HelmholtzZentrum münchen

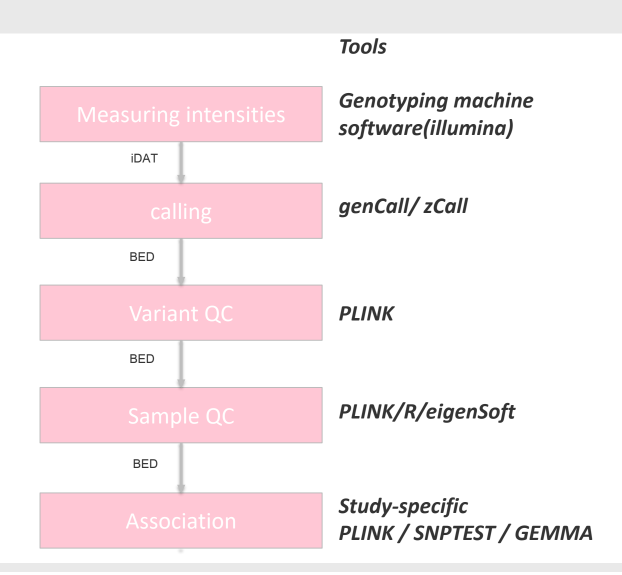
German Research Center for Environmental Health

Lecture 2

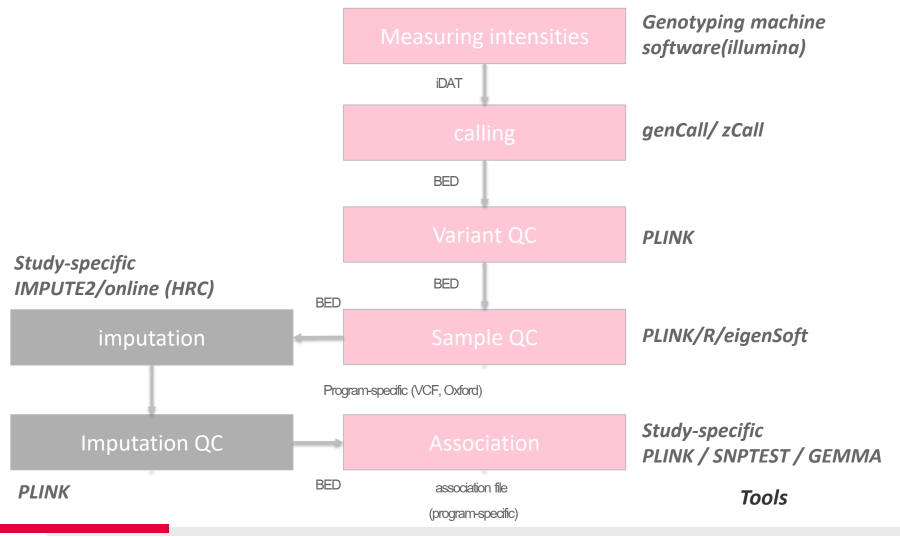
Current tools and best practices for performing genome-wide scans



