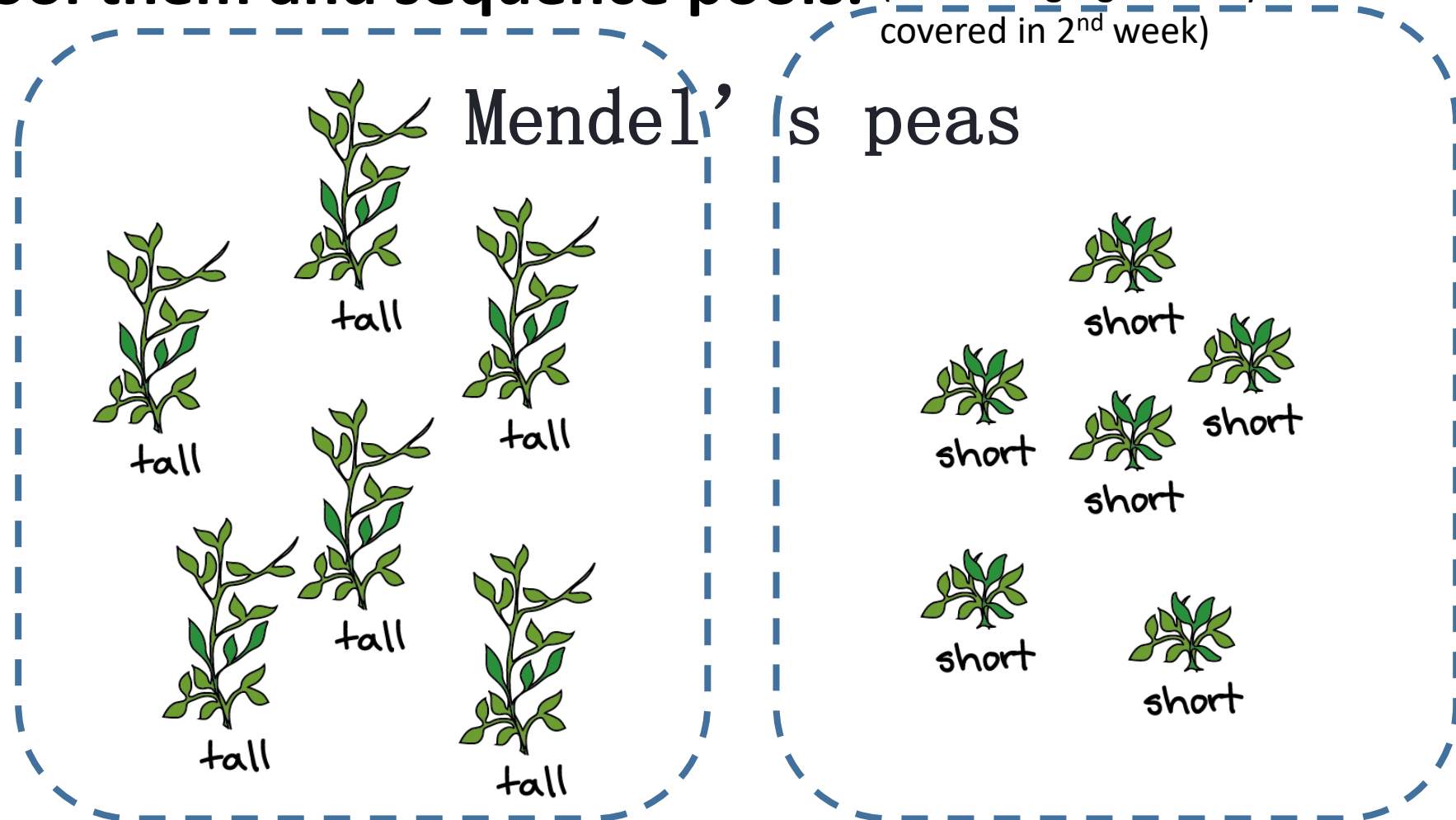


Bulked Segregant Analysis For Fine Mapping Of Genes

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

Pool them and sequence pools. (Bulked segregant analysis covered in 2nd week)

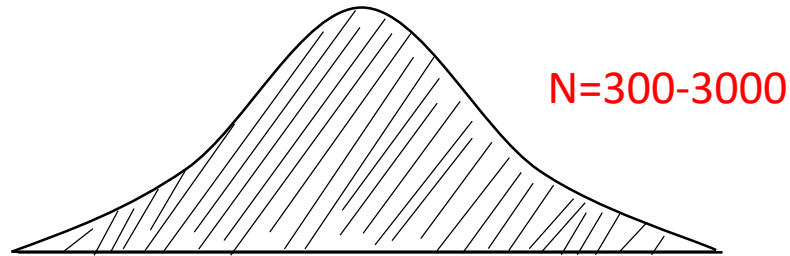


Outline

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

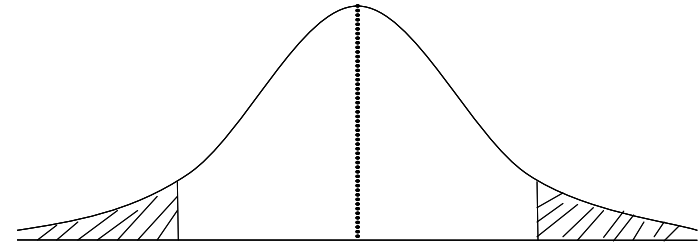
Compare BSA with traditional mapping strategy

Entire population
(all individual)
analysis

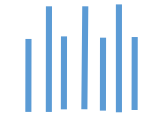


GWAS or linkage mapping

Phenotyping	entire population
Genotyping	entire population



Pool the samples
with extreme
phenotype

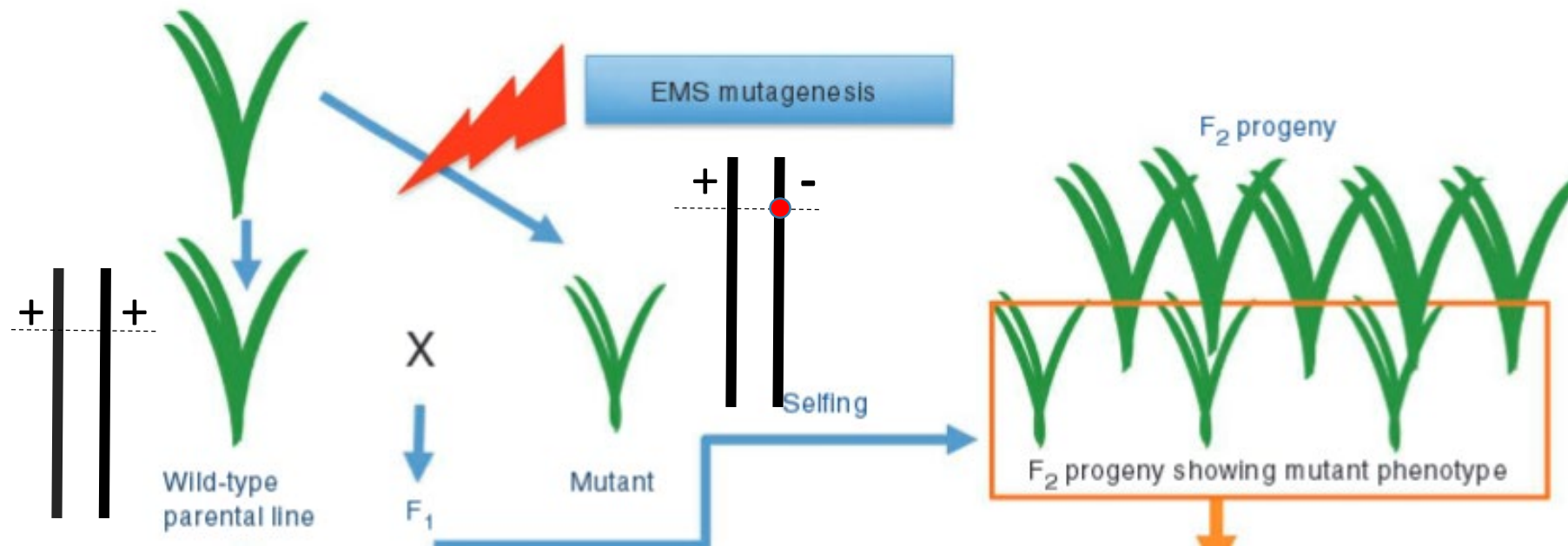


Phenotyping	entire population
Genotyping	two samples

Bulked Segregant Analysis (BSA)

