## Manual For FamSeq

## Synopsis

FamSeq vcf -vcfFile input.vcf -pedFile input.ped -output output.vcf

FamSeq LK -lkFile lk.txt -pedFile input.ped -output output.txt

## **Commands and Options**

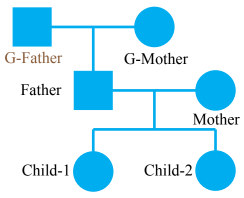
vcf

FamSeq vcf [-method 1] [-mRate 1e-7] [-v] [-a] [-l] [-vcfFile ] [-pedFile ] [-output ] [-LRC ] [-genoProbN] [-genoProbK] [-genoProbXK] [numBurnIn] [numRep]

Call variant when the input data is in a vcf file. The likelihood is extracted from the PL or GL section in the vcf file. The likelihood in the PL or GL section should be phred-scaled or log10 scaled.

Options:	DT	Description
-method	1	The method used in variant calling. It is an integer.
		1(default): Bayesian network. It works well when
		family size is less than seven. 2: Elston-Stewart
		algorithm. Use this method when family size is
-mRate	F	larger than 7 and the family has no loop. 3. MCMC. Mutation rate. It is a float. The default value is 1e-7.
-IIII\ale	,	Only record the position at which the genotype is
•		not RR in the output file. (R: reference allele, A: alternative allele).
-a		Record all the positions in the output file. If there is
		an indel at one position, FamSeq will write the
		same line in input vcf file to output vcf file. The
		number of lines in input vcf file and output vcf file
		are the same. If option -v is set, option -a will be
		discarded. If neither 'v' or 'a' is set, FamSeq will record all the positions except the indel positions.
-vcfFile	STR	The name of input vcf file. All the individuals must
VOI. 110	0771	be in one vcf file.
-pedFile	STR	The name of ped file that store pedigree
		information. The pedigree should be a full family,
		which means that everyone in the family has
		two parents except for the founders of the
		family. There are five columns in the ped file. The
		first column is individual id that should be larger than 0. The second and third column is mother's id
		and father's id. If the individual is the founder of the
		family, set the mother and father's id to 0. The forth
		column is gender. 1: male and 2: female. It will
		cause some errors at X chromosomes if the gender

is not set correctly. The last column is individual name in vcf/likelihood only format file. If there is no information of an individual in vcf/likelihood only format file, set the individual name to NA in the ped file. Example:



This is a family of six individuals. All individuals other than the grandfather were sequenced. The vcf file looks like the following:

#CH	ROM POS	ID	REF	ALT QUAI	FILTER INFO	FORMAT Child-1	Child-2	Father G-Mother	Mother
1	1337418 .	T		3924.52 PASS	AC=0;AF=0.00;	AN=10;DP=922;MQ=	35.47;MQ0=0	GT:DP 0/0:196	6 0/0:185 0/0:107 0/0:253 0/0:181
1	1337419 .	G		3947.61 PASS	AC=0;AF=0.00;	AN=10;DP=937;MQ=	35.48;MQ0=0	GT:DP 0/0:19	8 0/0:190 0/0:107 0/0:258 0/0:184
1	1337420 .	C		4081.22 PASS	AC=0;AF=0.00;	AN=10;DP=963;MQ=	35.51;MQ0=0	GT:DP 0/0:20	3 0/0:195 0/0:109 0/0:268 0/0:188
1	1337421 .	A		4024.57 PASS	AC=0;AF=0.00;	AN=10;DP=976;MQ=	35.53;MQ0=0	GT:DP 0/0:20:	5 0/0:196 0/0:112 0/0:273 0/0:190
1	1337422 .	A	-	4291.47 PASS	AC=0;AF=0.00;	AN=10;DP=993;MQ=	35.55;MQ0=0	GT:DP 0/0:200	5 0/0:196 0/0:116 0/0:281 0/0:194
1	1337423 .	A		4584.69 PASS	AC=0;AF=0.00;	AN=10;DP=1003;MQ	=35.57;MQ0=0	GT:DP 0/0:20	6 0/0:197 0/0:121 0/0:284 0/0:195
1	1337424 .	T		4478.19 PASS	AC=0;AF=0.00;	AN=10;DP=1007;MQ	=35.57;MQ0=0	GT:DP 0/0:20	8 0/0:197 0/0:122 0/0:285 0/0:195
1	1337425	Т		4678.45 PASS	AC=0:AF=0.00:	AN=10:DP=1016:MO	=35.59:MO0=0	GT:DP 0/0:20	8 0/0:199 0/0:123 0/0:289 0/0:197

Then we construct the corresponding ped file. Make sure the individual name in the ped file is the same as in the vcf file. The grandfather should be included in ped file with individual name NA, even though there is no information about him in the vcf file.