The GWAS analysis pipeline



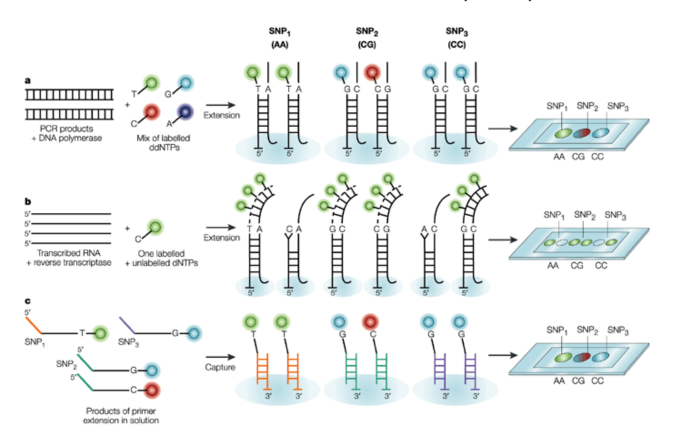
The (imputed) GWAS analysis pipeline

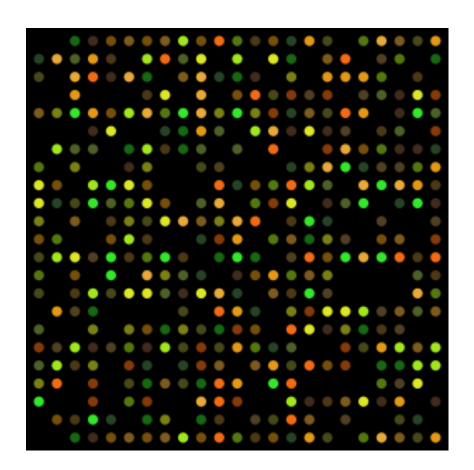


Genotyping data calling



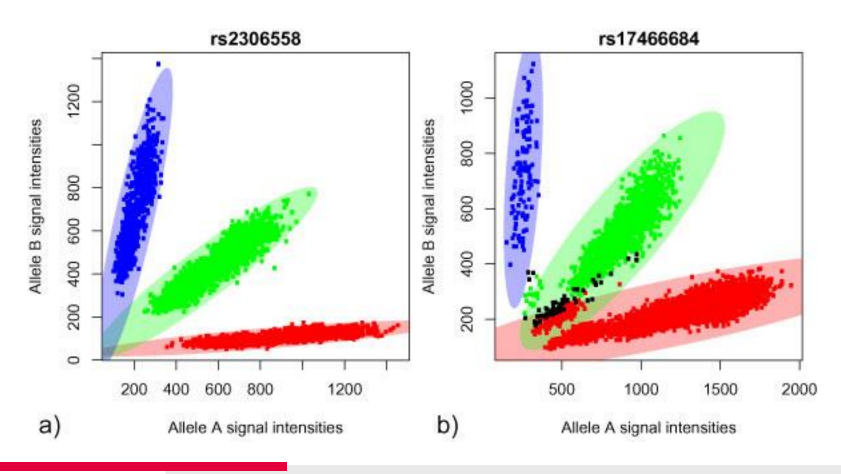
Did you say intensities?



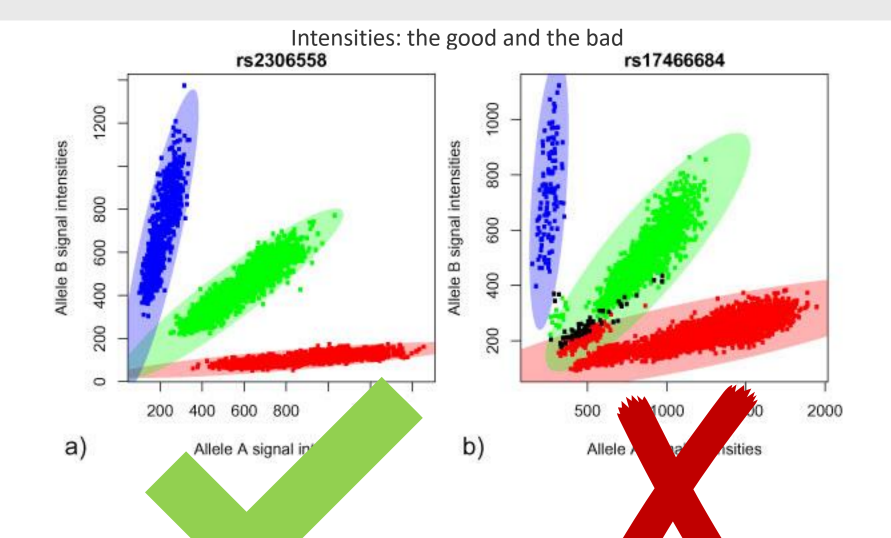


Nature Reviews | Genetics

Intensities: the good and the bad







Genotyping data storage



Which data types do we need?



Which data types do we need?

phenotype ~



Which data types do we need?

phenotype ~ genotype



Which data types do we need?

phenotype ~ genotype + covariates



Which data types do we need?

 $phenotype \sim genotype + covariates + structure$



Which data types do we need?