rapid discovery of genetic markers and trait mapping

1. Segregation in phenotype



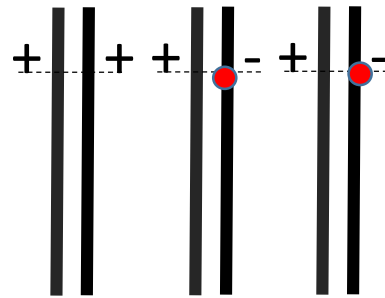
Bulked Segregant Analysis (BSA)

rapid discovery of genetic markers and trait mapping

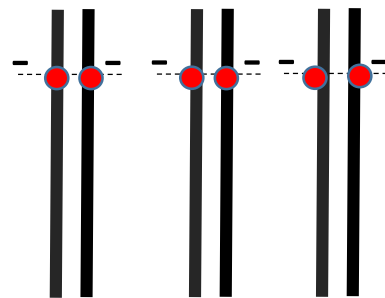
2. following Mendelian genetics or determined by a major effect loci



wide type



mutant



Linked sites

AGCTTGTCGCGCCTTATT

SNP index = $2/6 = 0.33$

AGCTTGTCGCGCCTTATT

SNP index = $6/6 = 1$

unlinked sites

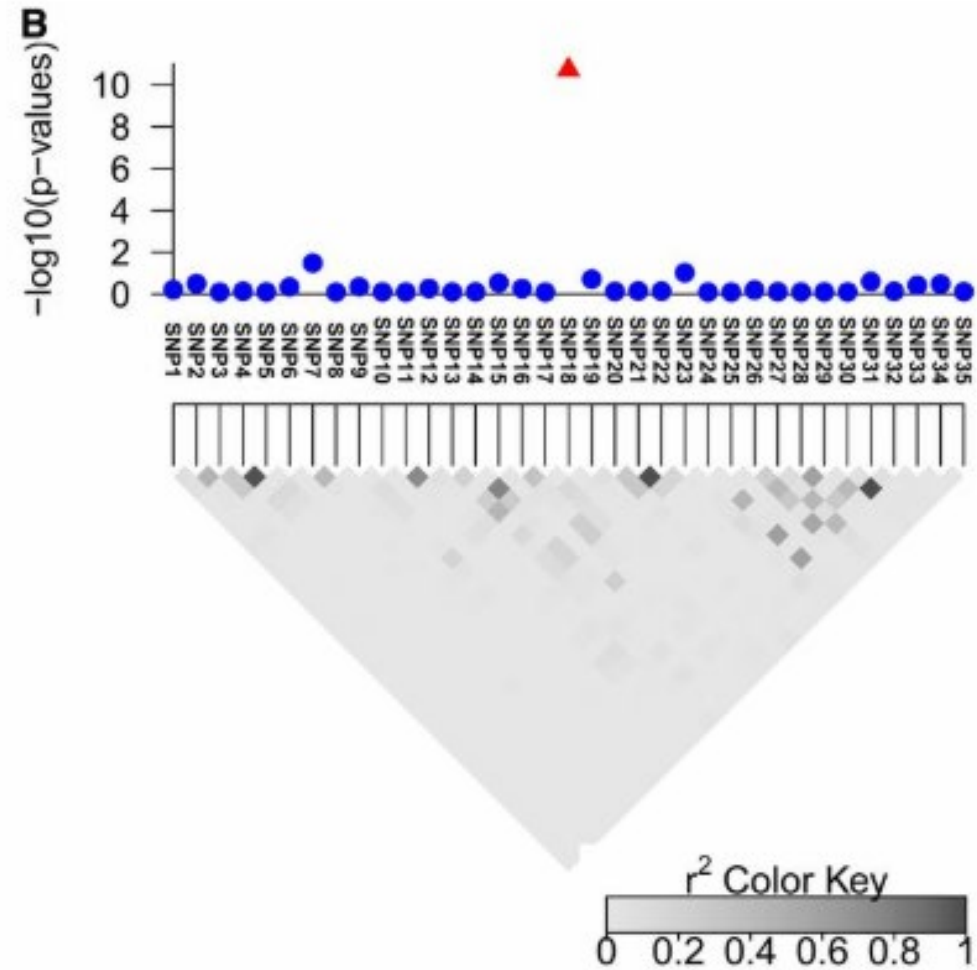
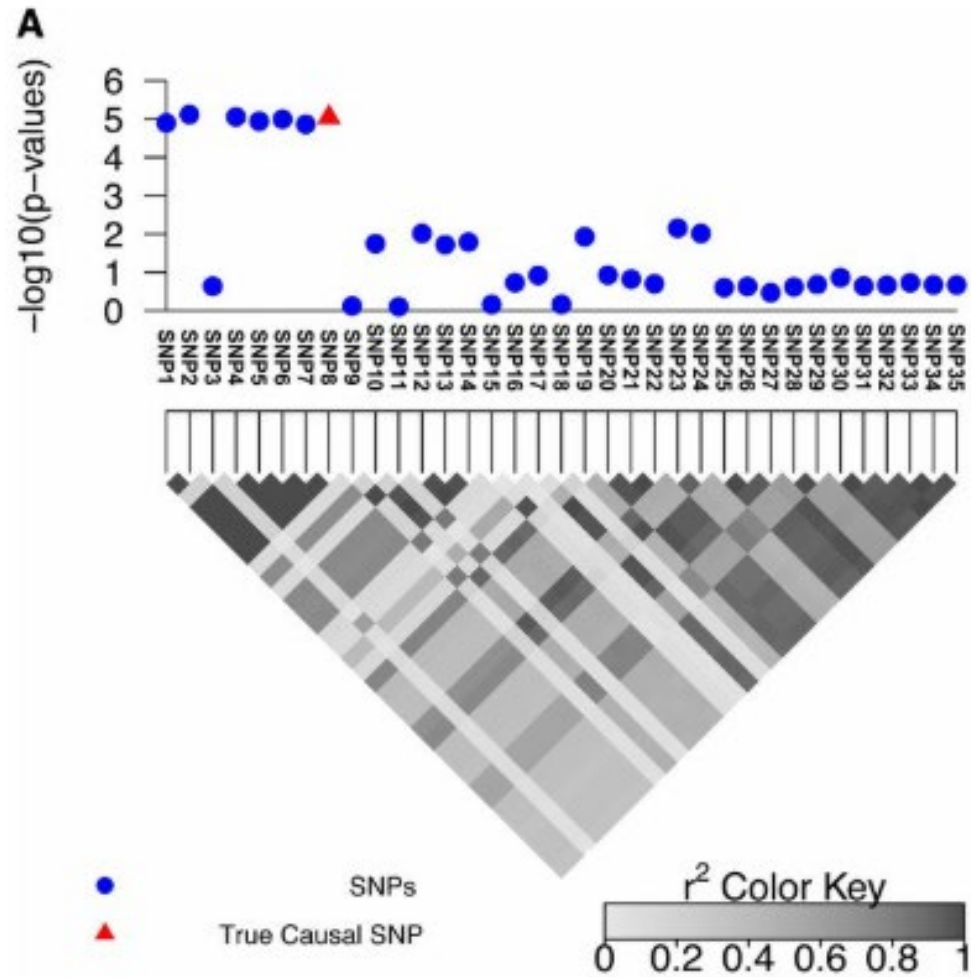
CAGGTATCGCGCTGGTT

