

# Bulked Segregant Analysis For Fine Mapping Of Genes

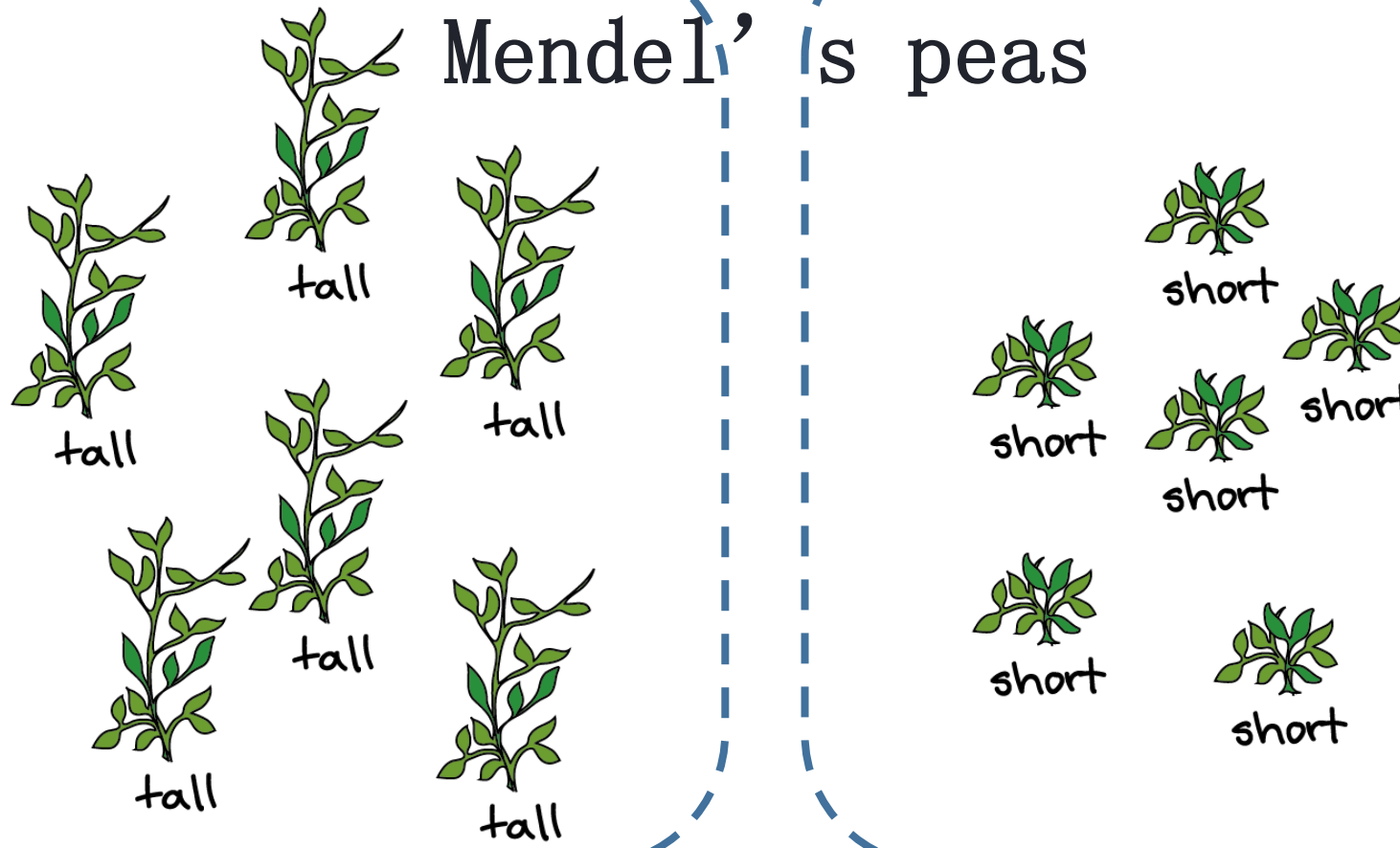
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Bioinformatics Facility  
Cornell University

Alternatively, if you do not have the budget to sequence each individual genome?

Pool them and sequence pools.

(Bulked segregant analysis covered in 2<sup>nd</sup> week)

Mendel's peas

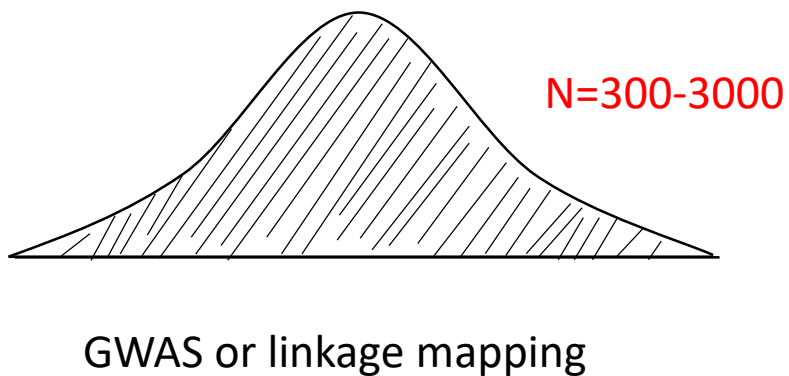


# **Outline**

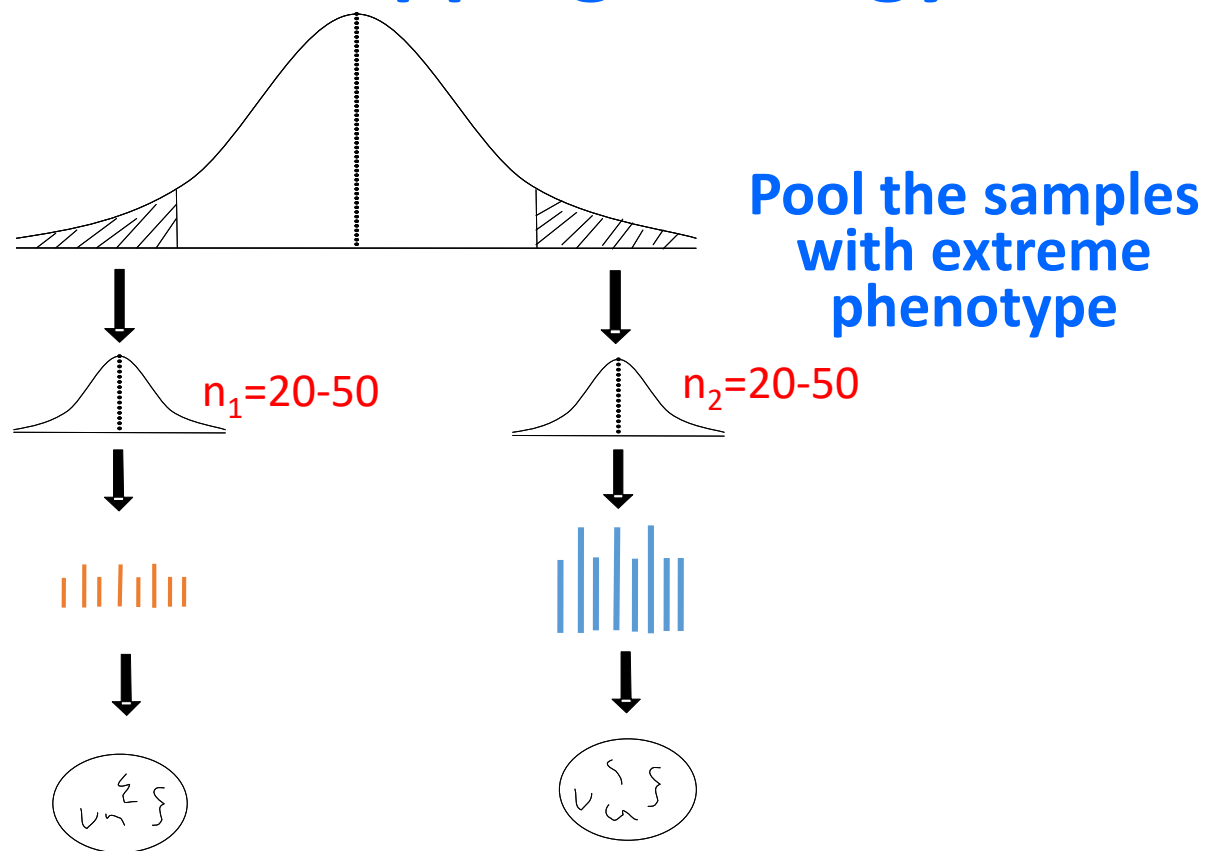
- **What is BSA?**
- **Keys for a successful BSA study**
- **Pipeline of BSA**
- **extended reading**

# Compare BSA with traditional mapping strategy

Entire population  
(all individual)  
analysis



Phenotyping	entire population
Genotyping	entire population

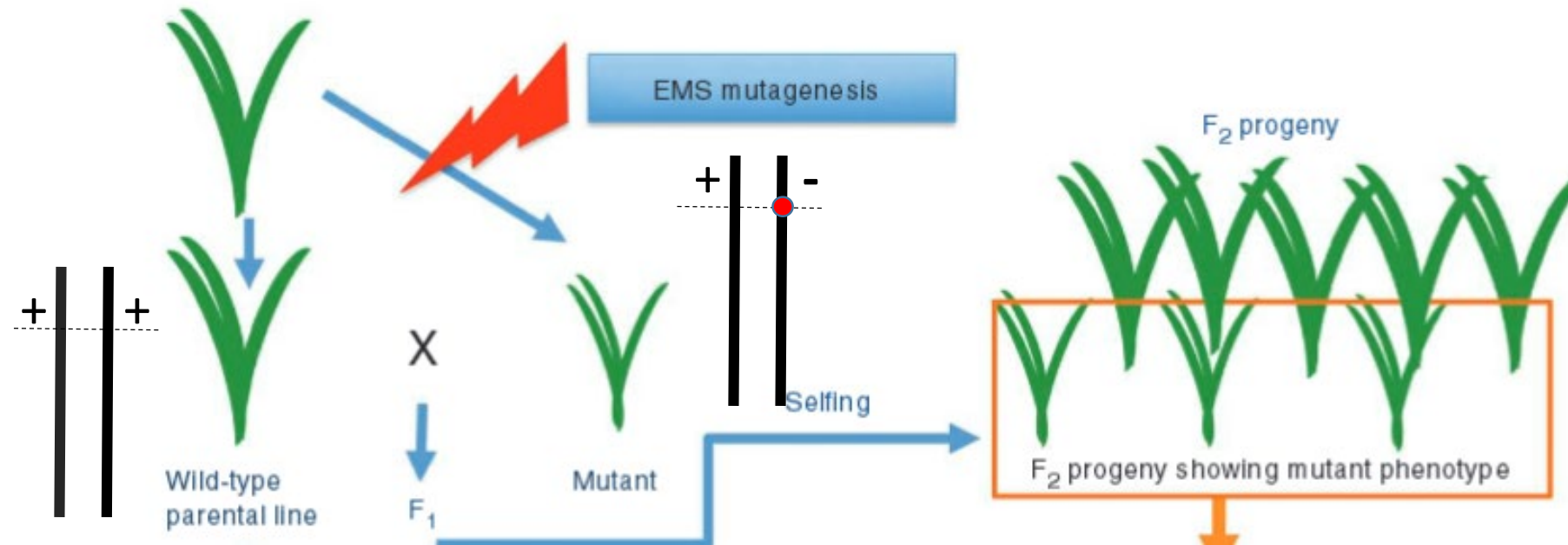


Phenotyping	entire population
Genotyping	two samples

# Bulked Segregant Analysis (BSA)

rapid discovery of genetic markers and trait mapping

## 1. Segregation in phenotype



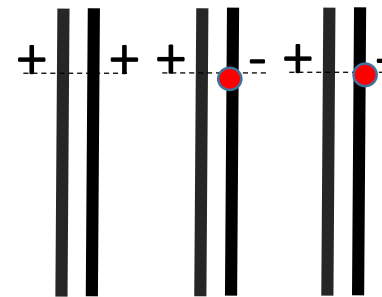
# Bulked Segregant Analysis (BSA)

rapid discovery of genetic markers and trait mapping

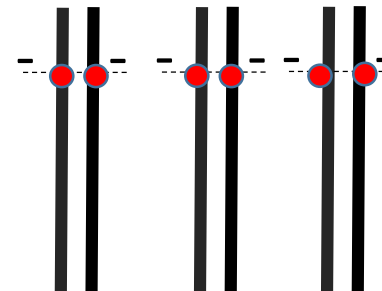
## 2. following Mendelian genetics or determined by a major effect loci



