# ANGSD formats

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# 1 SAF formats

SAF files are files that contain sample allele frequency. These are generated with -doSaf in main ANGSD. These contains either the loglikelihood ratio to the most likely category or the pp. This is determined if the -prior has been supplied. The first 8 bytes magic number determines which SAF version. If no magic number is present then version0 is assumed.

#### 1.1 version 0

First version of the SAF files were simply flat binary double files PREFIX.saf along with an associated PREFIX.saf.pos.gz which contains the gzip compressed 'chromosome position'. Assuming nChr number of chromosomes, then we have nChr+1 categories for each site. The number of sites can therefore be deduced either directly from the number of lines in the uncompressed output of the PREFIX.saf.pos.gz, or by using the filesize (fsize) of the PREFIX.saf

$$\frac{fsize}{size of(double)*(nChr+1)}.$$

### 1.2 version 1

Second iteration of the saf files now contains two raw files and an index file. Still under development. First 8 bytes in all three files is 8byte magic numer char[8] "safv3".

PREFIX.saf.gz bgzf compressed flat floats. With similar interpretation as version0. Each element is a cdatatype 'float' which is 4 bytes.

PREFIX.saf.pos.gz bgzf compressed flat integer. Representing the position. Each element is a cdatatype 'int' which is 4bytes

PREFIX.saf.idx uncompressed binary file containing blocks of data described in 1.2. This is preceded by a size\_t value which indicates the number of categories in the spectrum.

Note that it is not possible to deduce the number of sites directly from the compressed files.

$\operatorname{Col}$	$\mathbf{Field}$	$\mathbf{Type}$	Brief description
1	CLEN	$size_t$	Length of CHR (not including terminating null)
2	CHR	$char^*$	Reference sequence name. Length is CLEN
3	<b>NSITES</b>	$size_t$	Number of sites with coverage from reference CHR
4	OFF1	long int	CHR offset into the PREFIX.saf.pos.gz
5	OFF2	long int	CHR offset into the PREFIX.saf.gz

Table 1: Content of entry for a single reference name in the PREFIX.saf.idx file.

# 2 fst formats

This section describes the binary output generated by a **realSFS fst index pop1.saf.idx pop2.saf.idx -sfs prior** 

#### 2.1 fstv1

First iteration of the fst files contains two files. 1) PREFIX.fst.idx 2) PREFIX.fst.gz. First 8bytes is a magic number determining which binary version.

PREFIX.fst.idx flat file, described in table 2.1.1

PREFIX.fst.gz bgzf compressed binary file.

#### 2.1.1 PREFIX.fst.idx

The fst.idx has a very simple header 8bytes magicheader followed by a size\_t containing the number of samples for which we have generated fst results.

Col	Field	Type	Brief description
1	CLEN	$\mathrm{size\_t}$	Length of CHR (not including terminating null)
2	CHR	$char^*$	Reference sequence name. Length is CLEN
3	NSITES	$size_t$	Number of sites with coverage from reference CHR
4	OFF1	long int	CHR offset into the PREFIX.saf.pos.gz

# 2.1.2 PREFIX.fst.gz

Col	Field	Type	Brief description
1	POSI	int	Length of CHR (not including terminating null)
2	acoef1	double*	$\alpha$ coefficients from either reynolds estimator or Bhatia
3	bcoef2	double*	$\beta$ coeffficients from eithre Reynolds estimator or Bhatia

# 2.2 fstv2

PREFIX.fst.idx flat file, described in table 2.1.1

PREFIX.fst.gz bgzf compressed binary file.

# 2.2.1 PREFIX.fst.idx

## 2.2.2 PREFIX.fst.gz

FiXme Fatal: This is still under development

Col	Field	Type	Brief description
1	MAGIC	char[8]	magic nr = fstv2
2	NPOP	$\operatorname{size\_t}$	Number of populations
3	NPOP_LEN_i	size_t	length of population name_i
4	NPOP_i	$char[NPOP\_LEN]$	name of population_i.
5	NDIM_j	size_t	number of categories for pair $j$
6	PRIOR_j	$double[NDIM_{-j}]$	prior for pair $j$
6	AS_j	double[NDIM_j]	alfa for pair $j$
6	BS_j	$double[NDIM\_j]$	beta for pair $j$
7	CLEN	size_t	Length of CHR (not including terminating null)
8	CHR	char*	Reference sequence name. Length is CLEN
9	NSITES	$size\_t$	Number of sites with coverage from reference CHR
10	OFF1	long int	CHR offset into the PREFIX.saf.pos.gz

Table 2: Format of saf.idx file. i loops over number of populations. j loops over number of combinations, in this order pop1/pop2,pop1/pop3,pop1/pop4,pop2/pop3,....

Col	Field	Type	Brief description
1	POSI	int	Length of CHR (not including terminating null)
2	acoef1	$char^*$	Reference sequence name. Length is CLEN
3	bcoef2	$size_t$	Number of sites with coverage from reference CHR