 $phenotype \sim genotype + covariates + structure + \epsilon$



Which data types do we need?

phenotype $\sim \beta \times genotype + covariates + structure + \epsilon$



Which data types do we need?

$$phenotype \sim \beta \times genotype + covariates + structure + \epsilon$$

$$\begin{bmatrix} pheno_0 \\ \vdots \\ pheno_n \end{bmatrix} \qquad \begin{bmatrix} A/T \\ \vdots \\ T/T \end{bmatrix} \qquad \begin{bmatrix} male \\ \vdots \\ female \end{bmatrix} \begin{bmatrix} 22 \ years \\ \vdots \\ 65 \ years \end{bmatrix} \begin{bmatrix} r_{00} & \dots & r_{0n} \\ \vdots & r_{ij} & \vdots \\ r_{n0} & \dots & r_{nn} \end{bmatrix}$$

Which data types do we need?

phenotype ~
$$\beta \times genotype + covariates + structure + \epsilon$$

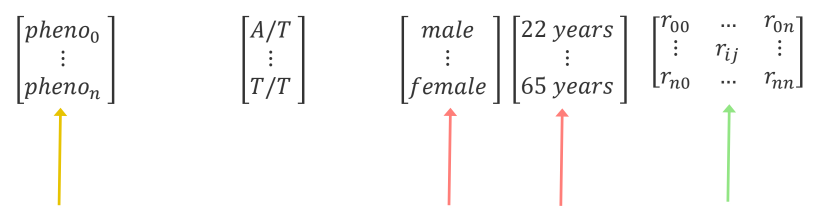
As we go from variant to variant...

$$\begin{bmatrix} pheno_0 \\ \vdots \\ pheno_n \end{bmatrix} \qquad \begin{bmatrix} A/T \\ \vdots \\ T/T \end{bmatrix} \qquad \begin{bmatrix} male \\ \vdots \\ female \end{bmatrix} \begin{bmatrix} 22 \ years \\ \vdots \\ 65 \ years \end{bmatrix} \begin{bmatrix} r_{00} & \dots & r_{0n} \\ \vdots & r_{ij} & \vdots \\ r_{n0} & \dots & r_{nn} \end{bmatrix}$$

Which data types do we need?

phenotype ~
$$\beta \times genotype + covariates + structure + \epsilon$$

As we go from variant to variant...

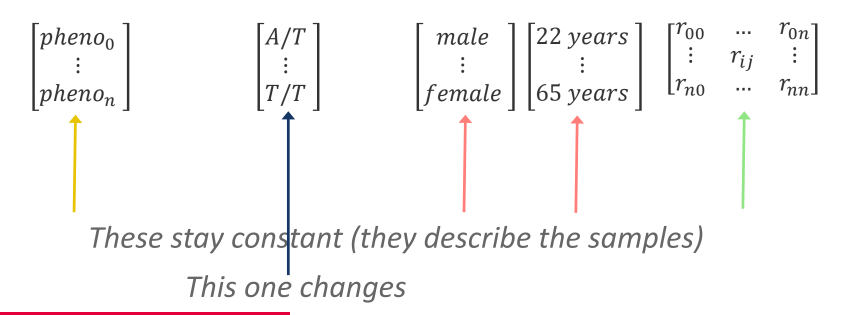


These stay constant (they describe the samples)

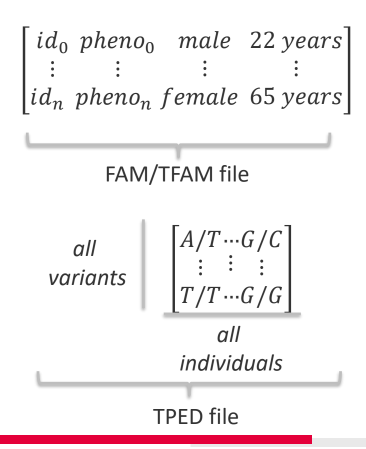
Which data types do we need?

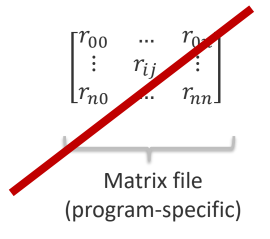
$phenotype \sim \beta \times genotype + covariates + structure + \epsilon$

As we go from variant to variant...