ID	mID	fID gender IndividualName				
1	0	0	2	G-Mother		
2	0	0	1	NA		
3	1	2	1	Father		
4	0	0	2	Mother		
5	4	3	2	Child-1		
6	4	3	2	Child-2		

#CHROM POS 1 1337418 . 1 1337419 . 1 1337420 . 1 1337421 . 1 1337422 . 1 1337423 . 1 1337424 . 1 1337425 .	T G C A A T T	3947.61 PASS A 4081.22 PASS A 4024.57 PASS A 4291.47 PASS A 4584.69 PASS A 4478.19 PASS A	C=0;AF=0.00 .C=0;AF=0.00 .C=0;AF=0.00 .C=0;AF=0.00 .C=0;AF=0.00 .C=0;AF=0.00 .C=0;AF=0.00	O FORMAT Child-1 Child-2 Father G-Mother Mother  0;AN=10;DP=922;MQ=35.47;MQ0=0 GT:DP 0/0:196 0/0:185 0/0:107 0/0:253 0/0:181 0;AN=10;DP=9337;MQ=35.51;MQ0=0 GT:DP 0/0:198 0/0:190 0/0:107 0/0:258 0/0:184 0;AN=10;DP=963;MQ=35.51;MQ0=0 GT:DP 0/0:203 0/0:195 0/0:109 0/0:268 0/0:188 0;AN=10;DP=976;MQ=35.55;MQ0=0 GT:DP 0/0:205 0/0:196 0/0:112 0/0:273 0/0:190 0;AN=10;DP=1003;MQ=35.57;MQ0=0 GT:DP 0/0:206 0/0:196 0/0:116 0/0:281 0/0:194 0;AN=10;DP=1003;MQ=35.57;MQ0=0 GT:DP 0/0:208 0/0:197 0/0:212 0/0:284 0/0:195 0;AN=10;DP=1001;MQ=35.57;MQ0=0 GT:DP 0/0:208 0/0:197 0/0:121 0/0:284 0/0:195 0;AN=10;DP=1016;MQ=35.59;MQ0=0 GT:DP 0/0:208 0/0:199 0/0:228 0/0:197
		-output	STR	Output file name. If FamSeq calls a variant at a
				position, it will add two tags (FGT: genotype called by FamSeq and FPP: posterior probability estimated by FamSeq) at column FORMAT in vcf file.
		-LRC	F	A likelihood ratio cutoff. If likelihood (most likely genotype)/sum(likelihood of all genotypes) is less than the cutoff, we use pedigree information to improve variant calling. The default value is 1, we call all variant using pedigree information. Set it to 0 to only use single individual based method. Any values in between will determine whether FamSeq or single method is used for variant calling at a position.
-genoProbN -genoProbK		FFF	Genotype probability of three kinds of genotype for autosome in population (Pr(G)) when this position is not in dbSNP. The default values are: 0.9985, 0.001 and 0.0005. The dbSNP position should be provided in column 'ID' in input vcf file.	
		-genoProbK	FFF	Genotype probability of three kinds of genotype for autosome in population (Pr(G)) when the position is in dbSNP. The default values are: 0.45, 0.1 and 0.45.
		-genoProbXN	FF	Genotype probability of two kinds of genotype for chromosome X for male in population (Pr(G)) when the variant is not in dbSNP. The default values are: 0.999 and 0.001.
		-genoProbXK	FF	Genotype probability of two kinds of genotype for chromosome X for male in population (Pr(G)) when the variant is in dbSNP. The default values are: 0.5 and 0.5.
		-numBurnIn	1	Number of burn in when the user chooses the MCMC method. The default value is 1,000 <i>n</i> , where <i>n</i> is the number of individuals in the pedigree.
		-numRep	1	Number of iteration times when the user chooses MCMC method. The default value is 20,000 <i>n</i> .

[-lkFile ] [-pedFile ] [-output ] [-LRC ] [-genoProbN] [-genoProbXK] [-genoProbXK]

Call variants when the input data is in a likelihood only format file.

Options: DT Description

-lkFile: STR The name of input likelihood file. The first row is the

individual name. The likelihood for each individual starts from the second row. Each column represents one individual. In each column, the likelihood for three kinds of genotype are sperated

by comma.

Child1	Child2	Father	G-Mother	Mother						
2.69e-06	5,0.209,0	.0494	5.5e-12,0.0038	7,0.153	3.49e-08,0.0193	3,0.003 0.1,0.	0985,1.38e-07	0.0239,0.11	5,9.18e-07	
0.427,0.	0331,1.2	4e-09	0.0032,0.272,9.	12e-05	0.000354,0.395,	,0.00254 0.1,0.	0985,1.38e-07	0.00466,0.3	332,0.000114	
0.203,0.	079,3.53	e-08	0.000781,0.38,0	0.00102	4.75e-07,0.137,0	0.0786 8.72e	-09,0.0479,0.183	6.33e-05,0.	341,0.0091	
0.376,0.	0432,3.1	1e-09	0.505,0.00191,2	2.02e-13	0.268,0.0165,2.4	45e-10 0.392	,0.000875,2.48e-14	0.0309,2.41	e-06,6.6e-21	
0.0493,0	0.0898,2.	13e-07	1.19e-07,0.0399	9,0.00772	0.000791,0.0085	5,1.13e-08 0.011	,0.209,1.21e-05	0.000354,0	.395,0.00254	
3.25e-07	7,0.107,0	.0541	3.34e-08,0.0059	98,9.33e-05	0.18,0.	0498,1e-08	2.69e-06,0.209,0.	0494 0.0	000861,0.209,0.000	)154

-lkType STR The likelihood type. There are four types of

likelihood: Normal (n), log10 scaled (log10), ln scaled (ln) and phred scaled (PS). The figure shown above is type n, without any scale.

DT: Data Type. *I*: integer. *F*: float value. *STR*: string.

## Output

FamSeq creates a new file by adding three columns to the original input file as the output file: GPP, FPP and FGT. GPP is the posterior probability calculated by single individual based method and FPP is the posterior probability calculated by FamSeq. These probabilities are all Phred-scaled. FGT is the genotype called by FamSeq.

Version: 1.0.3