# VCF\_creator: module of DiscoSnp++

## Mapping and VCF Creation features

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# **User part**

### VCF\_creator : one module in DiscoSnp++ Mapping and VCF Creation

**VCF\_creator** is designed to map on a genome the output of DiscoSnp++ the Single Nucleotide Polymorphism (SNP) and small indels. This module create a Variant Calling Format (VCF) from the output of DiscoSnp++ or from an alignment obtained with the software BWA \*.

### **Quick starting**

- Download and uncompress the DiscoSnp++
- Install BWA (you can add it to your PATH)
- Run the simple example:
  - XXX

# Running VCF\_creator

- The main script *run VCF creator.sh* has three modes according to your needs:
  - MODE 1: You don't have a reference genome but you want to create a vcf (it will summarize the DiscoSnp++ informations and will give the position of the variant on the upper path of DiscoSnp++ variant). The module will create a vcf from the output of DiscoSnp++:
    - ./run\_VCF\_creator.sh -p < disco\_file> -o < output> [-s < bwa\_errors\_in\_seed>]
  - MODE 2: You have a reference genome and you want to align your variants against a reference genome. The module will run BWA to make an alignment between your reference genome and the output of DiscoSnp++:
    - ./run\_VCF\_creator.sh -G <ref> -p <disco\_file> -o <output> [-B <path\_bwa>] [-l <seed size>] [-n <mismatch number>] [-s <bwa errors in seed>] [-w]
    - **MODE 3**: You already have an alignment (.sam file) and you want to create a vcf file:
      - ./run VCF creator.sh -f <sam file> -n <mismatch number> -o <output>

### **Options**

#### General options:

- -p: DiscoSnp++ output file (<file>.fasta) (Mandatory unless MODE 3)
- -o : ouput (<file>.vcf) (Mandatory)

<sup>\*: (</sup>Li H. and Durbin R. (2010) Fast and accurate long-read alignment with Burrows-Wheeler Transform. Bioinformatics, Epub. [PMID: 20080505]).

- -G: reference genome (<file.fasta>) (Only in MODE 2)
- B: bwa path ( /home/me/my\_programs/bwa-0.7.12/) (note that bwa must be precompiled) (Only in MODE 2) (not necessary if BWA is in the path)
- -f: alignment already done (<file>.sam) (Only in MODE 3)
- -h: show help
- -w : remove waste tmp files (bwa index files)
- -I : create of an output (VCF format) specific to IGV (Integrative Genomics Viewer) : sorting VCF by mapping positions and removing unmapped variants

BWA specifics options (- warning, bwa mapping running time highly depends on this parameter):

- -s: bwa option (-s): seed distance (optionnal, default value 0)
- -1: bwa option (-l): length of the seed for alignment (optionnal, default value 10)
- -n: bwa option (-n): maximal bwa mapping distance
  - Optionnal in MODE 1 and MODE 3 ( *default value 3*)
  - **Mandatory** in MODE 3, it's preferable to use the same value used during alignment by BWA (*default value 3*)

Sample example: XXX

### Output

- **Final results are in your <file>.vcf.** It's a Variant Call Format that will summarize all the mapping information and the header of DiscoSnp++ informations. Example :
- MODE 1:

