# NGS data anlysis course

**Association Analysis using PLINK** 

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### File formats: VCF

#### Tab delimited text file with a **header section**

```
##fileformat=VCFv4.2
##fileDate=20090805
##source=mvImputationProgramV3.1
##reference=file:///seg/references/1000GenomesPilot-NCBI36.fasta
##contig=<ID=20,length=62435964,assembly=B36,md5=f126cdf8a6e0c7f379d618ff66beb2da,species="Homo sapiens",taxonomy=x>
##phasing=partial
##INFO=<ID=NS.Number=1.Type=Integer.Description="Number of Samples With Data">
##INFO=<ID=DP.Number=1.Type=Integer.Description="Total Depth">
##INFO=<ID=AF.Number=A.Type=Float.Description="Allele Frequency">
##INFO=<ID=AA, Number=1, Type=String, Description="Ancestral Allele">
##INFO=<ID=DB,Number=0,Type=Flag,Description="dbSNP membership, build 129">
##INFO=<ID=H2.Number=0.Type=Flag.Description="HapMap2 membership">
##FILTER=<ID=q10.Description="Quality below 10">
##FILTER=<ID=s50,Description="Less than 50% of samples have data">
##FORMAT=<ID=GT, Number=1, Type=String, Description="Genotype">
##FORMAT=<ID=GQ.Number=1.Type=Integer.Description="Genotype Quality">
##FORMAT=<ID=DP.Number=1.Type=Integer.Description="Read Depth">
##FORMAT=<ID=HQ.Number=2.Type=Integer.Description="Haplotype Quality">
#CHROM POS
                                       QUAL FILTER INFO
                                                                                    FORMAT
                                                                                                NA00001
                                                                                                               NA00002
                                                                                                                             NA00003
      14370
            rs6054257 G
                                       29 PASS NS=3:DP=14:AF=0.5:DB:H2
                                                                                    GT:GQ:DP:HQ 0|0:48:1:51,51 1|0:48:8:51,51 1/1:43:5:.,.
      17330
                                           q10
                                                   NS=3;DP=11;AF=0.017
                                                                                    GT:GQ:DP:HQ 0|0:49:3:58,50 0|1:3:5:65,3 0/0:41:3
                               G.T
20
    1110696 rs6040355 A
                                       67 PASS NS=2:DP=10:AF=0.333.0.667:AA=T:DB GT:GD:DP:HD 1/2:21:6:23.27 2/1:2:0:18.2 2/2:35:4
     1230237 .
                                           PASS NS=3:DP=13:AA=T
                                                                                    GT:GQ:DP:HQ 0|0:54:7:56.60 0|0:48:4:51.51 0/0:61:2
     1234567 microsat1 GTC G.GTCT 50 PASS NS=3:DP=9:AA=G
                                                                                    GT:GD:DP 0/1:35:4
                                                                                                               0/2:17:2
                                                                                                                             1/1:40:3
```

#### May be compressed and indexed using tabix

# File formats: VCF

### Each variant is described by 8 fields

CHROM: chromosome

POS: position

ID: name

REF: reference base(s)

6 ALT: non-reference alleles

QUAL: quality score of the calls (phred scale)

FILTER: PASS / filtering\_tag

INFO: additional information

**Genotype data** for several samples may be included in a batch of additional columns (one for each sample) preceded by a FORMAT column which describes their format.

### File formats: VCF INFO column

May include several semicolon separated fields containing information about the variants coded in key value style:

Some reserved (but optional) keys are:

- AA ancestral allele
- AC allele count in genotypes, for each ALT allele, in the same order as listed
- AF allele frequency
- CIGAR cigar string describing how to align an alternate allele to the reference allele
- DB dbSNP membership
- MQ RMS mapping quality, e.g. MQ=52
- MQ0 Number of MAPQ == 0 reads covering this record

### File formats: PED & MAP

Classic format to represent genomic variants for several individuals

would be represented as TPED/TFAM files:

Some variants of the format are described depending on the software used to read or write them. Those variants may include *transposed* versions of the format which is closer to standard

## File formats: PED & MAP

#### PED file

- Family ID
- Individual ID
- Paternal ID
- Maternal ID
- 5 Sex (1=male; 2=female; other=unknown)
- Phenotype (1=unaffected; 2=affected; 0 missing; -9=missing)
- O ... genotypes ...

#### MAP file

- 1 chromosome (1-22, X, Y or 0 if unplaced)
- 2 rs... or SNP identifier
- Genetic distance (Morgans)
- Base-pair position (bp units)