Our first format: TPED





Our first format: TPED

FAM/TFAM file

FAMILY1 SAMPLE1 0 0 1 22 1.5 FAMILY2 SAMPLE2 0 0 2 65 2.1

$$all$$
 $variants$
 $\begin{bmatrix} A/T \cdots G/C \\ \vdots & \vdots & \vdots \\ T/T \cdots G/G \end{bmatrix}$
 all
 $individuals$

TPED file

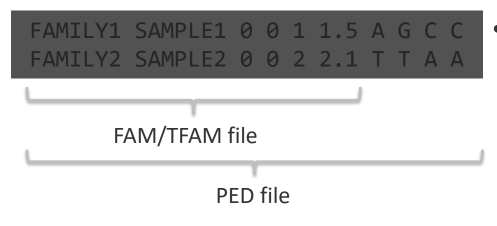
One of PLINK's traditional formats

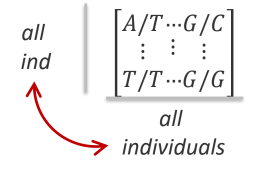
- Not used in practice
- Convenient for looping over SNPs
- Input --tfile
- Output --recode transpose

```
1 rs15933 0 752721 A G G G
1 1:846808 0 846808 C C T C
```



Another format: PED/MAP





- One of PLINK's traditional formats
 - Not used in practice
 - Convenient for looping over samples
 - Input --file
 - Output --recode

```
1 rs15933 0 752721
1 1:846808 0 846808
MAP file
```

Exercise 1: Format conversion

- In /Workshop_data/Lecture2/Exercise1
 - Convert cohort1.tped/tfam to cohort1.ped/map
 - Use the transpose.sh script provided or try to d.i.y

```
FAMILY1 SAMPLE1 0 0 1 1.5 A G C C FAMILY2 SAMPLE2 0 0 2 2.1 G G T C
```

- Convert the file to PED using PLINK
- Compare both files using diff



Exercise 1 : Solution

- cut -d' '-f1-4 cohort1.tped | tr ' ' '\t'> cohort1.map
- paste -d' 'cohort1.tfam <(./transpose.sh <(cut -d' '-f5- cohort1.tped)) > cohort1.ped
- plink --tfile cohort1 --recode --out fortest
- diff cohort1.ped fortest.ped



Exercise 2 : Storage

- Consider 3 different genotyping chips
 - 500,000 SNPs (Illumina OmniExpress)
 - 1,000,000 SNPs (ExomeChip)
 - 2,500,000 SNPs (Illumina Onmi 2.5)
 - How large is a PED file containing genetic information for 10,000 samples on each of these chips?



Exercise 2 : Storage

- Consider 3 different genotyping chips
 - 500,000 SNPs (Illumina OmniExpress)
 - 1,000,000 SNPs (ExomeChip)
 - 2,500,000 SNPs (Illumina Onmi 2.5)
 - 1 character = 1 byte
 - Each genotype = 2 alleles + 2 spaces = 4 characters

$$n_{SNPS} \times 4 \times n_{individuals} = 19 \ Gb$$
, 38Gb, 95Gb

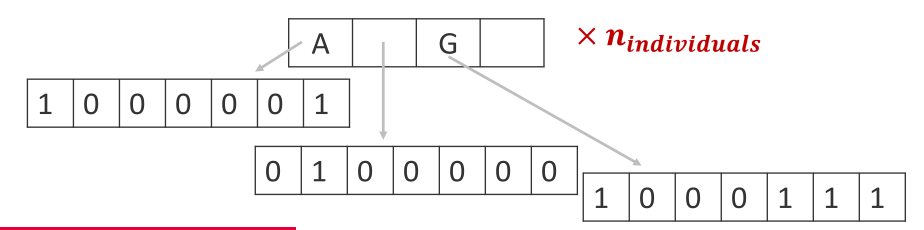
Binary formats

- 1 character = 1 byte
- Each genotype = 2 alleles + 2 spaces = 4 characters
- Can we make this better?