		ID	REF	ALT	QUAL	FILTE	3	INFO	FORMAT	G1	G2	G3
<b>G4</b> SNP_higher_path_	<b>G5</b> _13736	30	13736	Α	G			Ty=SNP	;Rk=0.99909;UL	.=.;UR=	.;CL=.;C	R=.
GT:DP:P	L:AD	0/0:5344	17:1947, <i>*</i>	160066,1	067676:5	3425,22	0/1:3:40	,9,20:1,2	1/1:94862:1895	477,284	1398,357	75:30,94832
0/1:5:34,9	9,54:3,2	1/1:6538	39:13066	61,19610	01,2493:1	9,65370						
SNP_higher_path_	_5442	30	5442_1	Α	G			Ty=SNP	;Rk=0.42995;UL	.=.;UR=	.;CL=.;C	R=.
GT:DP:P	L:AD	0 1:19:98	8,14,198	:12,7	0 1:18:1	38,12,13	8:9,9	0 1:8:66	,10,66:4,4	0 1:13	30:257,1	24,1813:104,26
0 0:224:1	4,598,43	324:220,4	4									
SNP_higher_path_	_5442	31	5442_2	G	Α			Ty=SNP	;Rk=0.42995;UL	.=.;UR=	.;CL=.;C	R=.
GT:DP:P	L:AD	0 1:19:9	8,14,198	:12,7	0 1:18:1	38,12,13	8:9,9	0 1:8:66	,10,66:4,4	0 1:13	30:257,1	24,1813:104,26
0 0:224:1	4,598,43	324:220,4	4									
INDEL_higher_pat			586	GGAC				-	Rk=0.40534;UL=	=.;UR=.;	CL=.;CF	₹=.
GT:DP:P	L:AD	0/1:127:	837,17,9	77:67,60	1/1:387:	6056,518	,408:52,	335	0/1:70:372,21,6	51:42,2	8	
0/1:126:1	395,53,4	477:40,86	6	0/1:102:	671,16,7	91:54,48						

#### MODE 2 and MODE 3:

#CHROM	POS	ID	REF	ALT	QUAL FILTER		INFO	FORM	IAT	G1	G2	G3
<b>G4</b>	G5											
Pseudomonas	109860	15103	С	G	. PASS							
Ty=SNI	P;Rk=0.6	5101;DT	=1;UL=.	;UR=.;CL	=.;CR=.;Genome=C;	Sd=-1	GT:DP:	PL:AD	1/1:4:84	1,16,4:0,	4 1/1:1:2	24,7,4:0,1
1/1:1:24	4,7,4:0,1	0/1:6:17	7,15,97:	5,1	0/1:10:124,14,44:3	,7						
Pseudomonas	4416118	31416_1	Α	G	. PASS							
Ty=SNI	P;Rk=0.4	1638;DT	=1;UL=.	;UR=.;CL	=.;CR=.;Genome=A;	Sd=1	GT:DP:	PL:AD	1 1:35:6	606,71,2	8:3,32	0
1:26:403,41,43:4	1,22	1 1:34:6	617,79,1	8:2,32	0 1:46:316,14,356:	24,22	0 1:43:5	47,36,12	28:11,32			
Pseudomonas	441611	91416_2	Α	С	. PASS							

```
Ty=SNP;Rk=0.41638;DT=1;UL=.;UR=.;CL=.;CR=.;Genome=A;Sd=1 GT:DP:PL:AD 1|1:35:606,71,28:3,32
1:26:403,41,43:4,22
                 1|1:34:617,79,18:2,32 0|1:46:316,14,356:24,22 0|1:43:547,36,128:11,32
Pseudomonas
           131283714361 C CGTAGAAGATCTC
                                                      PASS
      Ty=DEL;Rk=0.26896;DT=0;UL=.;UR=.;CL=.;Genome=.;Sd=-1 GT:DP:PL:AD 0/0:792:47,1978,15015:771,21
      0/0:783:32,2016,14979:766,17
      0/1:849:1303,868,12339:701,148
Pseudomonas
           17532726164
                        TGTGCAGGTCGAAGACTTGA
                                                            PASS
      Ty=INS;Rk=0.13677;DT=0;UL=.;UR=.;CL=.;CR=.;Genome=.;Sd=1
                                                      GT:DP:PL:AD
                                                                  0/0:1086:34,2817,20829:1064,22
      0/0:1486:124,3555,27823:1437,49
      0/0:1132:585.1967.19224:1033.99
```

- In this example, the three types of DiscoSnp++ variants : in red : simple SNP; in green close SNPs; in black INDEL.
- The VCF file has the following fields:
  - **CHROM**: chromosome id where the prediction is mapped, or allele id of the upper path if the variant is unmapped or if no reference genome is provided.
  - **POS** (1-based leftmost position):
    - If a reference genome is provided and if the variant is mapped on a unique position: the mapping position of the variant
    - If a reference genome is provided and if the variant is not uniquely mapped: one of the positions of the variant (1-based lefmost position)
    - Else (no reference genome provided or unmapped variant): position of the variant on the upper path of the discoSnp++ prediction (including the left extension)
  - **ID**: identification of the variant (used by DiscoSnp++). For the close SNPs, the SNP number is added to the ID. Example: 10388\_2