SNP index = $2/6 = 0.33$

CAGGTATCGCGCTGGTT

SNP index = $3/6 = 0.5$

Causal SNP and SNPs linked with causal SNP

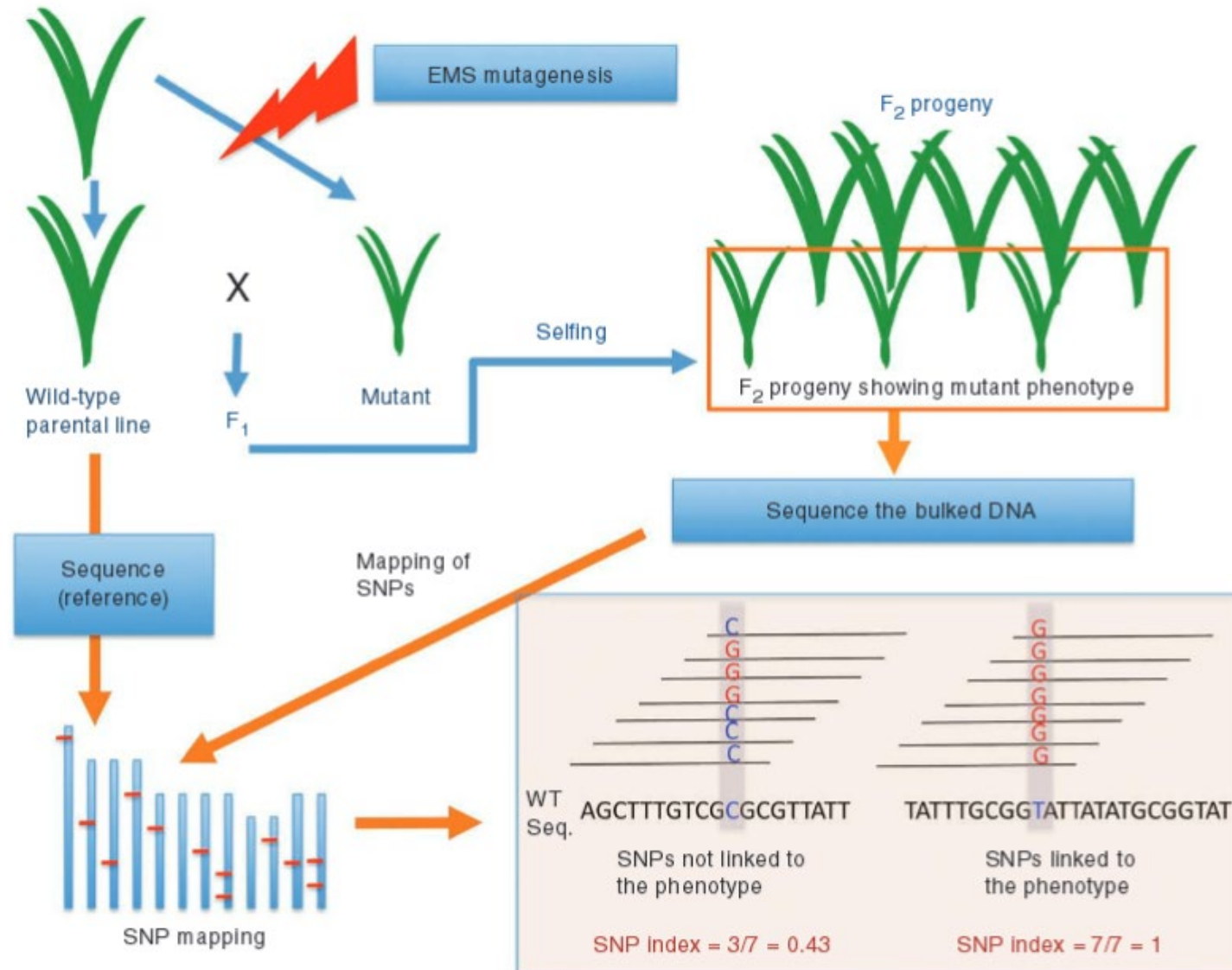


(copy from Hormozdiari, Farhad, et al *Genetics*, 2014)

Applicable populations

- EMS mutagenized population
- Mapping Population
- Natural Population

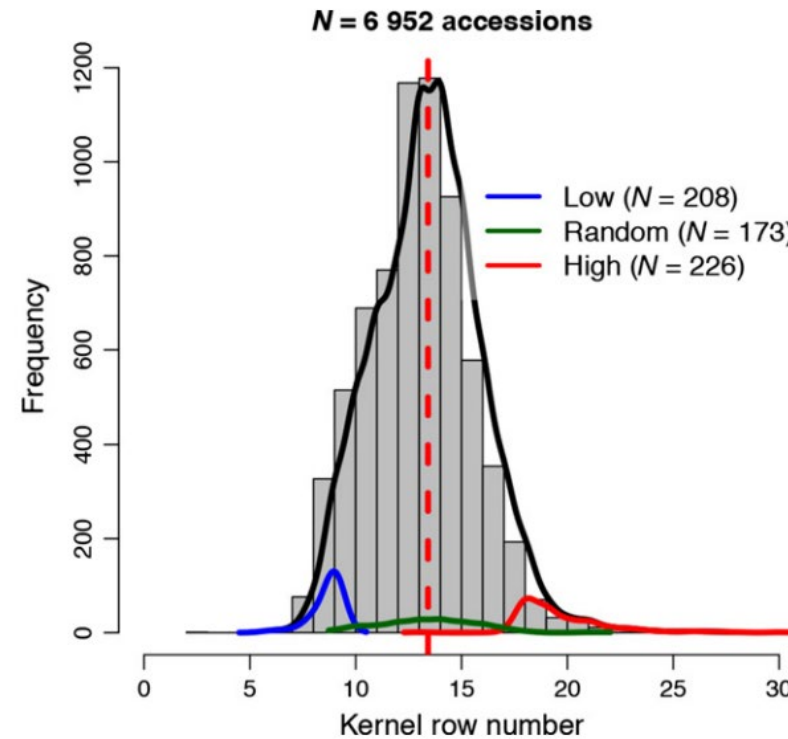
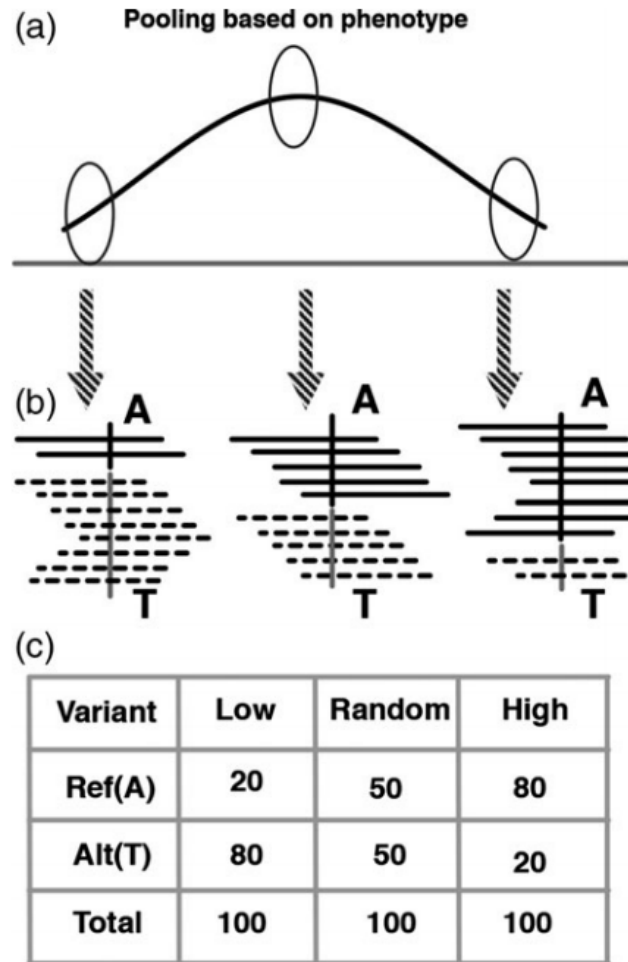
EMS mutagenized population



(Abe, 2011, NBT)