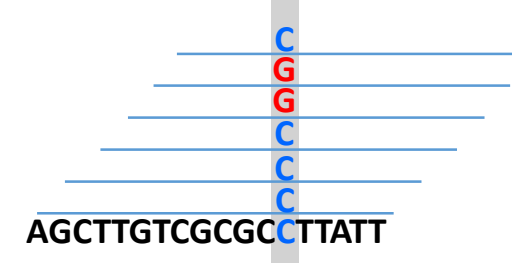
wide type



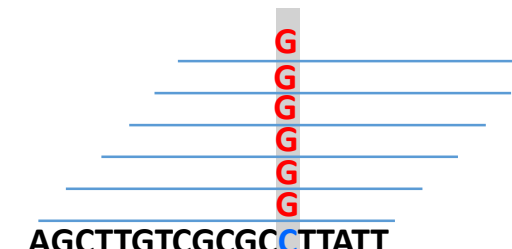
mutant



Linked sites

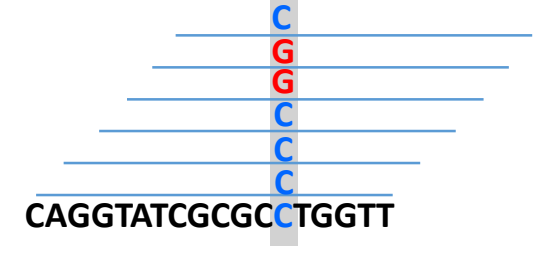


SNP index = 2/6 =0.33

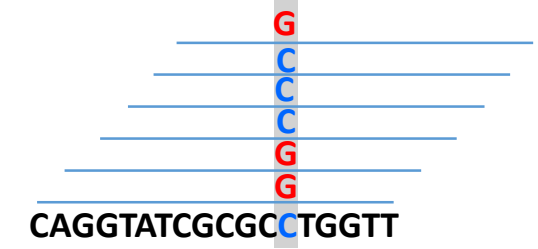


SNP index = 6/6 =1

unlinked sites

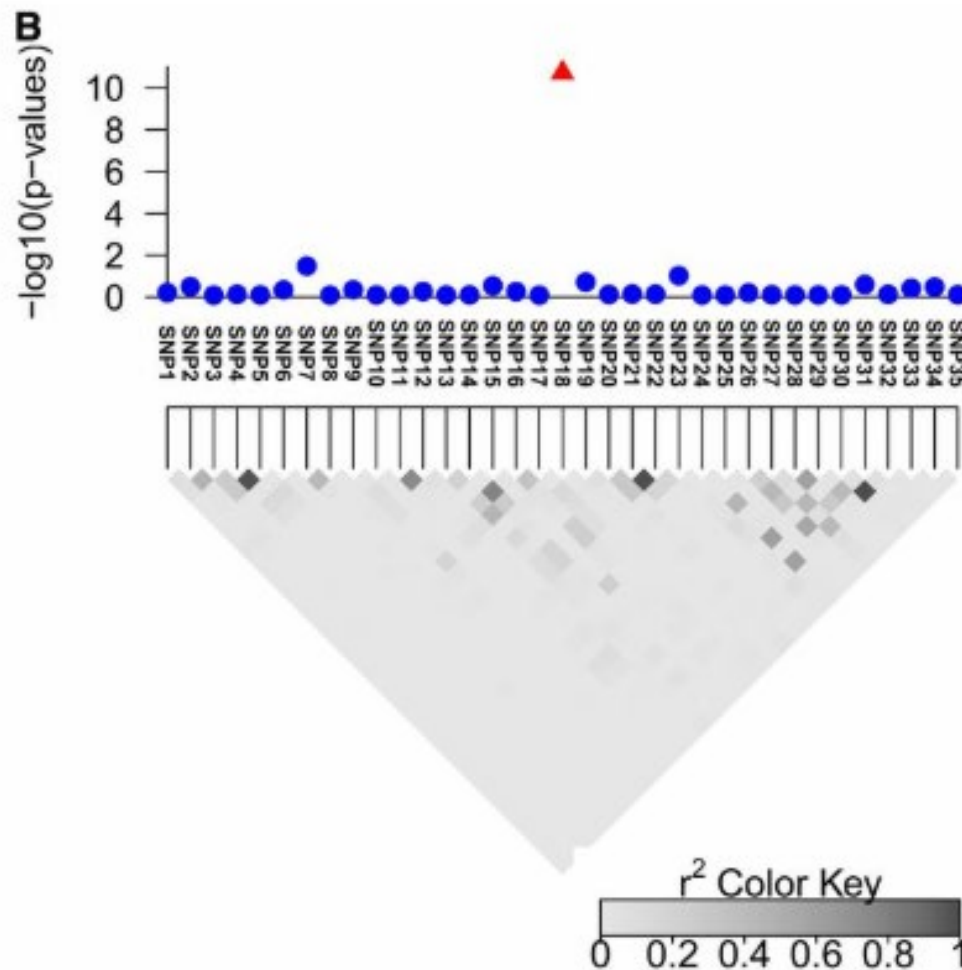
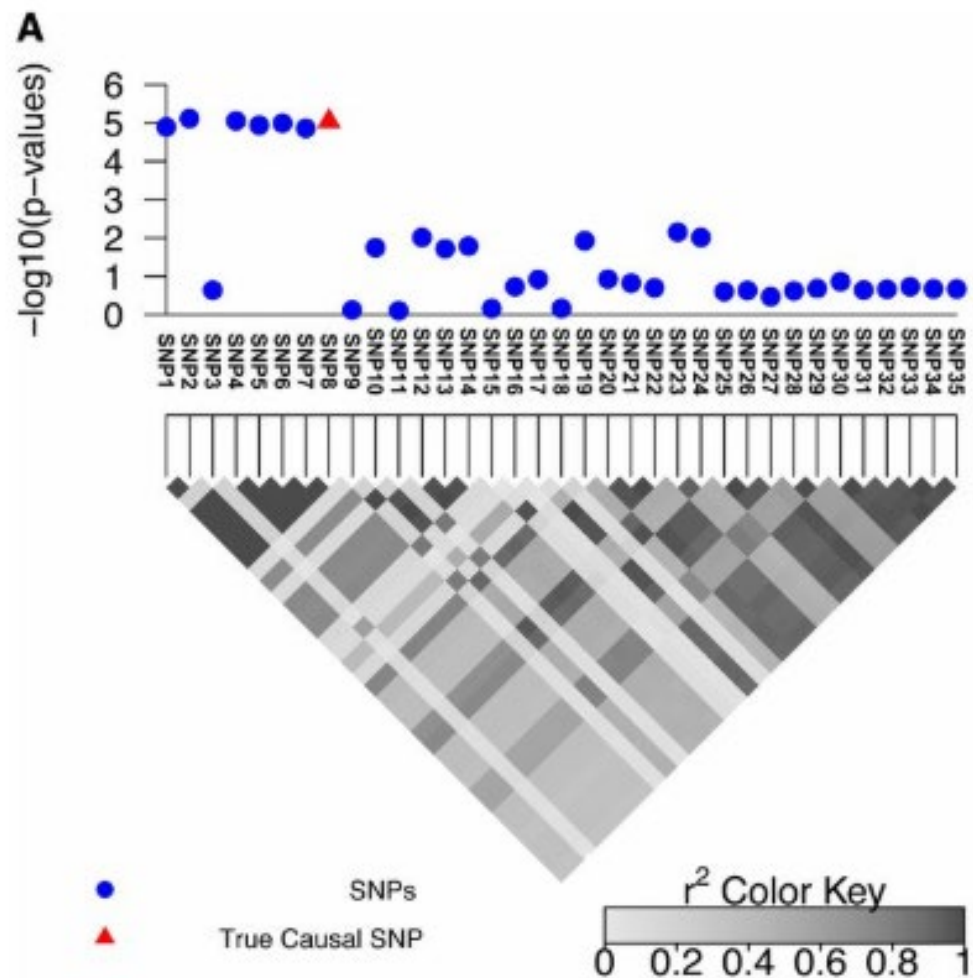


SNP index = 2/6 =0.33



SNP index = 3/6 =0.5

# Causal SNP and SNPs linked with causal SNP

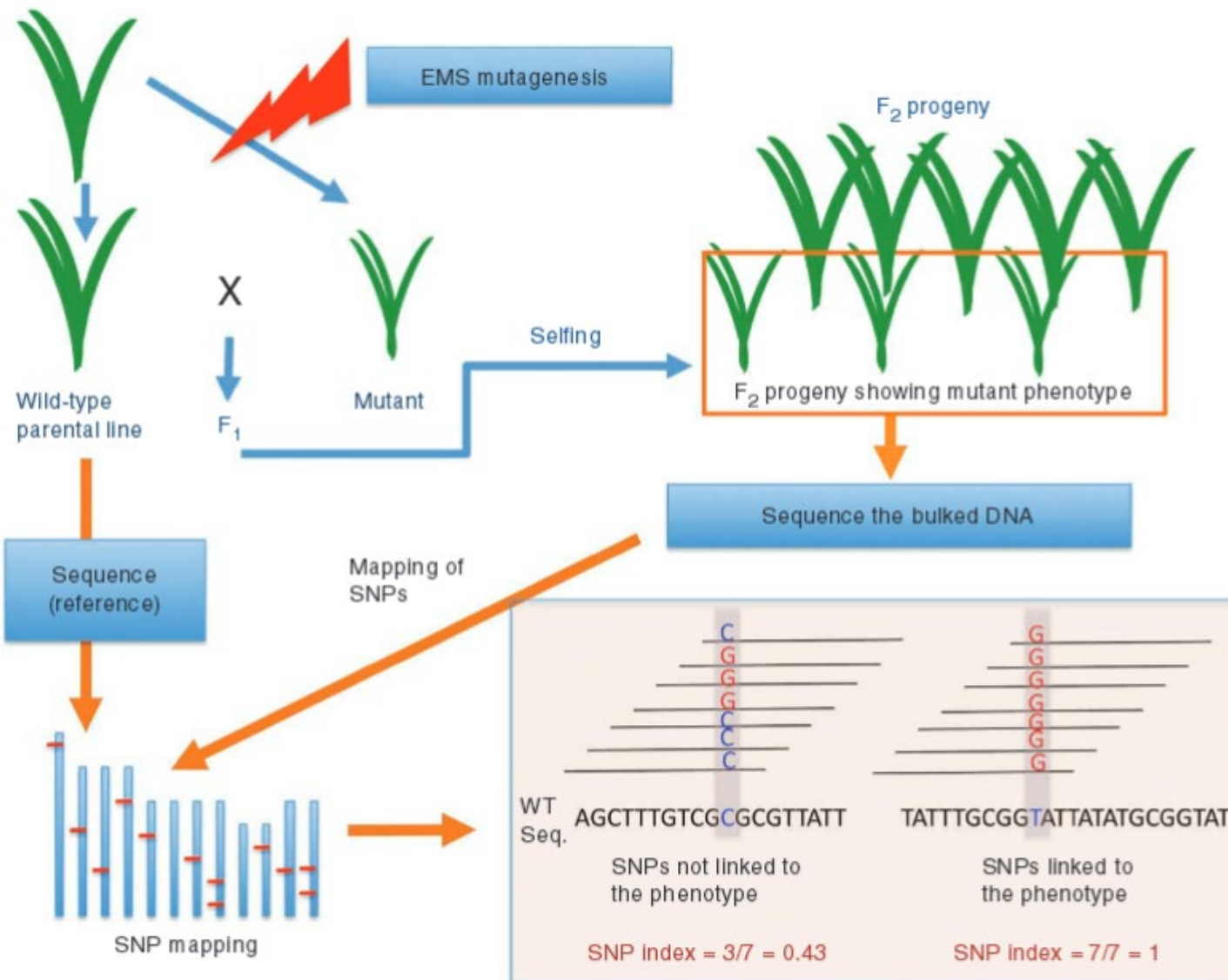


( copy from Hormozdiari, Farhad, et al *Genetics*, 2014)