#### • **REF**:

- If one of the two predicted allele maps this position : the corresponding variant
- Else, or if no reference genome provided: the lexicographically smallest of the two variants
- In case of close SNPs: the first is defined as previously described. The following SNPs are those located on the same path
- ALT: The variant non reported as the "REF" variant
- **QUAL**: "." (unused)

#### • FILTER:

- PASS if the variant is mapped at unique position
- MULTIPLE if the variant is mapped on multiple positions
- ".": if the variant is unmapped or if no reference genome is provided

#### • **INFO**:

- **Ty**: Type of variant
  - SNP: If the variant is a simple SNP or close SNPs
  - INS: If the variant mapped corresponds to the longuest path; the alt carries the deletion
  - DEL: If the variant mapped corresponds to the shortest path; the alt carries the deletion
- **Rk**: Rank of the prediction computed by DiscoSnp++ (if several datasets are used in DiscoSnp++, ranks the predictions according to their read coverage in each condition favoring SNPs that are discriminant between conditions value between 0 and 1)
- **DT**: If the variant is mapped on a unique position: distance of the mapping (number of mismatches). If the variant is unmapped or mapped on multiple positions: "-1"
- UL : Length of the left unitig ("." if not computed)
- UR: Length of the right unitig ("." if not computed)
- CL : Length of the left contig ("." if not computed)
- **CR**: Length of the right contig ("." if not computed)
- Genome: Applies only for SNPs when a reference genome is provided ("." for INDELs and when no reference genome provide or if the variant is unmapped).
   Reference nucleotide (!!nucleotide in the reference genome !! In general it is correspond to the REF field; could be different for close snps). Important
   Remark: If one of the two predictions matches the reference: equal to the "REF" field, else equal to the nucleotide of the reference genome.
- **Sd**: Applies only when a reference genome is provided ("." if no reference genome provided or if the variant is unmapped). Strand of the prediction mapping. "1": Forward; "-1": Reverse. **Important Remark**: Fields "REF", "ALT" and "Genome" are based on the mapped predictions. If Sd is 1 then these fields correspond to the DiscoSnp++ prediction, else if the Sd is "-1", then they correspond to the reverse complement of the DiscoSnp++ predictions.
- **FORMAT**: Description of the genotype fields (G1, G2, G3 ...)
  - **GT**: genotype, encodes as allele values with 0 corresponding to the reference and 1 to the alternative. About genotypes:
    - If the separator is a "/" the genotypes are unphased (INDEL, Simple SNP)
    - If the separator is a "|" the genotypes are phased (Close SNPs with the same ID )
  - **DP**: Cumulated depth across samples (sum)
  - **PL**: Phred-scaled Genotype Likelihood (given by DiscoSnp++)

• **AD**: Depth of each allele by sample

# **Examples**

- FILTER Fields:
  - PASS if the variant is mapped at unique position
  - MULTIPLE if the variant is mapped on multiple positions
  - ".": if the variant is unmapped or if no reference genome is provided

#CHROM G4	POS G5	ID	REF	ALT	QUAL	FILTER	INFO	FORMA	T	G1	G2	G3
Pseudomonas	194531	58816	С	G		PASS						
Ty=SNF	P;Rk=0.6	3574;DT=	:1;UL=.;l	JR=.;CL=	.;CR=.;G	enome=C;Sd=1	GT:DP:I	PL:AD	1/1:8:13	35,19,16	8:1,7	
0/0:2:4,	10,44:2,0	0/0:1:4,	7,24:1,0	0/1:8:30	,14,110:6	6,2 0/1:5:54	,9,34:2,3					
Pseudomonas	319075	4768	Α	G		MULTIPLE						
Ty=SNF	P;Rk=0.0	6837;DT=	:0;UL=.;U	JR=.;CL=	.;CR=.;G	enome=A;Sd=-1	GT:DP:I	PL:AD (	0/1:125	4:8182,	28,9459:6	659,595
0/1:123	5:9809,4	2,7594:56	32,673	0/1:903:	7217,37,	5521:409,494	0/1:473:	4217,52,2	520:19	4,279		
0/1:141	6:10513,	26,9395:6	880,736									