Extreme-phenotype GWAS using pooled samples

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



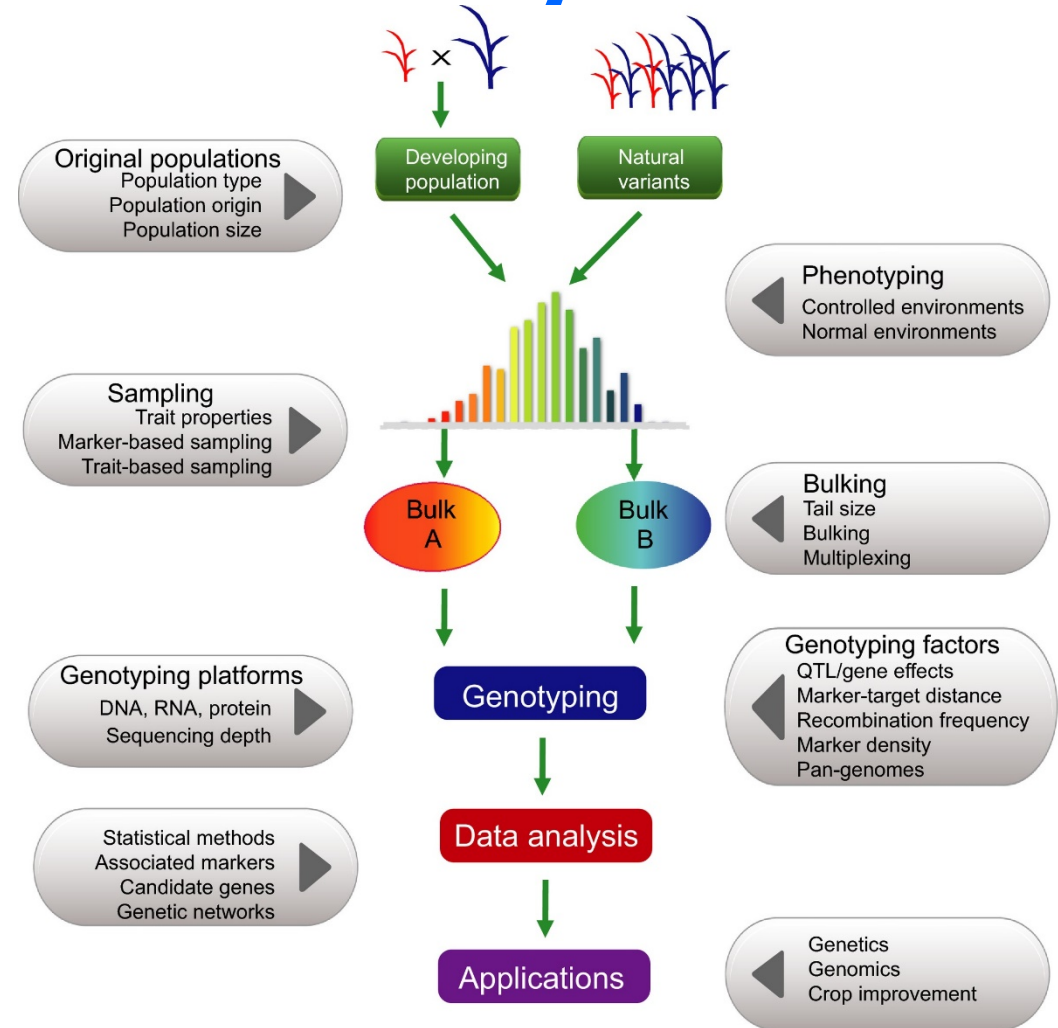
(Schnable, 2015 the Plant Journal)

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)

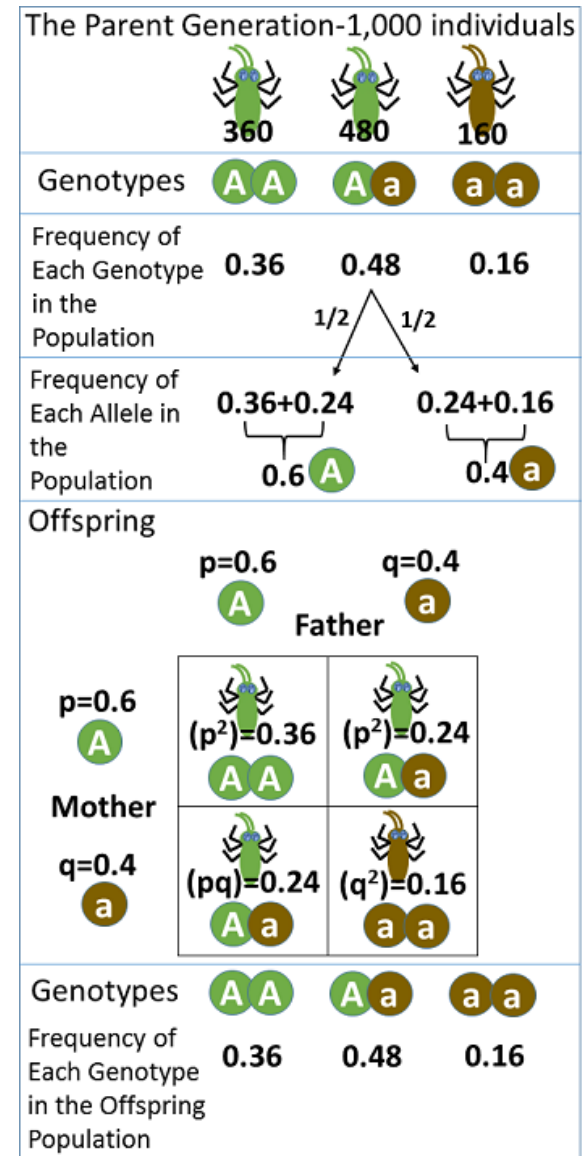
Beware of Variance Callings

Assumptions in Variant callers

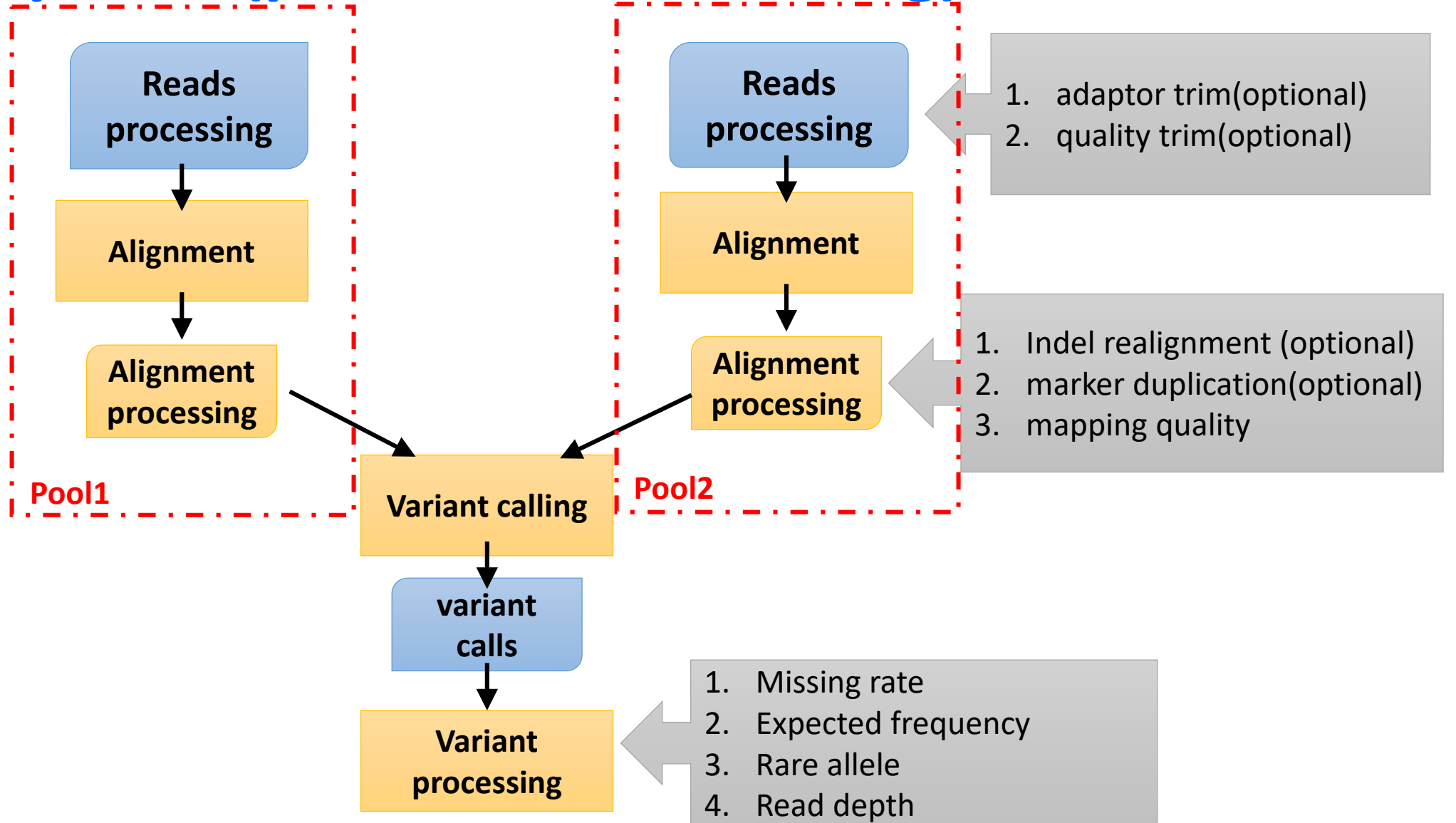
for example GATK :