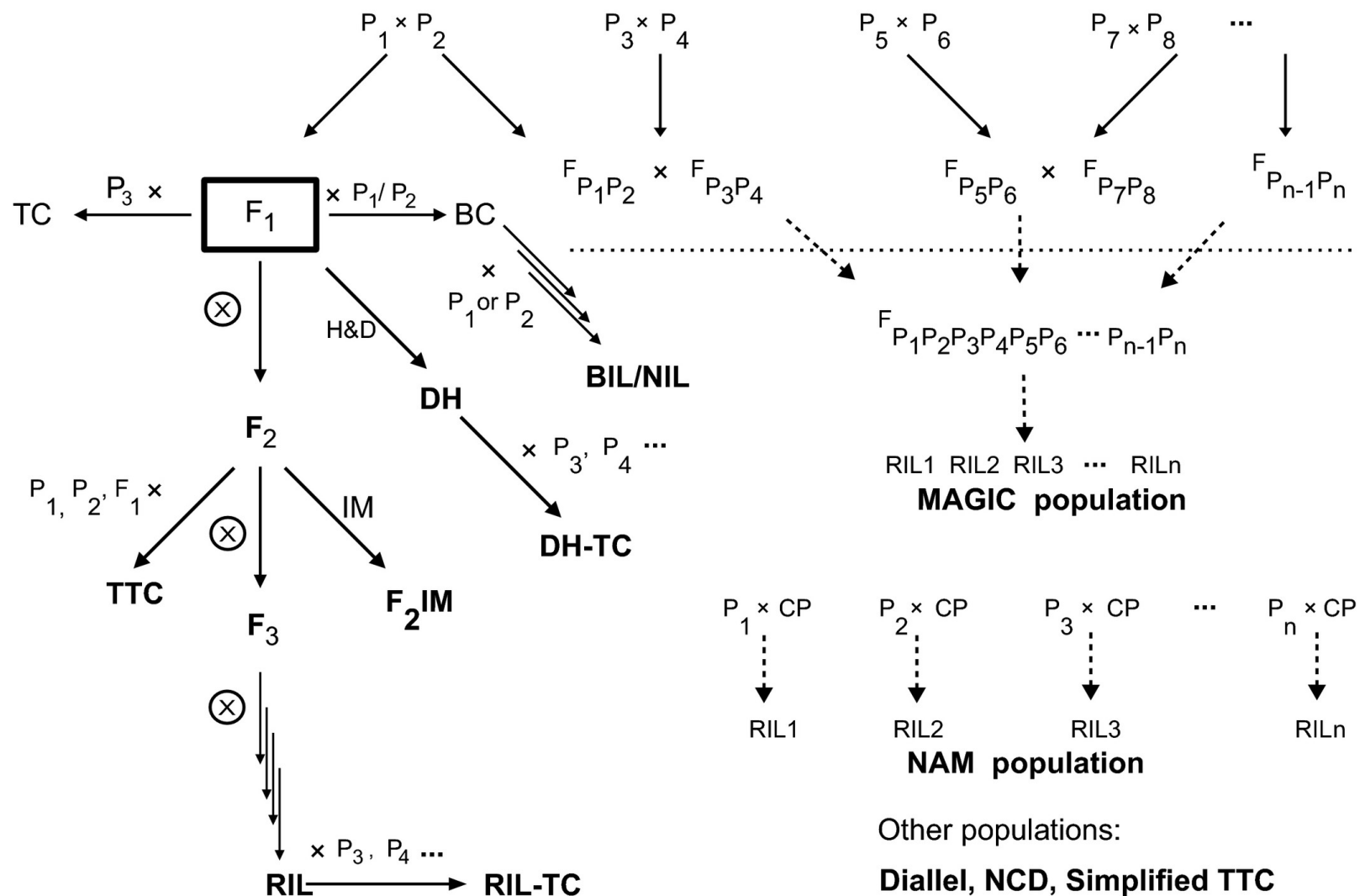
# Applicable populations

- EMS mutagenized population
- Mapping Population
- Natural Population

# EMS mutagenized population



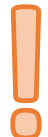
# Examples of Mapping Populations



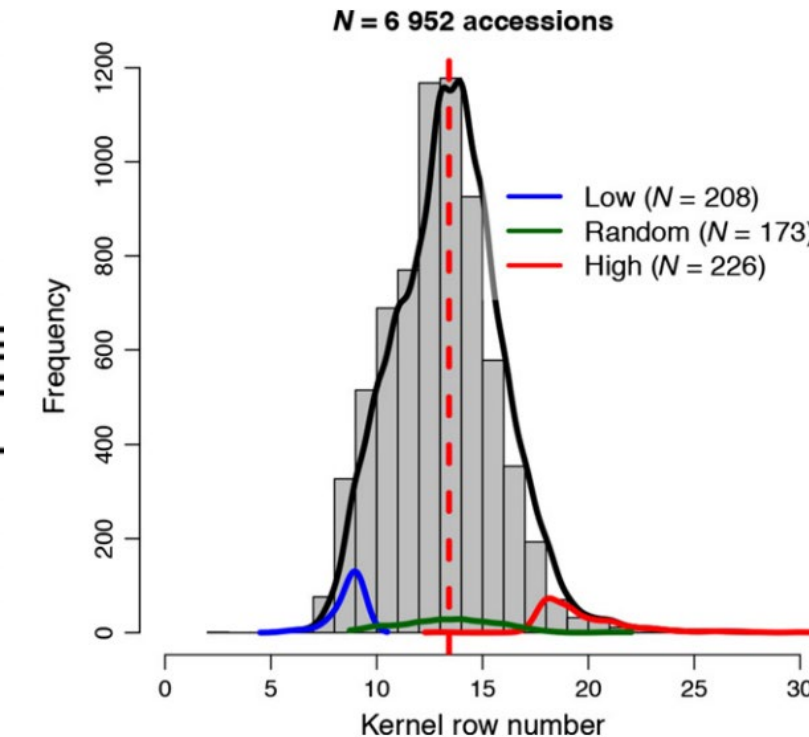
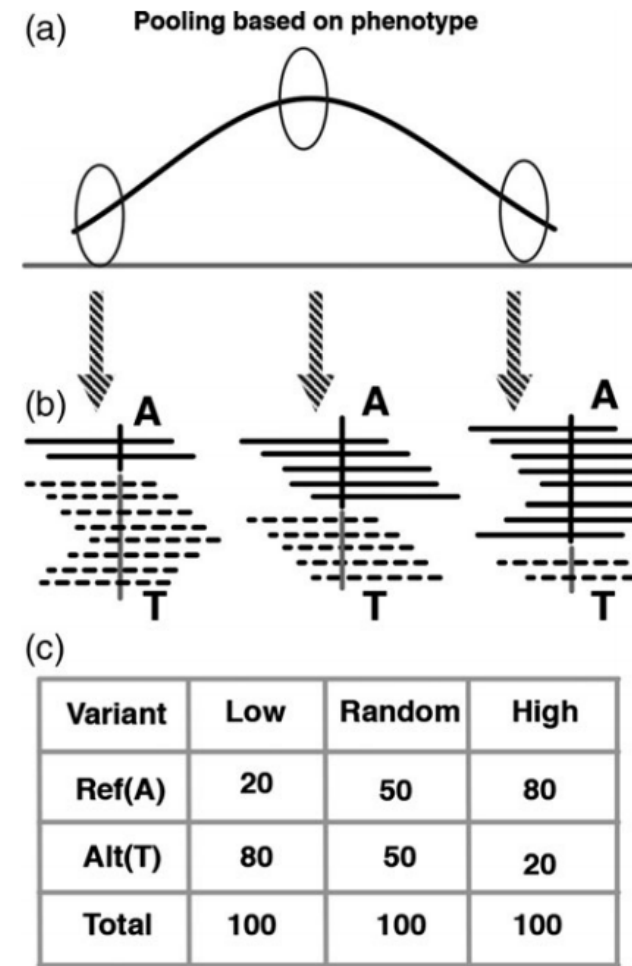
(Zou, 2016 the Plant Biotechnol J)



# Extreme-phenotype GWAS using pooled samples



1. complex genetic architecture of the trait.
2. complex genetic background and population structure



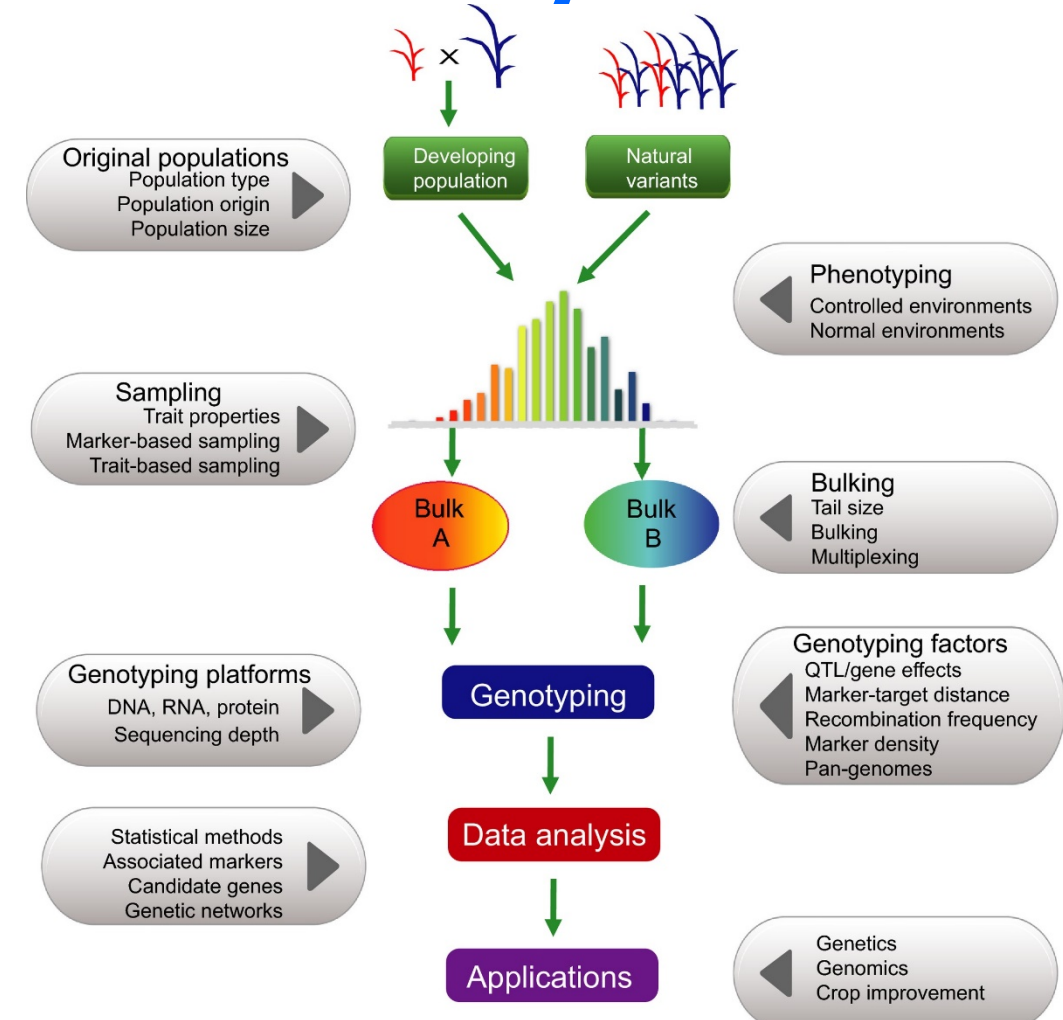
(Schnable, 2015 the Plant Journal)

