CellPhy - SC-Caller conversion

Alexey Kovlov, João M Alves*, Alexandros Stamatakis & David Posada

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1 Introductory Note

In the following supporting document, we will show how to easily transform a VCF file generated by the SC-Caller (a single-cell specific variant caller) into a CellPhy-GL readable VCF. Before we start, users should make sure that both BCFtools and Tabix from Samtools are installed and available.

2 VCF file conversion

SC-Caller VCF files can be converted by running the sc-caller-convert.sh script as follows:

```
$ ./sc-caller-convert.sh
Usage: ./sc-caller-convert.sh inputVCF SamplePrefix
Created by: Alexey Kovlov, Joao M Alves, Alexandros Stamatakis & David Posada - June 2020
$ ./sc-caller-convert.sh CRC_Cell-A.22.sccaller.vcf Cell-A
```

If everything went as expected, you should have generated a new VCF (Cell-A.PL-fixed.vcf) that is now ready for downstream analysis. Below we provide a detailed explanation of which changes are being made to the original VCF file.

2.1 Step-by-step explanation

While many callers currently being used with single-cell genomic data generate VCF files with a standard "PL" field (e.g., GATK-HaplotypeCaller, Monovar, Prosolo), it is important to highlight that the "PL" definition may differ from its standard meaning in different tools. Indeed, SC-Caller for instance uses the "PL" field to store not only the likelihood of heterozygous and alternative homozygous genotypes, but also the likelihood of sequencing noise and amplification artifacts.

Below we show an example of a VCF generated using SC-Caller (version 2.0.0).

```
$ grep -v "#" CRC Cell-A.22.sccaller.vcf | head -n 5
#CHROM POS ID REF ALT QUAL
                                            FORMAT CELLO01
                             FILTER INFO
                         13 .
  16149851
                  G
                                  NS=1
                                         GT:SO:AD:BT:GO:PI.
                                                             0/0:NA:0.6:0.6:13:91.54.13.0
                      Α
                      -1C 4
                                         GT:SO:AD:BI:GQ:PL
   16190307
                                  NS=1
                                                             0/0:NA:0,1:0.6:4:15,9,2,0
                      C,G 150 multiple-genotype
                                                        GT:SO:AD:BI:GO:PL 0/0:True:317.1:0.993:150:91.227.64.11570
   16193737
                                                 NS=1
                                         GT:SO:AD:BI:GQ:PL 0/0:True:136,1:0.6:150:39,93,310,3479
                                  NS=1
                      A 150 .
```

Looking at this output, you should notice that:

- SC-Caller renames the single-cell to "CELL001";
- The VCF contains calls other than single-nucleotide variants (i.e., SNVs);
- The "PL" field of **bi-allelic** sites is composed of 4 entries, as opposed to the 3 values usually observed in VCF files that were generated following the *format specifications**;
- The "PL" field will always contain 4 entries, regardless of the amount of alternative variants detected.

As a consequence, we will need to rename the sample to its proper ID and trim the VCF to exclude all indels and non-biallelic positions (as we won't be able to get the likelihood scores for all possible genotypes). This can be easily done using BCFtools.

```
# Rename sample in VCF
$ cat temp_rename
CELL001 Cell-A
$ bcftools reheader -s temp_rename CRC_Cell-A.22.sccaller.vcf -o temp.renamed.vcf
# Remove indels and non-biallelic sites
$ bcftools view --types snps -f "." temp.renamed.vcf -o temp.snvs.vcf
```

Once this is done, we should end up with a trimmed VCF that solely contains bi-allelic sites:

```
$ grep -v "##" temp.snvs.vcf | head -n 5
      POS ID REF ALT QUAL
                                             FORMAT Cell-A
              . G A 13 .
  16149851
                                 NS=1
                                         GT:SO:AD:BI:GQ:PL
                                                             0/0:NA:0,6:0.6:13:91,54,13,0
   16195864
                  G
                      Α
                         150 .
                                  NS=1
                                          GT:SO:AD:BI:GQ:PL
                                                             0/0:True:136,1:0.6:150:39,93,310,3479
   16195889
                      C
                          150 .
                                  NS=1
                                          GT:SO:AD:BI:GQ:PL
                                                             0/0:True:136,1:0.6:150:34,89,306,4496
                          150
                                          GT:SO:AD:BI:GQ:PL
                                                            0/0:True:132,1:0.6:150:34,87,298,4273
```

Afterwards, we need to transform the "PL" field at each site into the standard "PL" format. Following **SC-caller authors' suggestions**, we take the highest likelihood score of the first two values (i.e., sequencing noise, amplification artifact) as the phred-scaled genotype likelihood of the reference homozygous (0/0) genotype, and the remaining values as the likelihood for heterozygous (0/1) and alternative homozygous (1/1) genotypes, respectively.

Cautionary remark on SC-Caller developers suggestion:

It is perhaps important to mention that the SC-Caller authors suggest to take "the bigger number as the PL combined" as the 0/0 genotype likelihood. We interpreted this suggestion as to take the one with the "the highest likelihood" (which in this case would be the smallest integer value).

Our custom script will then essentially rename the previous "PL" field as "FPL" and add a standard "PL" field to our VCF to make it CellPhy readable:

```
$ grep -v "##" temp.PL-fixed.vcf | head
      POS ID REF ALT QUAL
                             FILTER INFO
                                            FORMAT Cell-A
              . G A 13 .
                                         GT:SO:AD:BI:GQ:FPL:PL
22 16149851
                                                                0/0:NA:0,6:0.6:13:91,54,13,0:54,13,0
   16195864
                 G
                     Α
                         150 .
                                 NS=1
                                         GT:SO:AD:BI:GQ:FPL:PL
                                                                0/0:True:136,1:0.6:150:39,93,310,3479:39,310,3479
   16195889
                         150 .
                                 NS=1
                                         GT:SO:AD:BI:GQ:FPL:PL
                                                                0/0:True:136,1:0.6:150:34,89,306,4496:34,306,4496
                     C
                  Т
   16195900
                 A G
                                 NS=1
                                         GT:SO:AD:BI:GQ:FPL:PL
                                                                0/0:True:132,1:0.6:150:34,87,298,4273:34,298,4273
                         150 .
```

3 Merging VCF files for downstream analyses

Once all single-cell VCF files are converted, we can easily merge them into a multi-sample VCF using BCFtools.

```
$ for i in *PL-fixed.vcf
do
bgzip $i
tabix -p vcf ${i}.gz
done
$ ls *.vcf.gz > listMERGE
$ bcftools merge -l listMERGE -O vcf -o CRC-allCells.vcf
```

The VCF is now ready for CellPhy (or for any additional filtering steps).