- assuming Hardy-Weinberg equilibrium
- diploidy

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

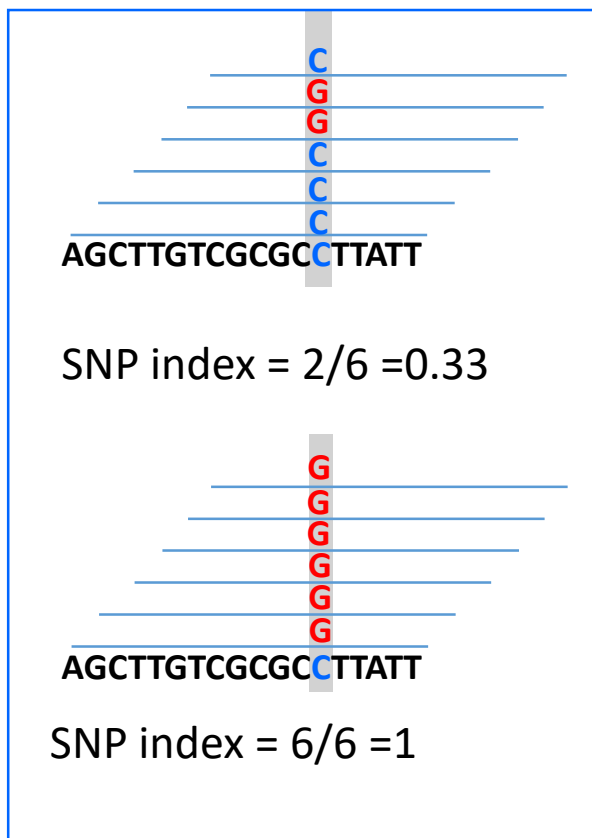


BSA Pipeline (part 1 variants calling)

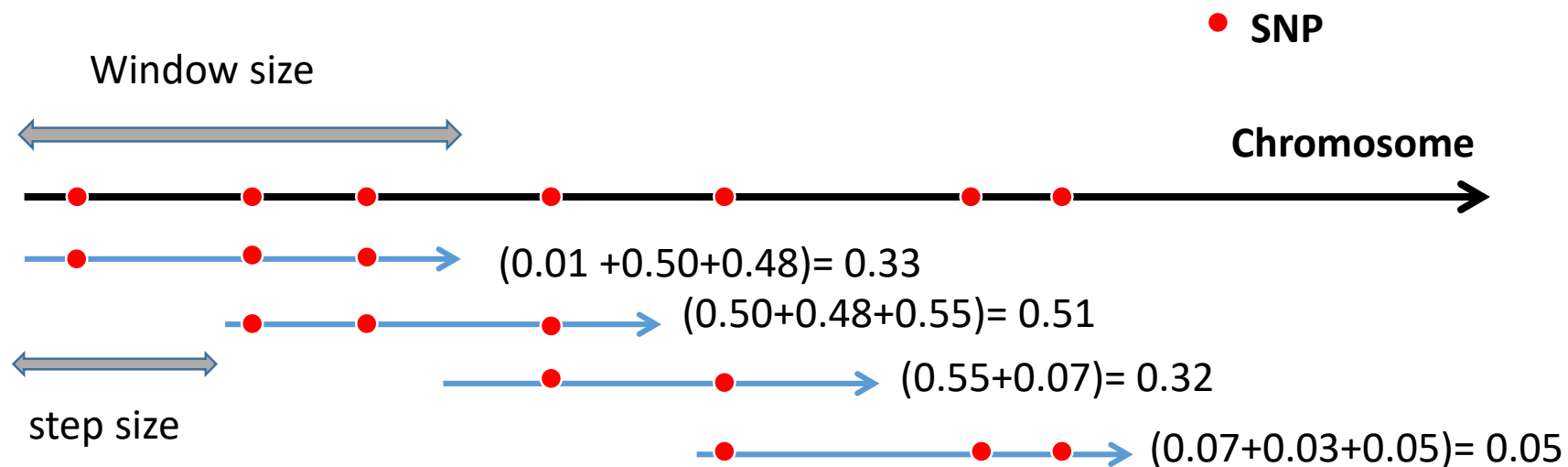


BSA Pipeline (part 2 Statistics and sliding window)