# Applicable genotyping platform

- Whole genome sequencing
  - High depth sequencing of each bulk (30 ~ 50 X for each pool is recommended, around one fold per each individual )
- RNA-seq based bulk segregant analysis

# Checklist for a successful BSA study

- 1. Genetic architecture and the phenotypic segregation
- 2. Population size, bulk size
- 3. Sequencing depth



# Checklist for a successful BSA study

- BSA has a high false positive rate. Ways to decrease false positives:
- 1. Using replicates or constructing multiple mapping families.
- 2. The genetic background and population characters:
  - EMS mutation: the original G:C pair -> A:T pair
  - Dominance or recessive from genetic study
  - Including parents in the bi-parental family.
- 3. Priori knowledge.
  - Chromosome candidate?
  - RNA-seq expression?
  - linkage map (low density markers)

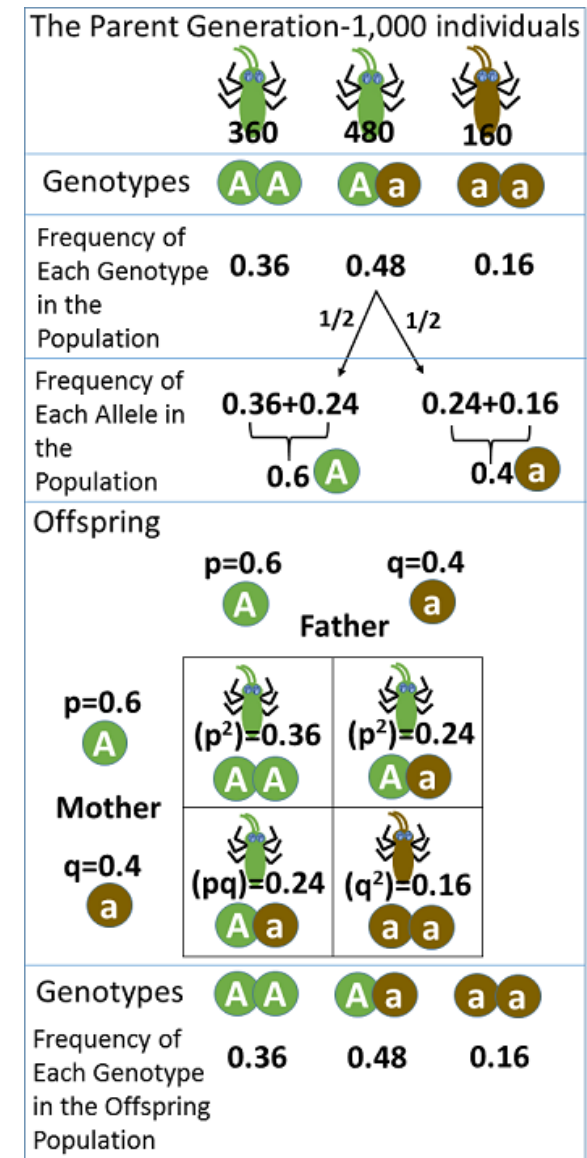
# Beware of Variance Callings

## Assumptions in Variant callers

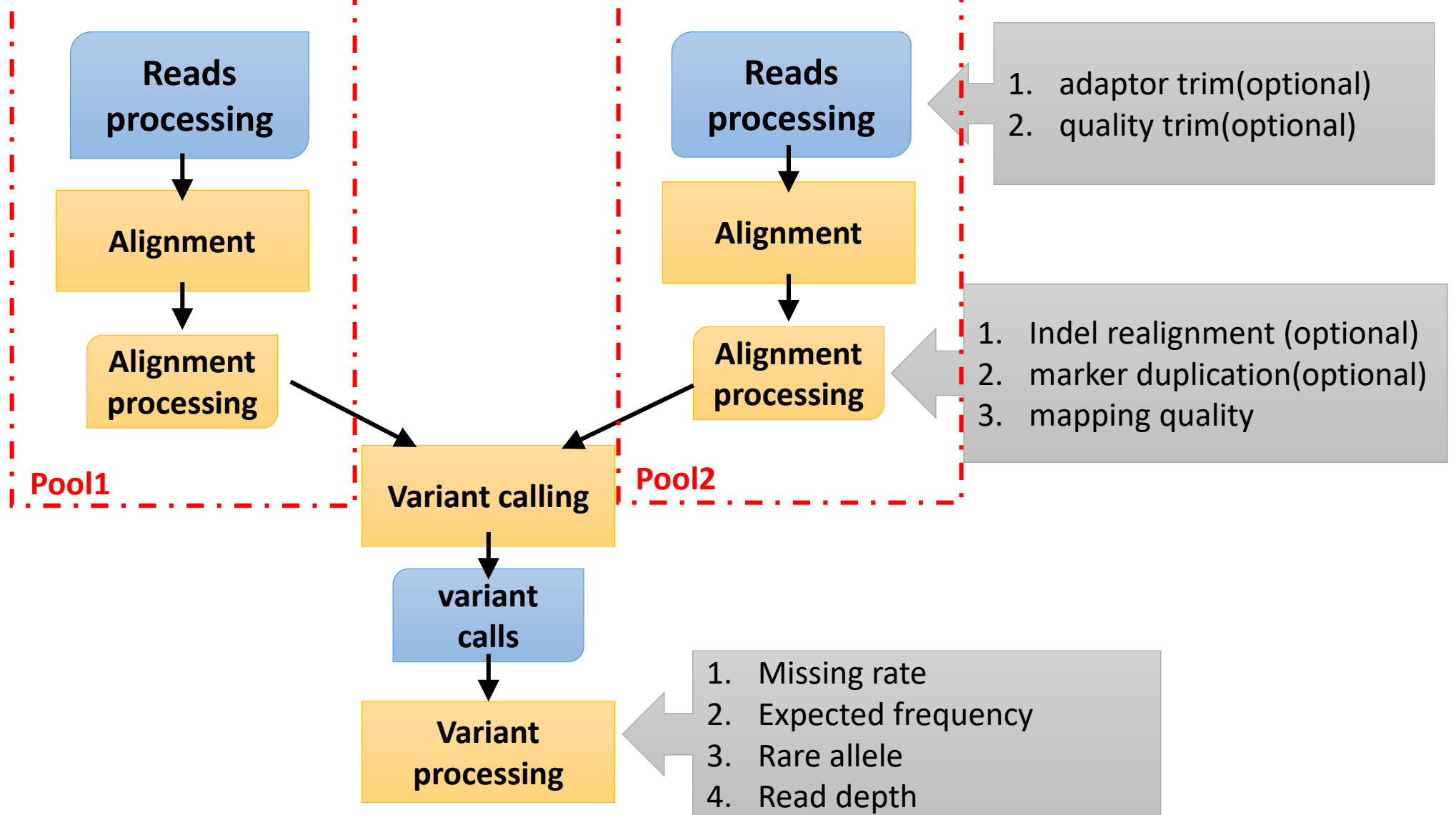
for example GATK :

- assuming Hardy-Weinberg equilibrium
- diploidy

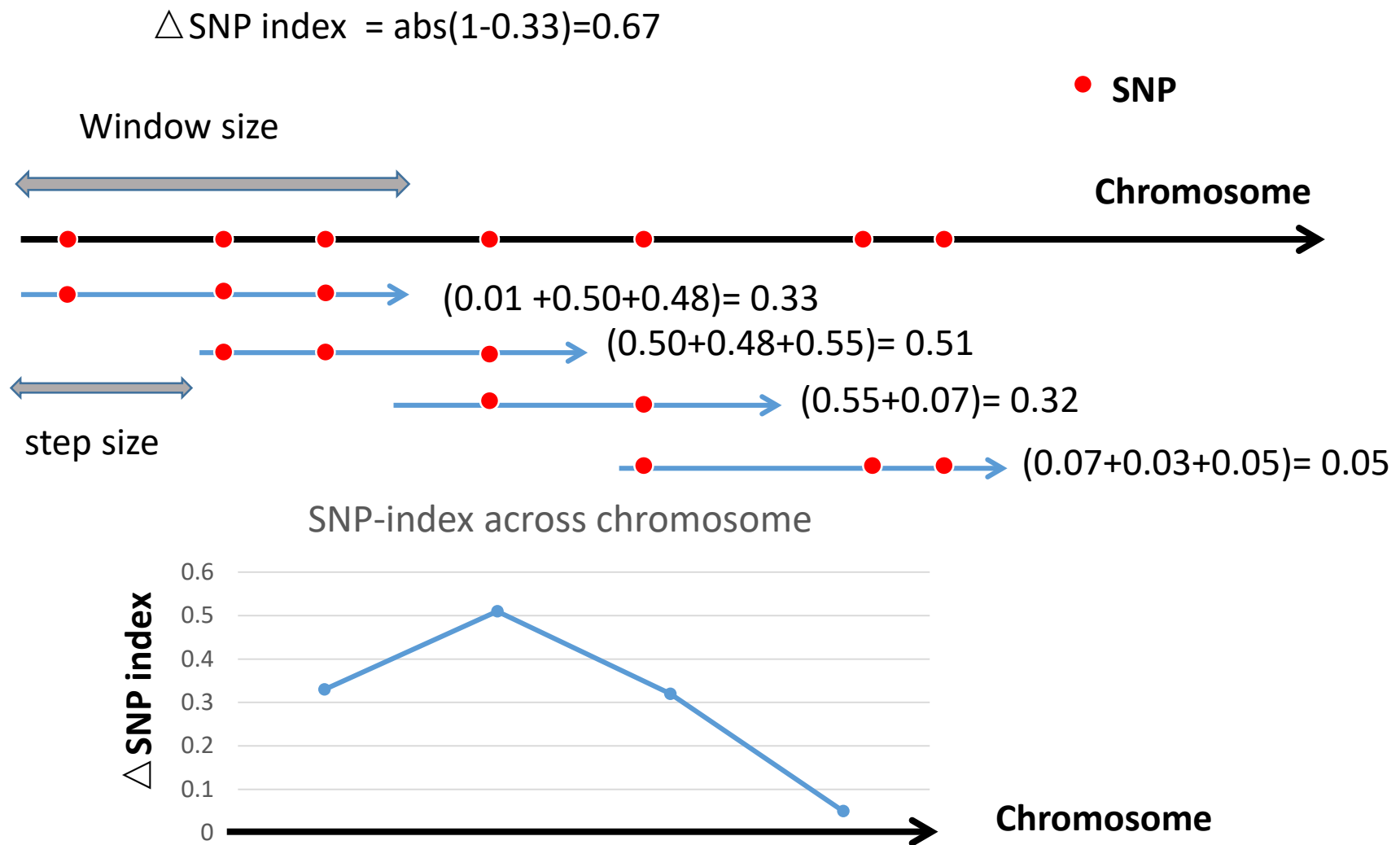
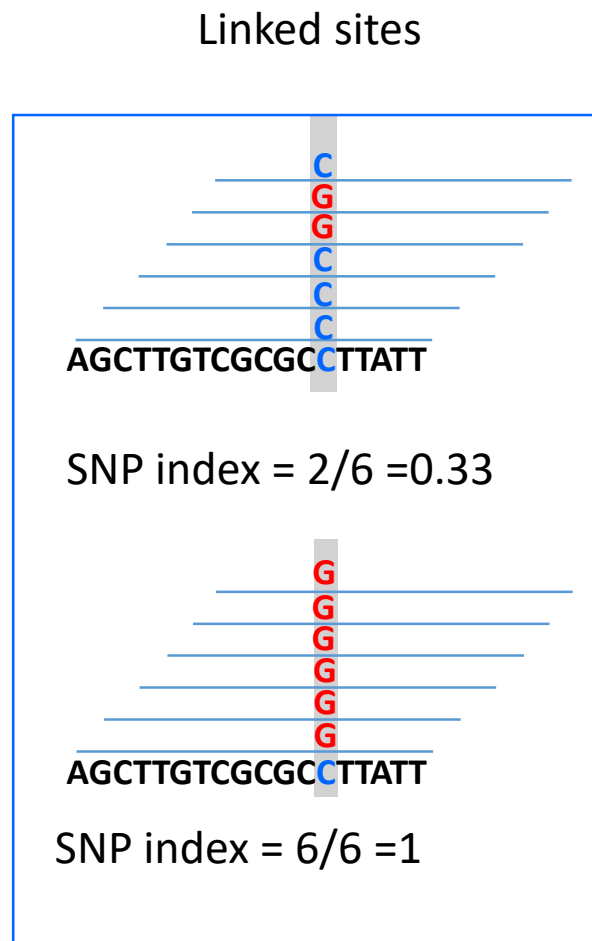
Using read depth directly, not the alleles that have been inferred.