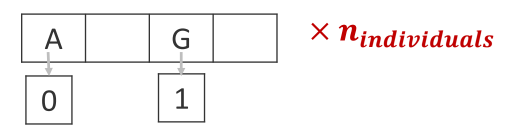
Binary formats

- 1 character = 1 byte
- Each genotype = 2 alleles + 2 spaces = 4 characters
- Can we make this better?
- 2 solutions
 - Compress using ZIP/GZIP
 - Use binary formats



Binary formats

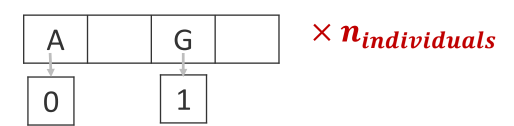
- 1 character = 1 byte
- Each genotype = 2 alleles + 2 spaces = 4 characters
- Can we make this better?
- 2 solutions
 - Compress using ZIP/GZIP
 - Use binary formats



Question: how smaller is the size now?

Binary formats

- 1 character = 1 byte
- Each genotype = 2 alleles + 2 spaces = 4 characters
- Can we make this better?
- 2 solutions
 - Compress using ZIP/GZIP
 - Use binary formats



Question: how smaller is the size now?

$$\frac{n_{SNPS} \times n_{individuals}}{4} = 1.1Gb$$

One (binary) format to rule them all: BED/BIM/FAM

```
FAMILY1 SAMPLE1 0 0 1 22 1.5 FAMILY2 SAMPLE2 0 0 2 65 2.1
```

```
1 rs15933 0 564862 C T
1 1:752566 0 752566 G A
```

FAM/TFAM file

BIM file

```
      10101111
      10101111
      10100010
      10111011
      10101000
      10000000

      00101011
      00100000
      10101000
      10001011
      00000011
      11111111

      11111111
      11111111
      111111110
      111111110
      111111111
```

BED file

- Input: --bfile
- Output: --make-bed
- Do not open a BED file with less, cat, head, or tail!
- If you absolutely want to look, xxd -b or od -c

Genotyping data: common operations



Common operations

Sample management

keep [file]	Keep samples in file
remove [file]	Remove samples in file

SNP management

extract [file]	Keep SNPs in file
exclude [file]	Remove SNPs in file

Extracting regions

chr [name]	Extract data on specified chromosome
from-bp [pos]	From specified position
to-bp [pos]	To specified position



Common operations

Variant QC

maf [threshold]	Keep variants with MAF>threshold
hwe midp [threshold]	Keep variants with HWE p>threshold

Sample QC

missing	Compute per-sample and per-variant missingness
check-sex	Check sexes by looking at chrX
genome	Compute relatedness, check for duplicates