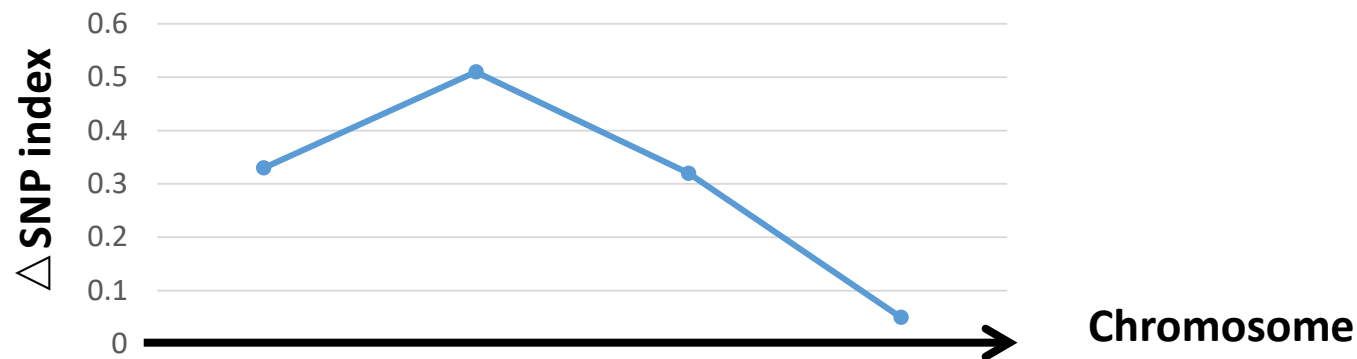
Linked sites



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



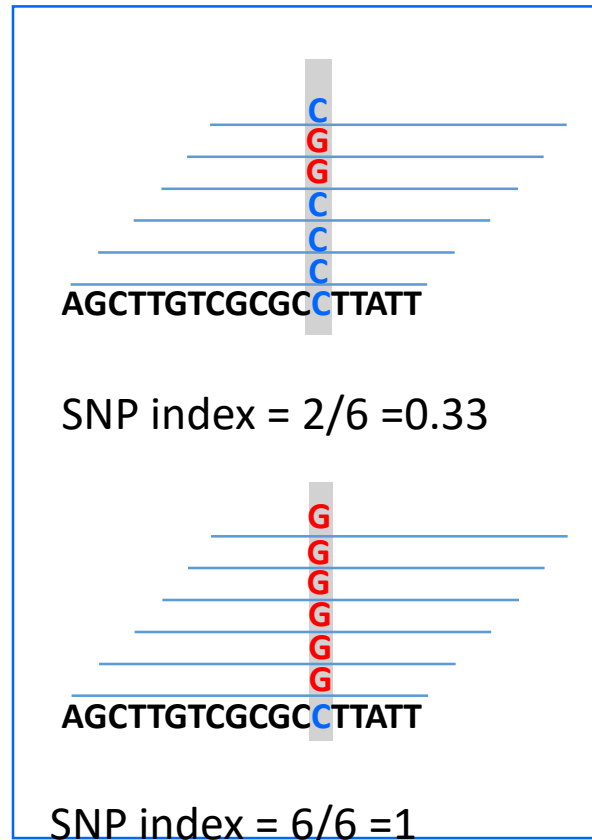
SNP-index across chromosome



Method 2. fishier exact test

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

Linked sites



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

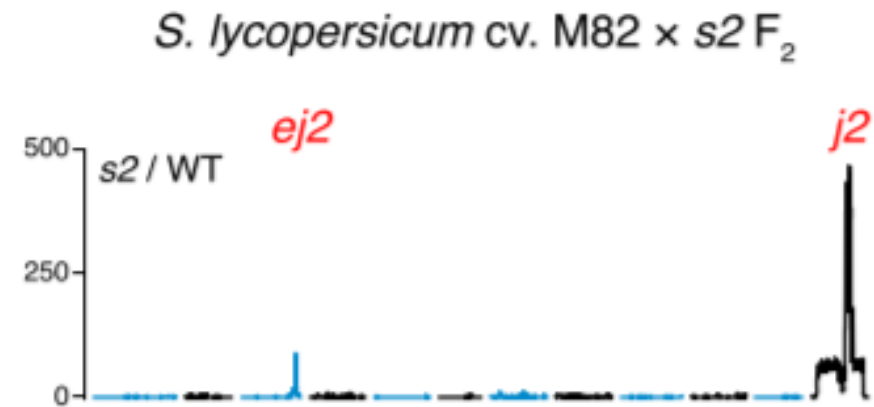
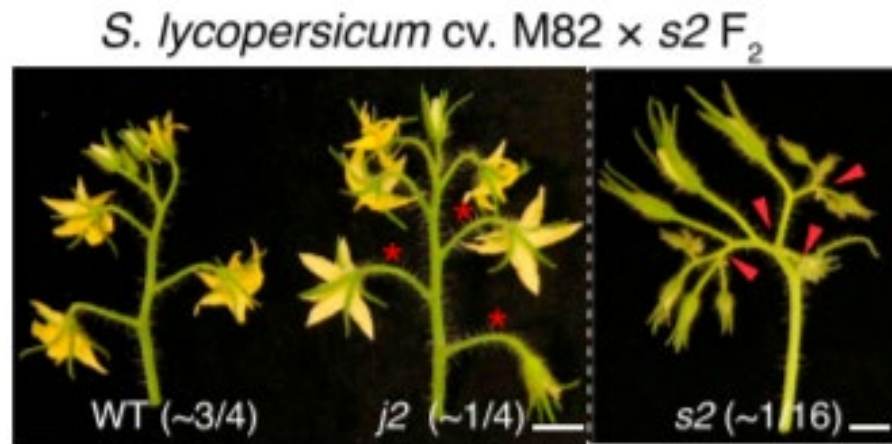
$$p = \frac{\binom{6}{4} \binom{6}{0}}{\binom{12}{4}}$$

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

```
F$p.value
```

```
0.06061
```

An exercise of BSA





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`

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
```

```
In -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

```
perl 00.src/01.variants_call.pl reads_table 03.bam/ 01.reference/reference.fasta
```



1. table with sample name and reads location.

2. The directory for the output. The output directory can not be an exist directory.

3. The reference file.

Reads_table is a tab delimited txt file

```
[chengzou@cbsuvitisgen2 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
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
```

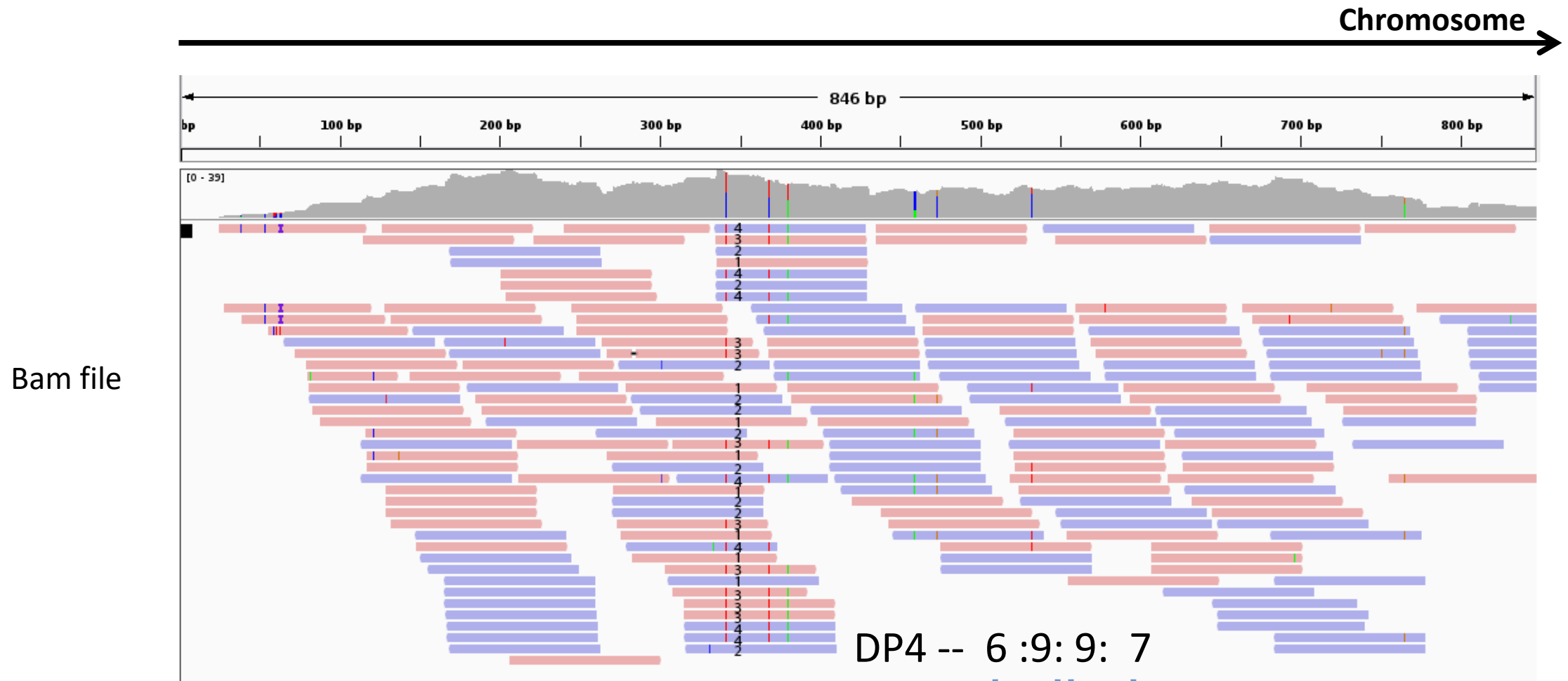
GT: Genotype

PL: list of Phred-scaled genotype likelihoods

DP: Number of high-quality bases

AD: Allelic depths

Definition of DP4 and AD



DP4 is Number of 1) forward ref alleles; 2) reverse ref; 3) forward non-ref; 4) reverse non-ref alleles

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 supported 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 -0 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@cbsuvisgen2 23.BSA_test]$ perl 00.src/01.variants_call.pl reads_table 04.bam/ 01.reference/reference.fast
a
[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; m
axMemory: 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

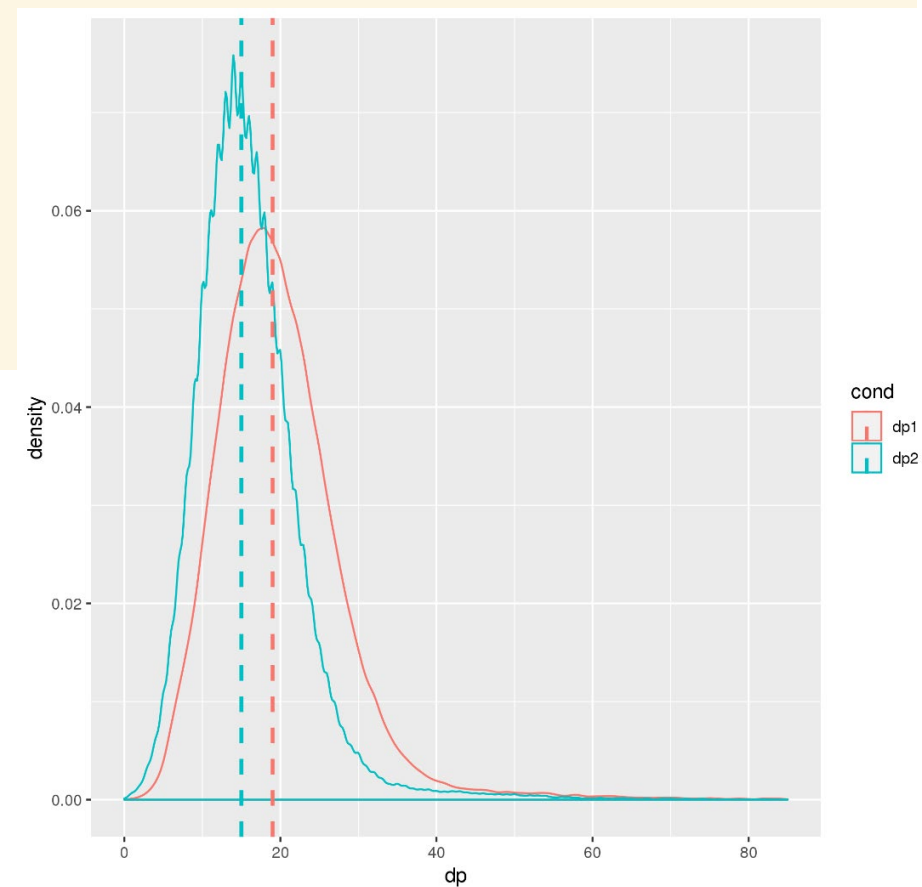
```
[chengzou@cbsuvitisgen2 03.bam]$ ls -l
total 2031636
-rw-rw-r-- 1 chengzou chengzou      1123 Nov 27 14:00 bwalog
-rw-rw-r-- 1 chengzou chengzou 10181013 Nov 27 14:00 filter.vcf.gz
-rw-rw-r-- 1 chengzou chengzou  4218553 Nov 27 14:06 filter.vcf.txt
-rw-rw-r-- 1 chengzou chengzou   955320 Nov 27 13:05 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

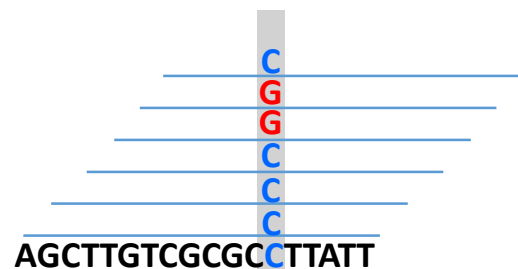
Examples:

```
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.
```

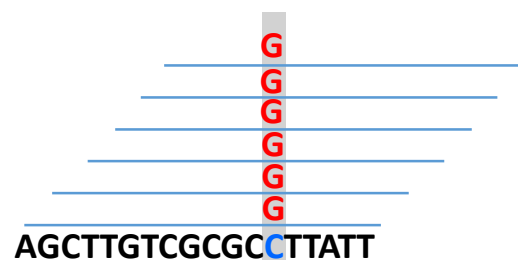
```
bcftools filter -i 'FORMAT/DP[1]<30 & FORMAT/DP[2]<34' filter.vcf.gz
-O z -o filter2.vcf.gz
```

Summary statistics 1. Δ SNP index

Linked sites



SNP index = $2/6 = 0.33$

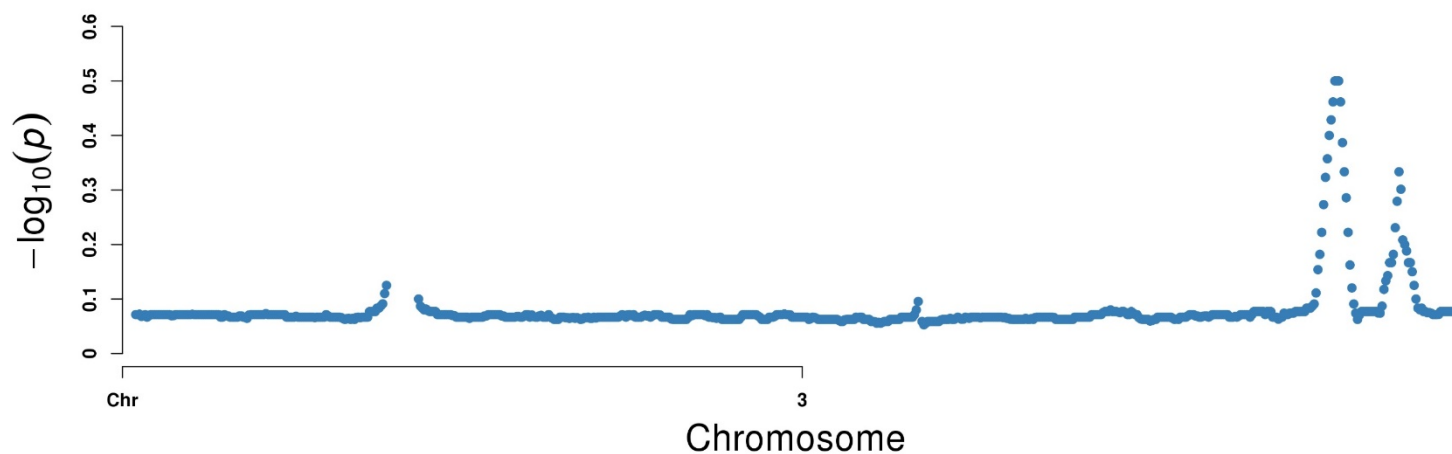


SNP index = $6/6 = 1$

```
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
```

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

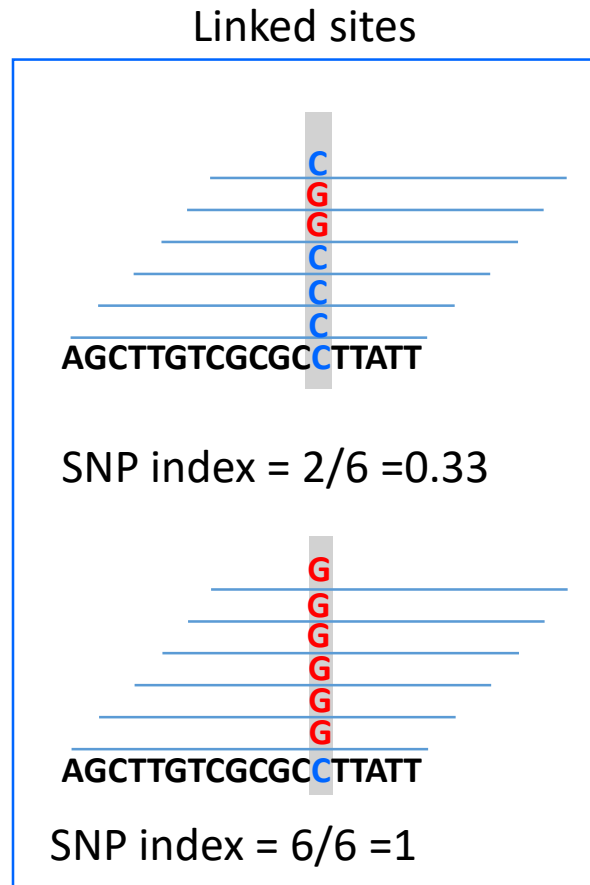
Manhattan plot of signal_filter.vcf.txt.abs_diff_window.txt.jpg



Summary statistics 2. ratio of allele frequency

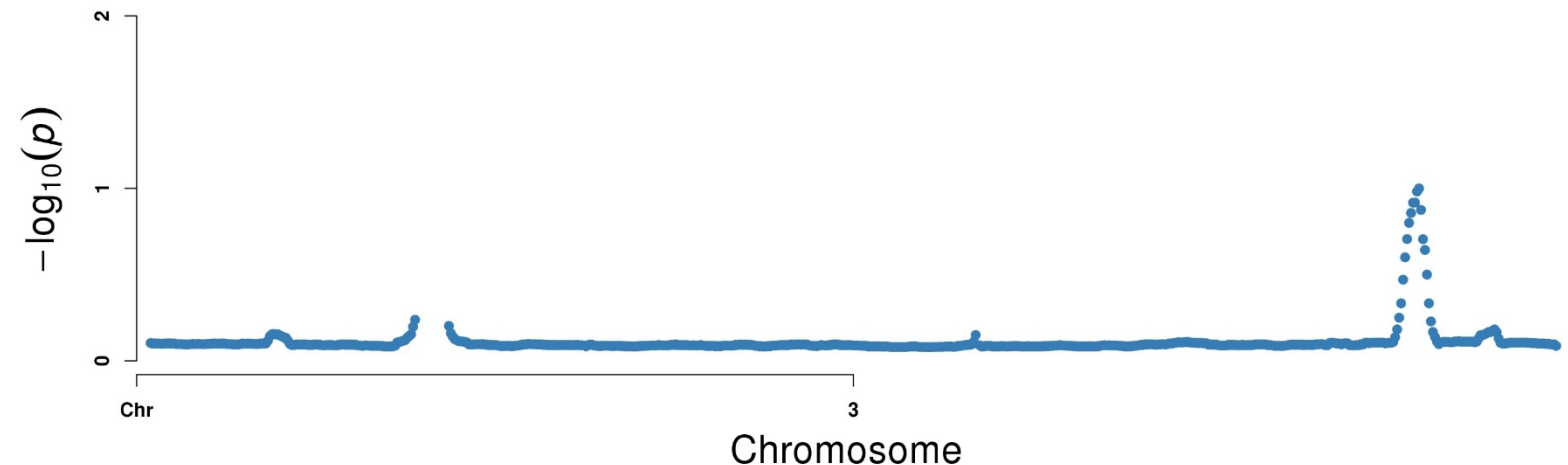
- 1. Compare the ratio, and sliding window to find the peaks.