# BSA Pipeline (part 1 variants calling)

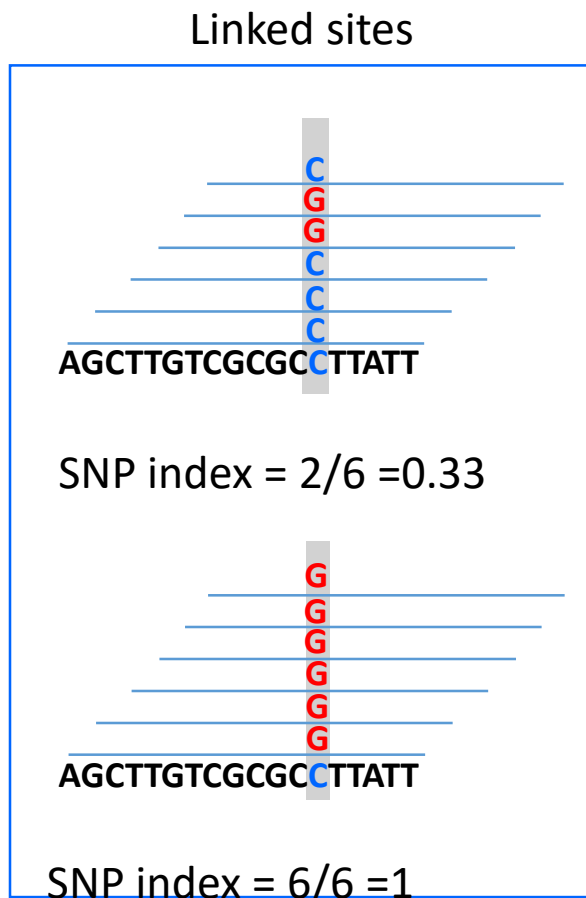


# BSA Pipeline (part 2 Statistics and sliding window)



## Method 2. fishier exact test

- 2. Compare fishier exact test to test if the read depth in each buck are significantly different or not.

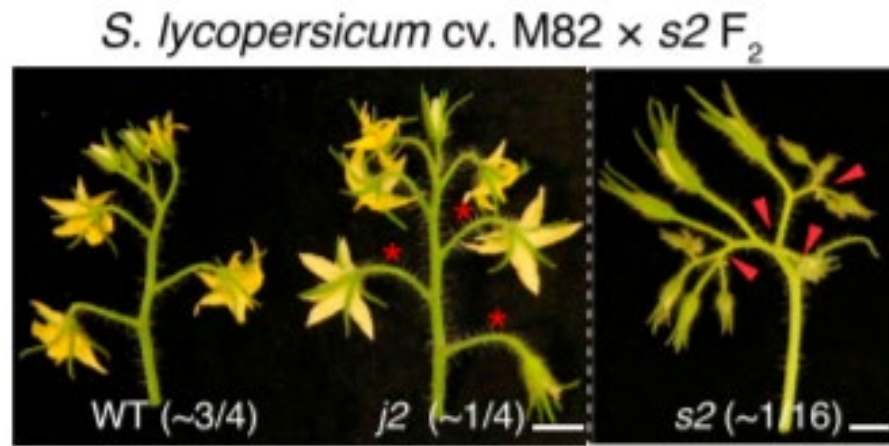


	Ref allele	Alt allele	Row total
WT	4	2	6
Mutant	0	6	6
Column total	4	8	12

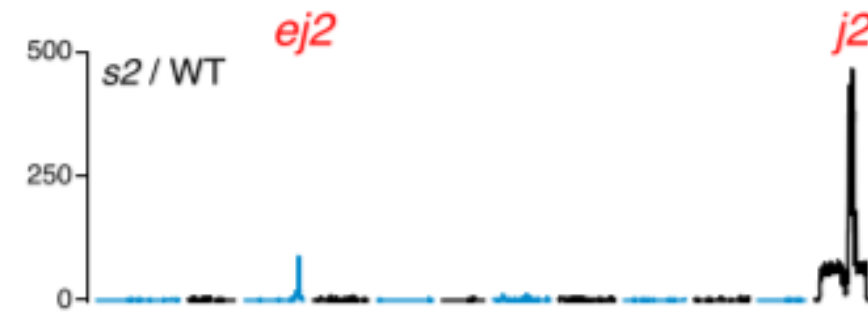
$$p = \frac{\binom{6}{4} \binom{6}{0}}{\binom{12}{4}}$$

```
F=fisher.test(rbind(c(4,2),c(0,6)),  
              alternative="two.sided")  
F$p.value  
0.06061
```

# An exercise of BSA



*S. lycopersicum* cv. M82 × s2 F<sub>2</sub>



# Download reads and reference genome

The Sequence Read Archive (SRA) on NCBI is the most commonly used website to store the high-throughput sequencing data.

- fastq-dump --split-files --gzip SRR5274882
- fastq-dump --split-files --gzip SRR5274880
- wget [ftp://ftp.ensemblgenomes.org/pub/plants/release-35/fasta/solanum\\_lycopersicum/dna/Solanum\\_lycopersicum.SL2.50.dna.toplevel.fa.gz](ftp://ftp.ensemblgenomes.org/pub/plants/release-35/fasta/solanum_lycopersicum/dna/Solanum_lycopersicum.SL2.50.dna.toplevel.fa.gz)

**Do not run. Data has been downloaded.**

To speed up the calculations, the data has been down-sampled using reads that were mapped to chr3 only in the test data. If you are interested in testing the entire data, you can download it from NCBI.

# Copy the data under your directory

```
cp -r /shared_data/BSA_workshop_2018/* ./
```

```
tree -A
```

```
[chengzou@cbsuvitisgen2 upload_test]$ tree -A
.
├── 00.src
│   ├── 01.variants_call.pl
│   ├── check_depth.R
│   ├── Difference_window.R
│   ├── Fisher_window.R
│   ├── plot_signal.R
│   └── Ratio_window.R
├── 01.reference
│   └── Solanum_lycopersicum.SL2.50.dna.toplevel.fa
└── 02.reads
    ├── mut_1.fq.gz
    ├── mut_2.fq.gz
    ├── wt_1.fq.gz
    └── wt_2.fq.gz
    └── command_lines.sh
    └── reads_table

3 directories, 13 files
```

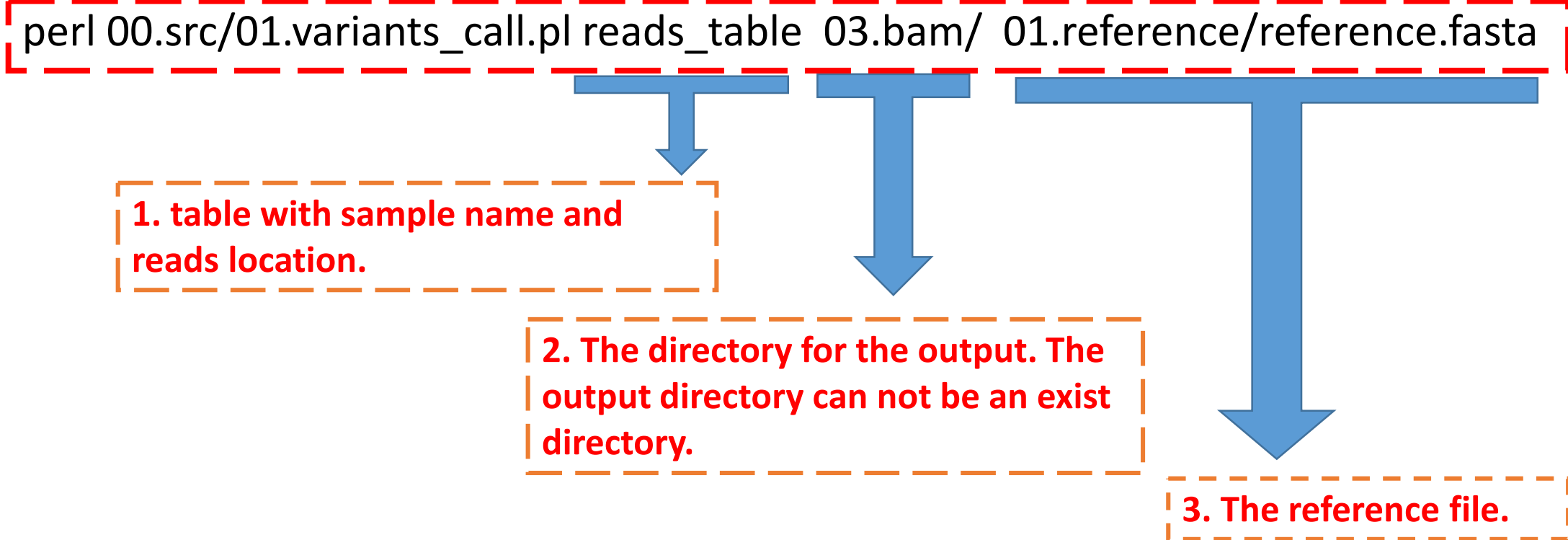
# Index the genome

```
cd 01.reference  
ln -s Solanum_lycopersicum.SL2.50.dna.toplevel.fa reference.fasta  
bwa index reference.fasta  
java -jar /programs/picard-tools-2.9.0/picard.jar  
CreateSequenceDictionary R=reference.fasta  
samtools faidx reference.fasta
```

# It takes about ten minutes to finish

```
[bwt_gen] Finished constructing BWT in 233 iterations.  
[bwa_index] 580.43 seconds elapse.  
[bwa_index] Update BWT... 4.67 sec  
[bwa_index] Pack forward-only FASTA... 4.33 sec  
[bwa_index] Construct SA from BWT and Occ... 253.99 sec  
[main] Version: 0.7.13-r1126  
[main] CMD: bwa index reference.fasta  
[main] Real time: 850.328 sec; CPU: 850.037 sec
```

# Variance calling



## **Reads\_table is a tab delimited txt file**

```
[chengzou@cbsuvititisgen2 upload]$ head reads_table
mut      02.reads/mut_1.fq.gz      02.reads/mut_2.fq.gz
wt       02.reads/wt_1.fq.gz      02.reads/wt_2.fq.gz
```

# Step 1: Align the reads, sort and index the results

```
bwa mem -t 8 -M -R '@RG\tID:mut\tSM:mut' 01.reference/reference.fasta  
03.bam /fixed6.mut_1.fq.gz 04.bam/fixed6.mut_2.fq.gz | samtools sort -@ 8 -o  
03.bam /mut.sorted.bam - 2>> 03.bam/bwalog  
java -jar /programs/picard-tools-2.9.0/picard.jar BuildBamIndex INPUT= 03.bam  
/mut.sorted.redup.bam QUIET=true VERBOSITY=ERROR
```

```
bwa mem -t 8 -M -R '@RG\tID:wt\tSM:wt' 01.reference/reference.fasta 03.bam  
/ fixed.wt_1.fq.gz 04.bam/fixed.wt_2.fq.gz | samtools sort -@ 8 -o  
03.bam/wt.sorted.bam - 2>> 03.bam//bwalog  
java -jar /programs/picard-tools-2.9.0/picard.jar BuildBamIndex  
INPUT=04.10bam//wt.sorted.redup.bam QUIET=true VERBOSITY=ERROR
```

-M : mark shorter split hits as secondary (*for Picard compatibility*).