• **VCF for IGV :** If you want to use IGV (Integrative Genomics Viewer) to visualize your data, VCF\_creator can create a vcf specific to IGV (0-based, sorted by positions, and without unmapped variants)

#CHROM	POS	ID	REF	ALT	QUAL	FILTER	INFO	FORMA	T	G1	G2	G3	
<b>G4</b>	G5												
Pseudomonas	1206	7667_1	G	Т		PASS							
Ty=SN	P;Rk=0.1	0501;DT=	:1;UL=.;U	IR=.;CL=	.;CR=.;C	Genome=G;Sd=-1	GT:DP:	PL:AD (	0 1:115:	1172,36	,513:41,7	'4	0
1:142:1154,19,8	75:64,78	0 1:90:7	07,16,58	7:42,48	0 1:111	:1190,43,452:37,7	4	0 1:102:9	60,26,5	21:40,62	<u> </u>		
Pseudomonas	1210	7667_2	Т	G		PASS							
Ty=SN	P;Rk=0.1	0501;DT=	:1;UL=.;U	IR=.;CL=	.;CR=.;C	Genome=T;Sd=-1	GT:DP:	PL:AD (	0 1:115:	1172,36	,513:41,7	'4	0
1:142:1154,19,8	75:64,78	0 1:90:7	07,16,58	7:42,48	0 1:111	:1190,43,452:37,7	4	0 1:102:9	60,26,5	21:40,62	<u> </u>		
Pseudomonas	1214	8588	T	G		PASS							
,		•		•		Genome=T;Sd=1	GT:DP:	PL:AD (	0/0:1059	9:32,275	5,20328:	1038,21	
		8,25081:1 ,19237:97		0/0:830:	21,2204	,16026:816,14	0/0:907	14,2486,1	7676:89	6,11			
Pseudomonas	1217	12549_2	2A	G		PASS							
Ty=SNI	P;Rk=0.0	5280;DT=	:1;UL=.;U	IR=.;CL=	.;CR=.;G	Genome=A;Sd=1	GT:DP:	PL:AD (	0 1:284:	1659,27	,2378:160	0,124	0
1:379:2586,19,2	766:194,	185	0 1:231:	1509,19,	1768:12	2,109 0 1:321:	1852,30,	2710:182,	139	0			
1:260:1820,17,1	860:131,	129											
Pseudomonas	1237	10749	T	G		PASS							
Ty=SN	P;Rk=0.0	2921;DT=	:0;UL=.;U	IR=.;CL=	.;CR=.;C	Senome=T;Sd=1	GT:DP:	PL:AD (	0/0:995:	14,2806	,19551:98	87,8	
0/0:125	2:15,355	0,24642:1	1243,9	0/0:784:	14,2240	,15461:779,5	0/0:812	31,2420,1	6195:81	1,1			
	•	,18615:93	38,6										
Pseudomonas	1243	860	T	G		PASS							
· · · · · · · · · · · · · · · · · · ·						, , - ,	GT:DP:				5,19811:	1000,8	
		5,24542:1		0/0:775:	19,2258	,15366:772,3	0/0:782	13,2213,1	5380:77	6,6			
		,18240:91											
Pseudomonas		14601		Α		PASS							
	•			•		Senome=C;Sd=1	GT:DP:			,	6,29017:	1455,3	
	•	0,32264:1 1,24253:1	-	0/0:1044	4:20,301	7,20654:1039,5	0/0:101	3:48,3054,	20264:1	013,0			
Pseudomonas	6702	8511_1	Т	G		PASS							

Ty=SNP;Rk=0.05088;DT=1;UL=.;UR=.;CL=.;CR=.;Genome=T;Sd=1 GT:DP:PL:AD 0|1:200:1360,17,1479:103,97 0| 1:196:1177,22,1616:109,87 0|1:118:843,16,843:59,59 0|1:182:1041,25,1560:104,78 0|1:160:1040,18,1239:85,75