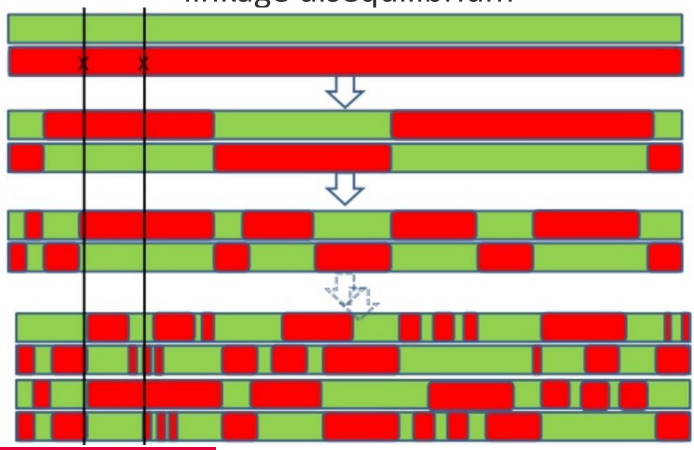


Linkage disequilibrium

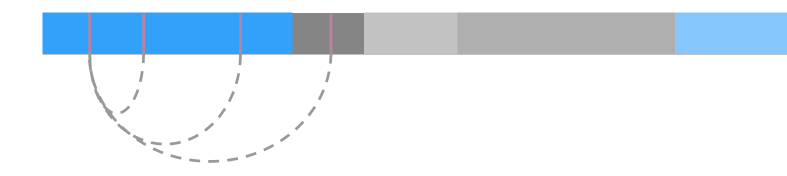


Linkage Disequilibrium

The strange beautiful world of linkage disequilibrium

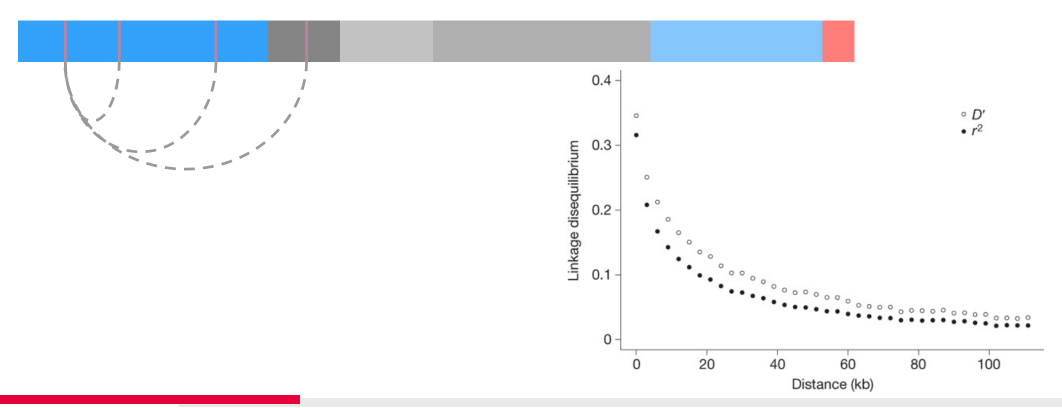


The strange beautiful world of linkage disequilibrium





The strange beautiful world of linkage disequilibrium



LD between 2 or more SNPs

```
plink --r2 --ld-snps rs1234,rs4567
plink --r2 --ld-snp-list [file]
```

Pairwise LD in a region

```
plink --r2 --ld-window 10 --ld-window-kb 1000 -
-ld-window-r2 0.2
```

LD-pruning (only independent SNPs)

```
plink --indep 50 5 2

plink --indep-pairwise 50 5 0.2

Plink --indep-pairphase 50 5 0.2
```



Exercise 3: Stretching the PLINK muscle

- In /Workshop_data/Lecture2/Exercise3
 - How many common (MAF>5%) variants are there on chromosome 11 in the `cohort1` dataset?
 - How many variants are in LD (r2>0.4) with 21:28759840 on chromosome 21 in a 1Mbp window?



Exercise 3: Stretching the PLINK muscle

```
plink --bfile cohort1 --maf 0.05 --chr 11 --out chr11 --make-bed
```

wc -l chr11.bim

```
plink --bed cohort1.bed --bim cohort1.bim --fam cohort1.fam --r2 --ld-snp 21:28759840 --ld-window-kb 1000000 --ld-window 1000000 -ld-window-r2 0.4
```

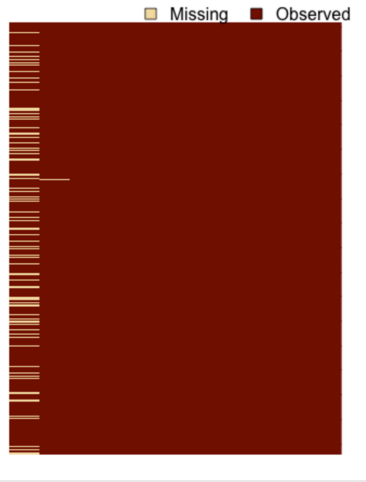
wc -l plink.ld



QC steps

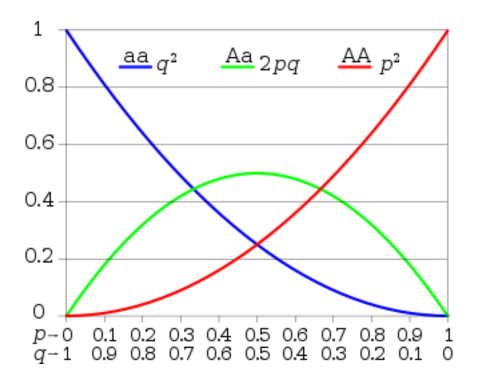
Variant QC: which variants do we want

to remove?