## Step 2: Filtering the alignments, mpileup and variance calling

```
 samtools mpileup -t AD,DP \
 | -C 50 \
 | -Q 20 \
 | -q 40 \
 | -f 01.reference/reference.fasta \
 | 03.bam/mut.sorted.redup.bam \
 | 03.bam/wt.sorted.redup.bam \
 | -v \
 | bcftools call --consensus-caller --variants-
 | only --pval-threshold 1.0 -O z -o Out.vcf.gz
```

- t LIST optional tags to output  
DP,AD,ADF,ADR,SP,INFO/AD,INFO/AD  
F,INFO/ADR
  - C adjust mapping quality;  
recommended:50 (unique hit of the reads)
  - Q skip bases with baseQ/BAQ smaller than INT [13]
  - q skip alignments with mapQ smaller than INT [0]
  - f faidx indexed reference sequence file
- } input bam files
- v generate genotype likelihoods in VCF format

# vcf file of variance calling result

```
##bcftools_viewVersion=1.8+htslib-1.8
##bcftools_viewCommand=view -m2 -M2 -O z -o 03.bam/filter.vcf.gz -; Date=Tue Nov 27
14:00:32 2018
#CHROM POS ID REF ALT QUAL FILTER INFO FORMAT mut wt
3 357 . A C 4.34172 PASS DP=11;VDB=0.1;SGB=0.0047313
;RPB=0.5;MQB=0.222222;BQB=0.777778;MQ0F=0;AF1=0.271323;AC1=1;DP4=9,0,2,0;MQ=46;FQ=
5.28671;PV4=1,0.320328,0.0449975,1 GT:PL:DP:AD 0/1:35,0,119:8:6,2 0
/0:0,9,76:3:3,0
3 539 . A C 3.81791 PASS DP=13;VDB=0.84;SGB=-2.48712
;RPB=0.5;MQB=0.5;MQSB=0.838008;BQB=0.5;MQ0F=0;AF1=0.495023;AC1=2;DP4=6,4,1,1;MQ=50;
FQ=5.75671;PV4=1,1,0.00809854,1 GT:PL:DP:AD 0/1:15,0,147:7:6,1 0/1:21,0,
74:5:4,1
3 762 . C T 9.96297 PASS DP=11;VDB=0.72;SGB=-2.48712
;RPB=0.666667;MQB=1;MQSB=0.450401;BQB=0.666667;MQ0F=0;AF1=0.495209;AC1=2;DP4=3,3,2,
0;MQ=43;FQ=12.6728;PV4=0.464286,0.209877,0.284691,1 GT:PL:DP:AD 0/1:14,0,
140:6:5,1 0/1:30,0,26:2:1,1
```

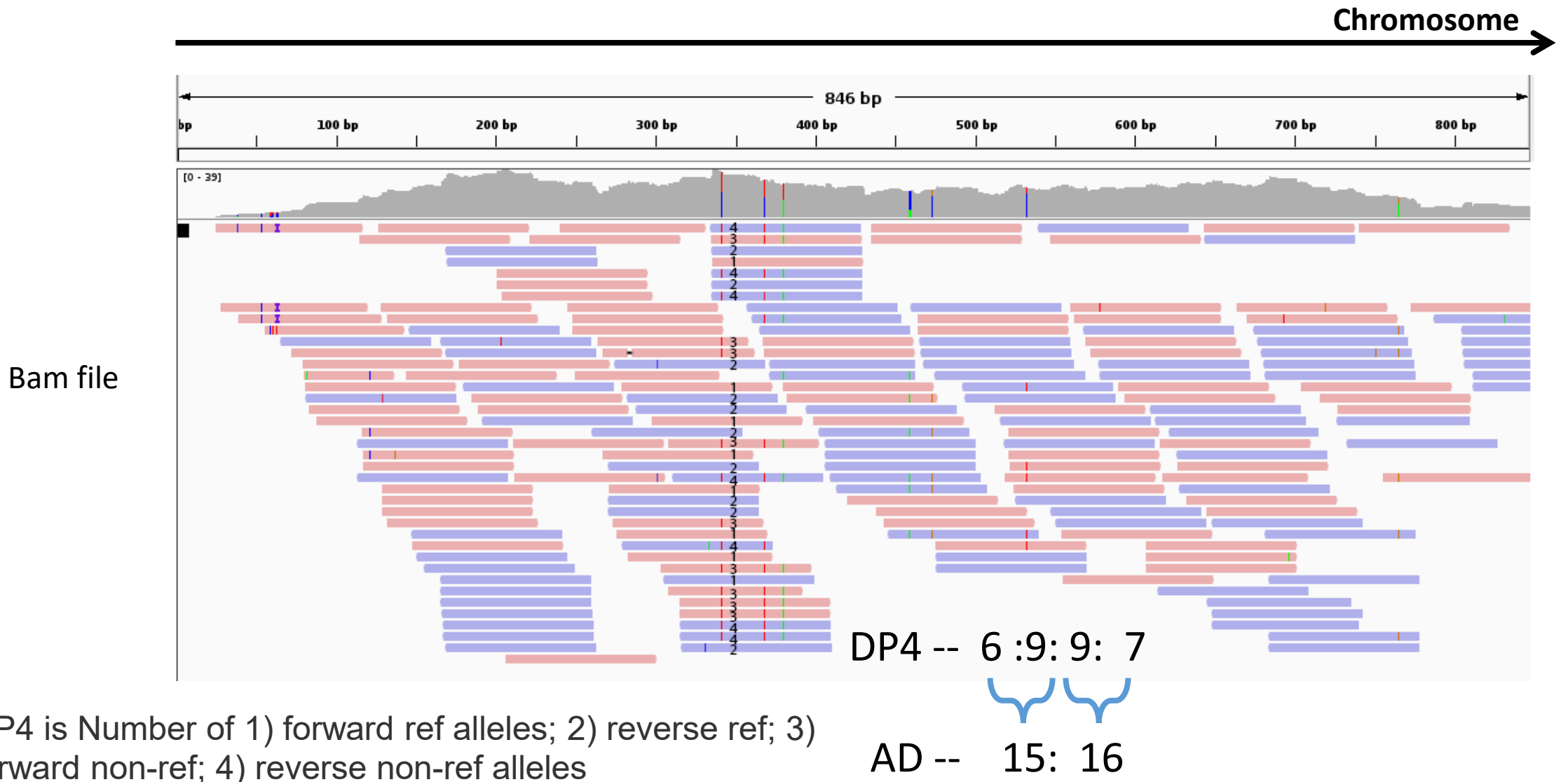
GT: Genotype

PL: list of Phred-scaled genotype likelihoods

DP: Number of high-quality bases

AD: Allelic depths

# Definition of DP4 and AD



## Step 3: Filtering the variances

```
bcftools filter \  
-g10 \  
-G10 \  
-i '(DP4[0]+DP4[1])>1 & (DP4[2]+DP4[3])>1  
& FORMAT/DP[]>5' Out.vcf.gz \  
| bcftools view \  
-m2 -M2  
-  
-O z  
-o 03.bam/filter.vcf.gz
```

- g filter SNPs within <int> base pairs of an indel
- G filter clusters of indels separated by <int> or fewer base pairs allowing only one to pass
- i expression of Variance that will be included:  
 $(DP4[0]+DP4[1])>1 \& (DP4[2]+DP4[3])>1$   
Both reference allele and alternative allele must be support by at least 2 reads.  
FORMAT/DP[]>5 for each sample, there must be more than five reads covering this site.
- m2 -M2 to only view biallelic SNPs
- O format of the output file
- o name of the output file

## Step 4: Extract information for downstream analysis

```
bcftools query \
-i 'TYPE="SNP"' \
-f '%CHROM\t%POS\t%REF\t%ALT{0}\t%DP[\t%AD]\n' \
03.bam/filter.vcf.gz | sed 's/[,]/\t/g' - 
>03.bam/filter.vcf.txt
```

## Final result in vcf format-- filter.vcf.gz

```
##bcftools_viewVersion=1.8+htslib-1.8
##bcftools_viewCommand=view -m2 -M2 -O z -o 03.bam/filter.vcf.gz -; Date=Tue Nov 27
14:00:32 2018
#CHROM POS ID REF ALT QUAL FILTER INFO FORMAT mut wt
3 357 . A C 4.34172 PASS DP=11;VDB=0.1;SGB=0.0047313
6;RPB=0.5;MQB=0.222222;BQB=0.777778;MQ0F=0;AF1=0.271323;AC1=1;DP4=9,0,2,0;MQ=46;FQ=
5.28671;PV4=1,0.320328,0.0449975,1 GT:PL:DP:AD 0/1:35,0,119:8:6,2 0
/0:0,9,76:3:3,0
3 539 . A C 3.81791 PASS DP=13;VDB=0.84;SGB=-2.48712
;RPB=0.5;MQB=0.5;MQSB=0.838008;BQB=0.5;MQ0F=0;AF1=0.495023;AC1=2;DP4=6,4,1,1;MQ=50;
FQ=5.75671;PV4=1,1,0.00809854,1 GT:PL:DP:AD 0/1:15,0,147:7:6,1 0/1:21,0,
74:5:4,1
3 762 . C T 9.96297 PASS DP=11;VDB=0.72;SGB=-2.48712
;RPB=0.666667;MQB=1;MQSB=0.450401;BQB=0.666667;MQ0F=0;AF1=0.495209;AC1=2;DP4=3,3,2,
0;MQ=43;FQ=12.6728;PV4=0.464286,0.209877,0.284691,1 GT:PL:DP:AD 0/1:14,0,
140:6:5,1 0/1:30,0,26:2:1,1
```

# Final result in txt format -- filter.vcf.txt

```
[chengzou@cbsuvitisgen2 04.10bam]$ less filter.vcf.txt
```

3	357	A	C	11	6	2	3	0
3	539	A	C	13	6	1	4	1
3	762	C	T	11	5	1	1	1
3	860	C	T	35	15	1	12	3
3	906	G	T	41	19	1	15	3
3	949	T	A	42	22	1	13	1
3	1369	A	C	25	11	1	5	1
3	1449	A	C	29	11	1	5	1
3	1454	C	A	30	15	1	11	1
3	1485	T	G	28	7	2	7	0
3	1488	T	G	27	9	1	8	2
3	1524	T	C	27	8	1	7	2

Chr	Pos	Ref	Alt	total DP	Mut_ref	Mut_alt	WT_ref	WT_alt
-----	-----	-----	-----	----------	---------	---------	--------	--------

# The running log

```
[chengzou@cbsuvitisgen2 23.BSA_test]$ perl 00.src/01.variants_call.pl reads_table 04.bam/ 01.reference/reference.fasta
[M::bwa_idx_load_from_disk] read 0 ALT contigs
[M::process] read 537238 sequences (80000058 bp)...
[M::process] read 537556 sequences (80000264 bp)...
[M::mem_pestat] # candidate unique pairs for (FF, FR, RF, RR): (15, 172278, 39, 8)
[M::mem_pestat] analyzing insert size distribution for orientation FF...
[M::mem_pestat] (25, 50, 75) percentile: (493, 624, 1867)
[M::mem_pestat] low and high boundaries for computing mean and std.dev: (1, 4615)
[M::mem_pestat] mean and std.dev: (966.93, 857.13)
[M::mem_pestat] low and high boundaries for proper pairs: (1, 5989)
[M::mem_pestat] analyzing insert size distribution for orientation FR...
```

```
INFO    2018-11-19 13:10:40    MarkDuplicates After output close freeMemory: 13338438088; totalMemory: 13466861568; maxMemory: 19088801792
[Mon Nov 19 13:10:40 EST 2018] picard.sam.markduplicates.MarkDuplicates done. Elapsed time: 4.42 minutes.
Runtime.totalMemory()=13466861568
[mpileup] 2 samples in 2 input files
Note: none of --samples-file, --ploidy or --ploidy-file given, assuming all sites are diploid
<mpileup> Set max per-file depth to 4000
```