```
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



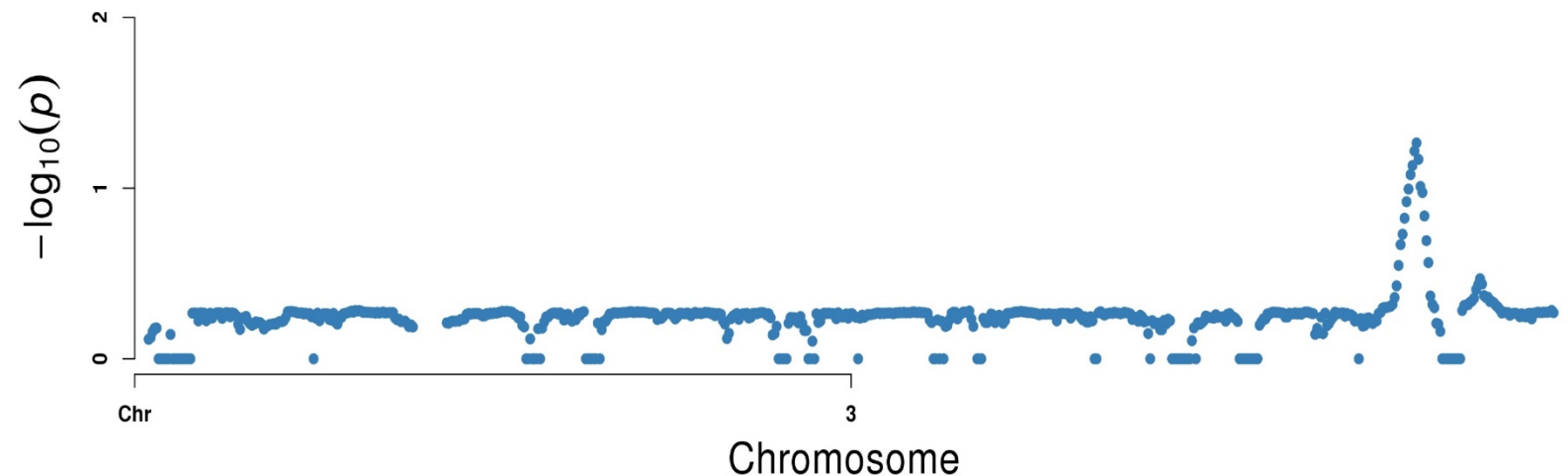
Summary statistics 3. fishier exact test

Linked sites



```
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
```

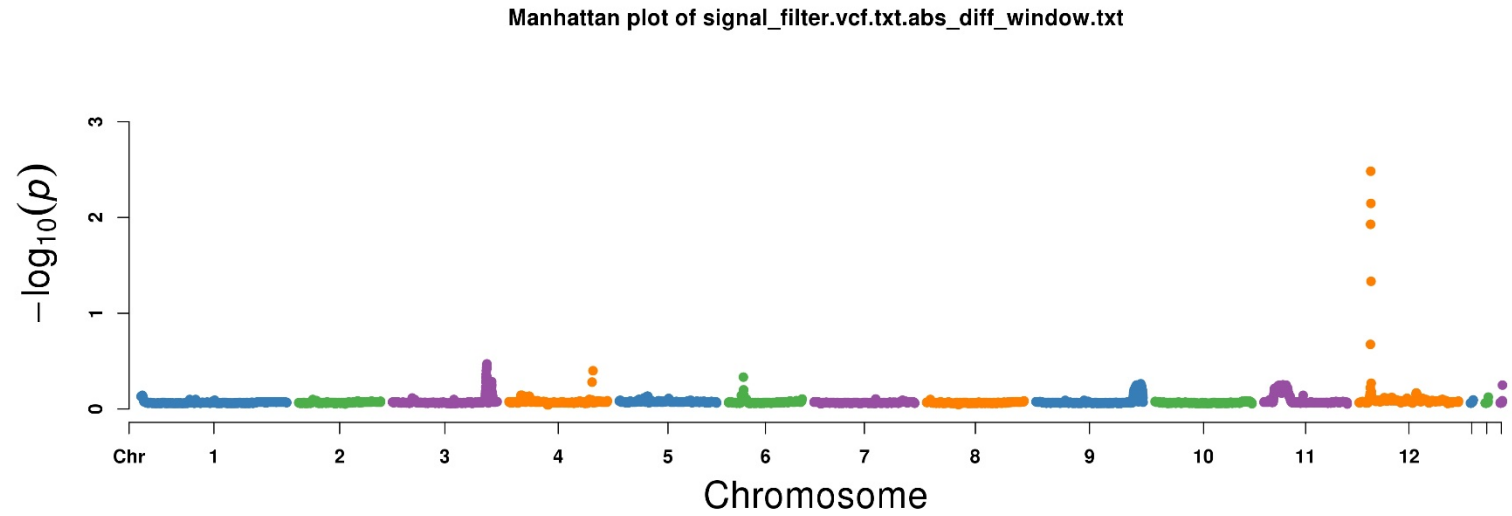
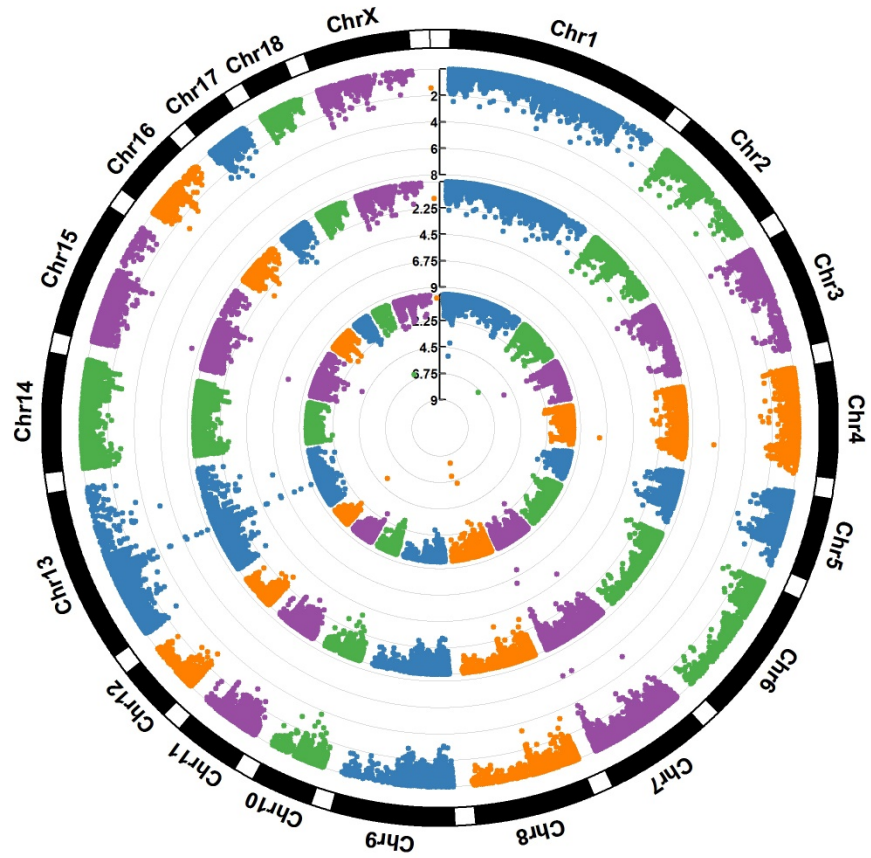
Manhattan plot of signal_filter.vcf.txt.fisher_window.txt.jpg



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)