	Field	Descrption	Type	Value
magic		Magic string	char[4]	BCF\4
I_seqnm		Length of concatenated sequence names	int32_t	
seqnm		Concatenated names, NULL padded	char[l_seqnm]	
l_smpl		Length of concatenated sample names	int32_t	
smpl		Concatenated sample names	char[l_smpl]	
I_meta		Length of the meta text (double-hash lines)	$int32_t$	
meta		Meta text, NULL terminated	char[l_meta]	
List of records until the end of the file				
	seq_id	Reference sequence ID	int32_t	
	pos	Position	int32_t	
	qual	Variant quality	float	
	l_str	Length of str	int32_t	
	str	ID+REF+ALT+FILTER+INFO+FORMAT, NULL padded	char[l_str]	
	Blocks of data; #blocks and formats defined by FORMAT (table)

Field	Type	Description
DP	uint16_t[n]	Read depth
GL	float[n*G]	Log10 likelihood of data; $G = \frac{A(A+1)}{2}$, $A = \#\{alleles\}$
GT	$uint8_t[n]$	missing<<7 phased<<6 allele1<<3 allele2
$_{\tt GT}$	$uint8_t+uint8_t[n*P]$	Generic GT; the first int equals the max ploidy P . If the
		highest bit is set, the allele is not present (e.g. due to
		different ploidy between samples).
GQ	$uint8_t[n]$	Genotype quality
HQ	$uint8_t[n*2]$	Haplotype quality
$_{-}$ HQ	$uint8_t+uint8_t[n*P]$	Generic HQ
IBD	$uint32_t[n*2]$	IBD
$_{ t L}{ t IBD}$	$uint8_t+uint32_t[n*P]$	Generic IBD
PL	$uint8_t[n*G]$	Phred-scaled likelihood of data
PS	$uint32_t[n]$	Phase set
Integer	$int32_t[n*X]$	Fix-sized custom Integer; X defined in the header
Numeric	double[n*X]	Fix-sized custom Numeric
String	uint32_t+char*	NULL padded concat. strings (int equals to the length)

- A BCF file is in the BGZF format.
- All multi-byte numbers are little-endian.
- In a string, a missing value '.' is an empty C string "\0" (not ".\0")
- For GL and PL, likelihoods of genotypes appear in the order of alleles in REF and then ALT. For example, if REF=C, ALT=T, A, likelihoods appear in the order of CC, CT, TT, CA, TA, AA (NB: the ordering is different from the one in the original BCF proposal).
- Predefined FORMAT fields can be missing from VCF headers, but custom FORMAT fields are required to be explicitly defined in the headers.
- A FORMAT field with its name starting with '_' is specific to BCF only. It gives an alternative binary representation of the corresponding VCF field, in case the default representation is unable to keep the genotype information, for example, when the ploidy is not 2 or there are more than 8 alleles.