# Result of the run

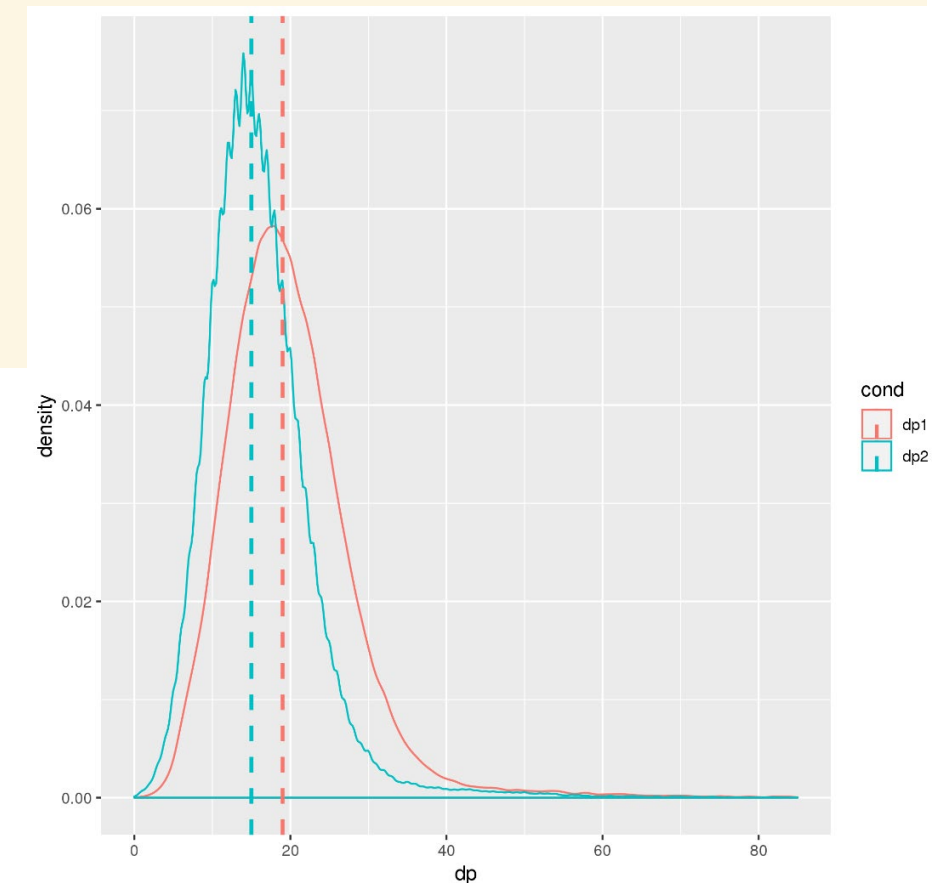
```
[chengzou@cbsuvitisgen2 03.bam]$ ls -l
total 2031636
-rw-rw-r-- 1 chengzou chengzou      1123 Nov 27 14:00 bwalog
-rw-rw-r-- 1 chengzou chengzou    10181013 Nov 27 14:00 filter.vcf.gz
-rw-rw-r-- 1 chengzou chengzou    4218553 Nov 27 14:06 filter.vcf.txt
-rw-rw-r-- 1 chengzou chengzou    955320 Nov 27 13:05 mut.sorted.bai
-rw-rw-r-- 1 chengzou chengzou   1035205166 Nov 27 13:04 mut.sorted.bam
-rw-rw-r-- 1 chengzou chengzou   137945951 Nov 27 14:00 Out.vcf.gz
-rw-rw-r-- 1 chengzou chengzou    918520 Nov 27 13:17 wt.sorted.bai
-rw-rw-r-- 1 chengzou chengzou   890951273 Nov 27 13:17 wt.sorted.bam
```

# Check distribution of the depth in each pool

```
R --vanilla --slave --args filter.vcf.txt < ../00.src/check_depth.R
```

```
[chengzou@cbsuvitisgen2 04.10bam]$ R --vanilla --slave --args filter.vcf.txt < ../00.src/check_depth.R
  Min. 1st Qu. Median Mean 3rd Qu. Max.
0.00 14.00 19.00 20.46 24.00 3051.00
  Min. 1st Qu. Median Mean 3rd Qu. Max.
0.00 12.00 15.00 16.69 19.00 3264.00
cond dp.median
1  dp1      19
2  dp2      15
Warning message:
Removed 487 rows containing non-finite values (stat_density).
```

**SNP with total read depth that is larger than two times of the average is not desired.**



# Further filtering by depth distribution

## Examples:

```
MIN(DV)>5
MIN(DV/DP)>0.3
MIN(DP)>10 & MIN(DV)>3
FMT/DP>10 & FMT/GQ>10 .. both conditions must be satisfied within one sample
FMT/DP>10 && FMT/GQ>10 .. the conditions can be satisfied in different samples
QUAL>10 | FMT/GQ>10 .. true for sites with QUAL>10 or a sample with GQ>10, but selects only samples with GQ>10
QUAL>10 || FMT/GQ>10 .. true for sites with QUAL>10 or a sample with GQ>10, plus selects all samples at such sites
TYPE="snp" && QUAL>=10 && (DP4[2]+DP4[3] > 2)
COUNT(GT="hom")=0
MIN(DP)>35 && AVG(GQ)>50
ID=@file .. selects lines with ID present in the file
ID!=@~/file .. skip lines with ID present in the ~/file
MAF[0]<0.05 .. select rare variants at 5% cutoff
POS>=100 .. restrict your range query, e.g. 20:100-200 to strictly sites with POS in that range.
```

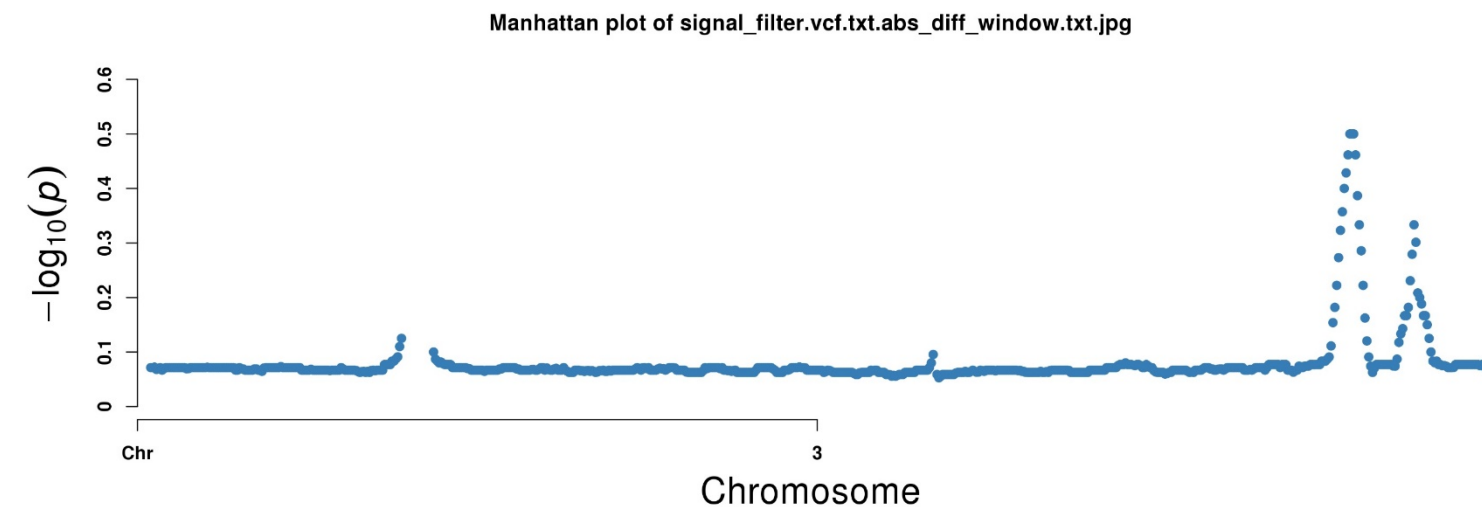
```
bcftools filter -i 'FORMAT/DP[1]<30 & FORMAT/DP[2]<34' filter.vcf.gz
-O z -o filter2.vcf.gz
```

# Summary statistics 1. $\Delta$ SNP index



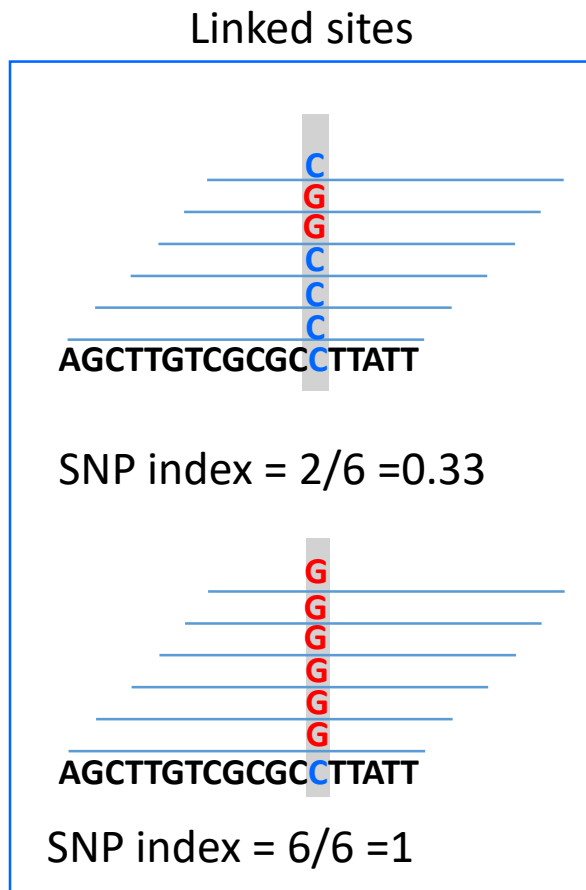
```
R --vanilla --slave --args filter.vcf.txt < ../00/src/Difference_window.R  
R --vanilla --slave --args filter.vcf.txt.abs_diff_window.txt  
< ../00/src/plot_signal.R
```

$$\Delta \text{SNP index} = \text{abs}(1-0.33)=0.67$$



## Summary statistics 2. ratio of allele frequency

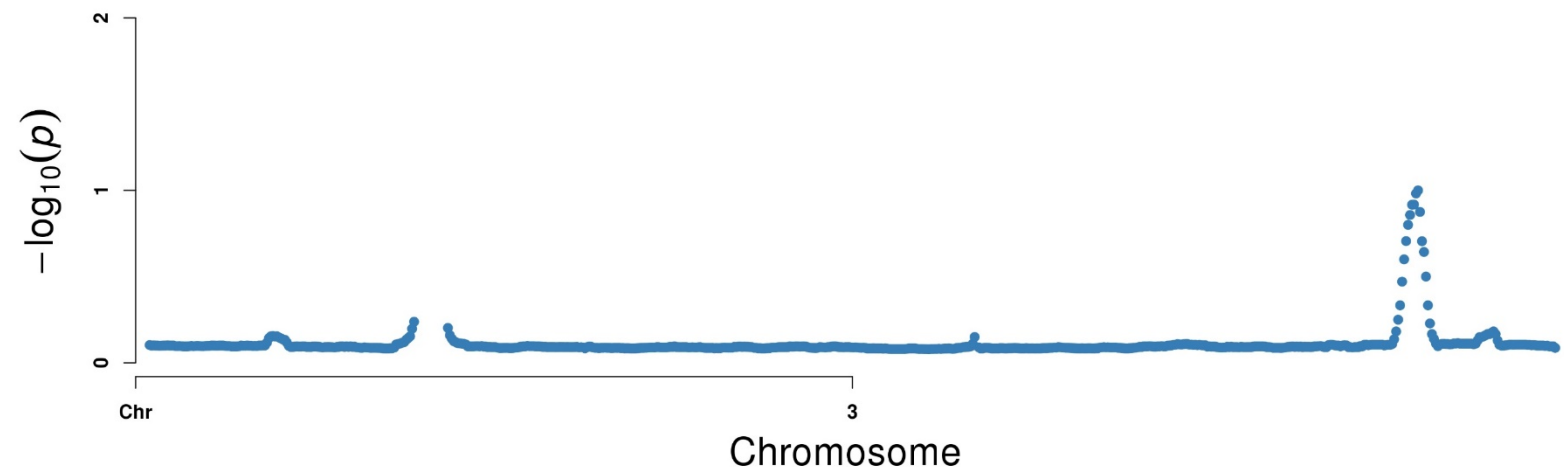
- 1. Compare the ratio, and sliding window to find the peaks.



