utilities, which will split a input BED file into multiple BED files of equal size. This is to be used to parallelize large WGS jobs.
- Made compatible with MuTect2 from GATK4.
- Removed long options for the SomaticSeq.Wrapper.sh script because it's more readable this
  way.

# 5 Contact Us

For suggestions, bug reports, or technical support, please post in  $\frac{1}{2}$  https://github.com/bioinform/somaticseq/issues. The developers are alerted when issues are created there. Alternatively, you may also email li\_tai.fang@roche.com.