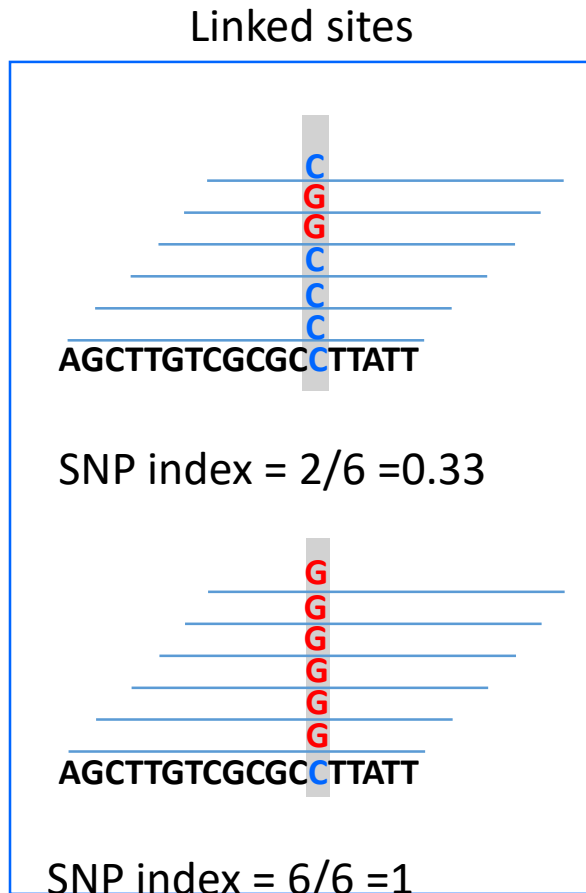
```
R --vanilla --slave --args filter.vcf.txt < ../00/src/Ratio_window.R  
R --vanilla --slave --args filter.vcf.txt.ratio_window.txt  
< ../00/src/plot_signal.R
```

$$\text{Ratio of SNP index} = \frac{1}{0.33} = 3$$

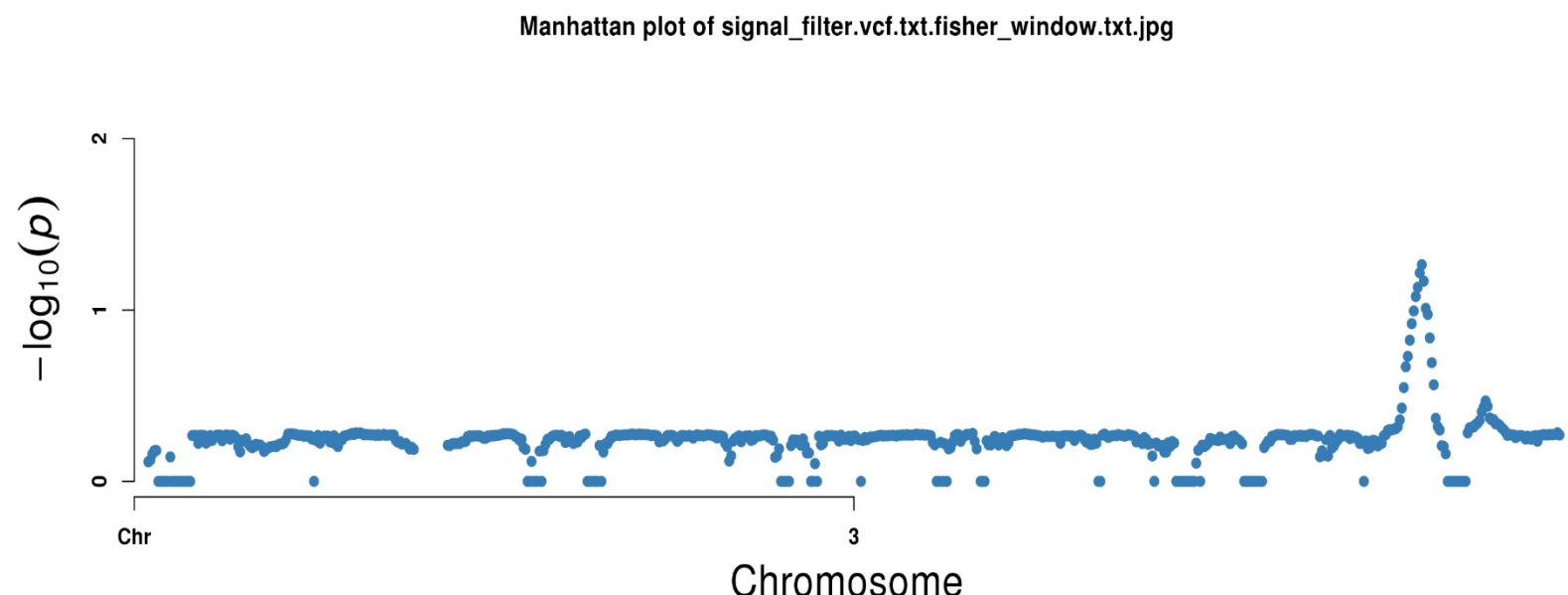
Manhattan plot of signal\_filter.vcf.txt.ratio\_window.txt.jpg



# Summary statistics 3. fishier exact test



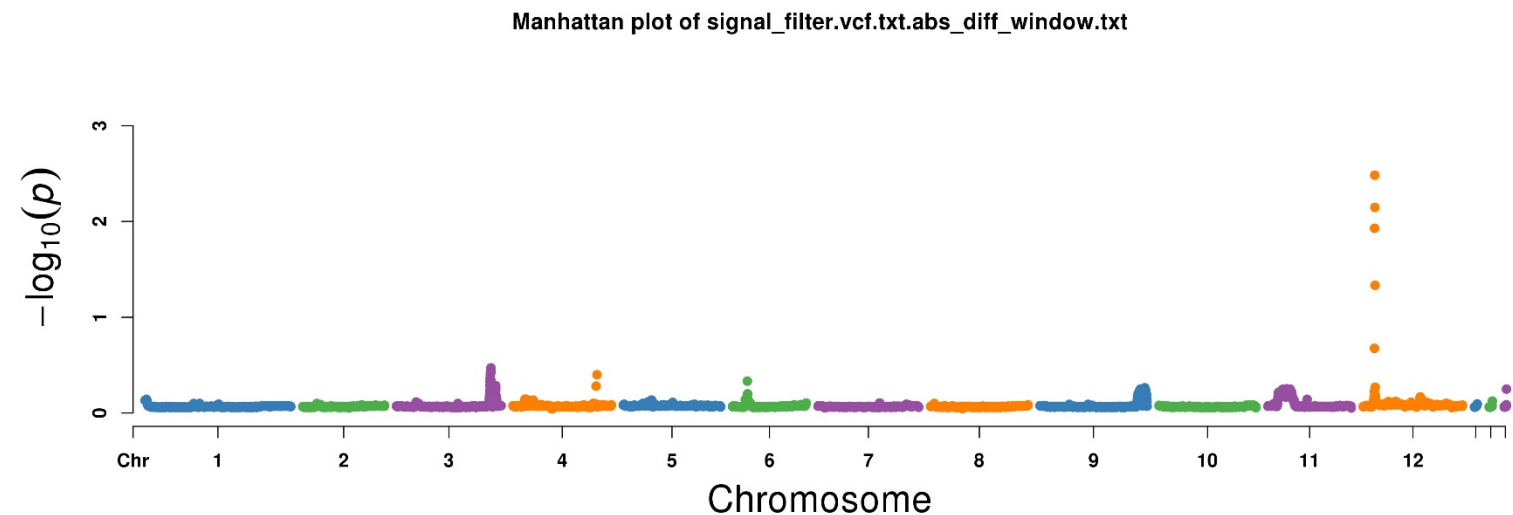
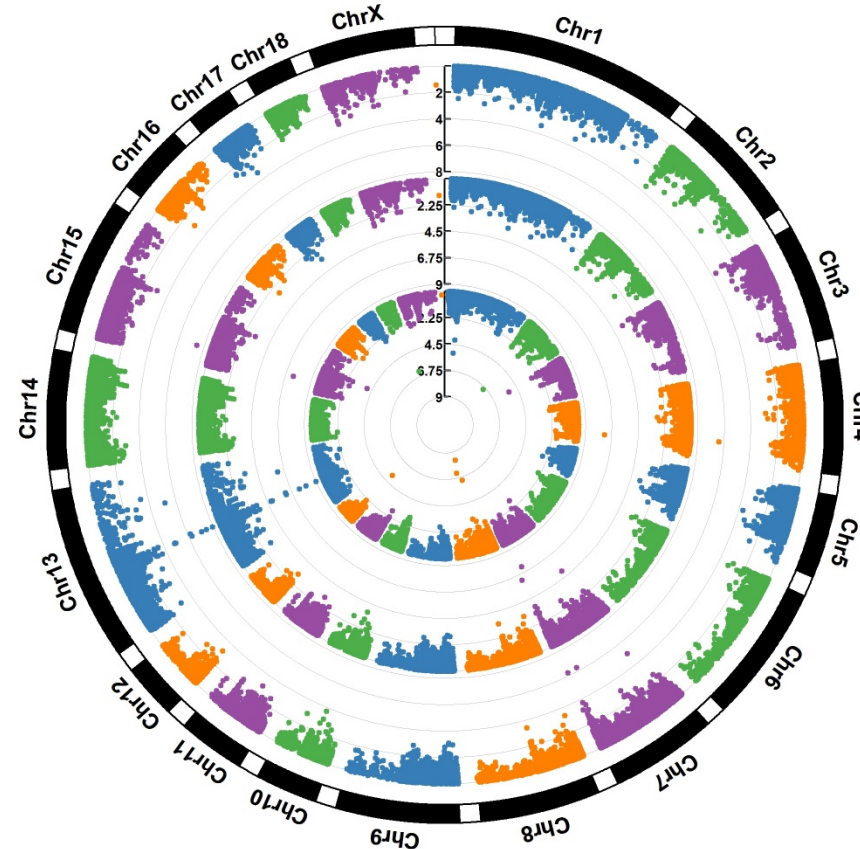
```
R --vanilla --slave --args filter.vcf.txt < ../00/src/Fisher_window.R  
R --vanilla --slave --args filter.vcf.txt.fisher_window.txt  
< ../00/src/plot_signal.R
```



## A few notes of the R script:

- Window size in the script is 1 Mbp, steps is 100 kpb
- Only considering contigs >1 Mbp
- Chr name can be any characters, with or without “chr”
- You can manually modify the result ( `filter.vcf.txt.abs_diff_window.txt` ) to get rid of undesired scaffolds or contigs.

# More plotting options



- <https://github.com/YinLiLin/R-CMplot>

## Further reading

MutMap (Abe, A. et al., 2012)

QTL-seq (Takagi, H. et al., 2013)

MutMap+ (R Fekih et al., 2013)

MutMap-Gap (Takagi, H. et al., 2013)

BSR-Seq (Sanzhen, Liu et al., 2013 )