Variant QC: which variants do we want to remove?

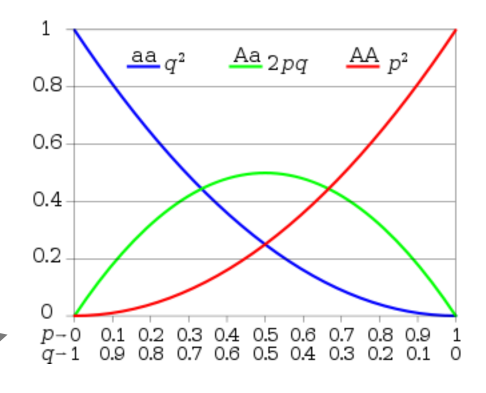
 Calling is not perfect: some genotypes are missing



Variant QC: which variants do we want to remove?

 Calling is not perfect: some genotypes are missing

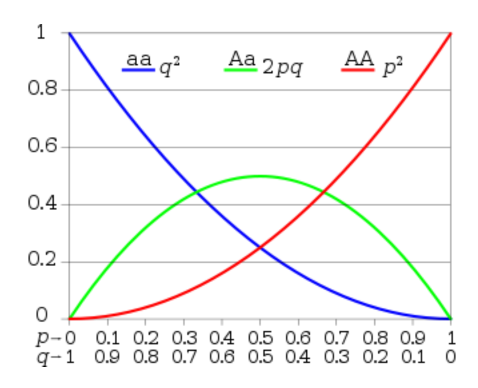
If we see that 40% of all alleles are a, what is the proportion of aa, Aa, AA?





Variant QC: which variants do we want to remove?

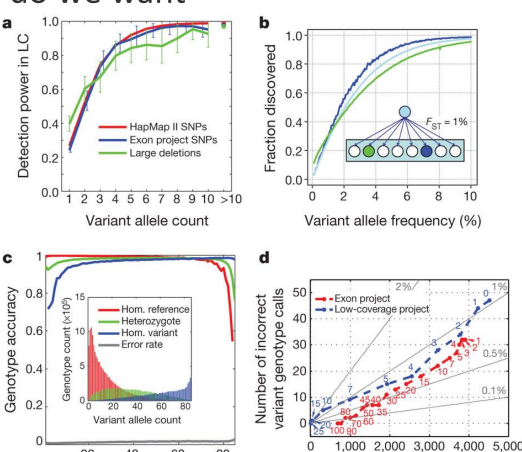
- Calling is not perfect: some genotypes are missing
- Variants violating Hardy-Weinberg equilibrium are improbable



Variant QC: which variants do we want

to remove?

- Calling is not perfect: some genotypes are missing
- Variants violating Hardy-Weinberg equilibrium are improbable

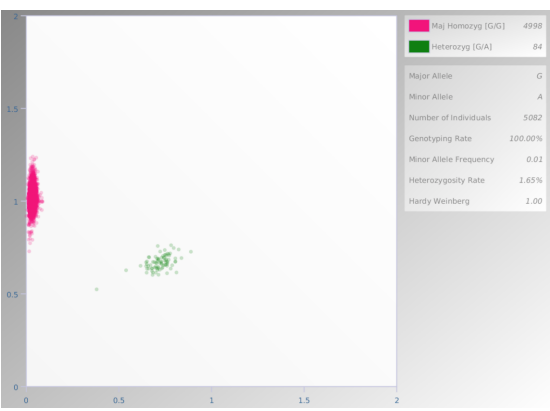


Variant allele count

Number of variant genotype calls

Variant QC: which variants do we want to remove?

- Calling is not perfect: some genotypes are missing
- Variants violating Hardy-Weinberg equilibrium are improbable
- Rare variants are difficult to call





Sample QC: which individuals do we want to remove?

All the different ways in which our samples could be the wrong ones

What are some defining sample characteristics?



Sample QC: which individuals do we want to remove?

All the different ways in which our samples could be the wrong ones

What are some defining sample characteristics?





Sample QC: which individuals do we want to remove?

All the different ways in which our samples could be the wrong ones

What are some defining sample characteristics?

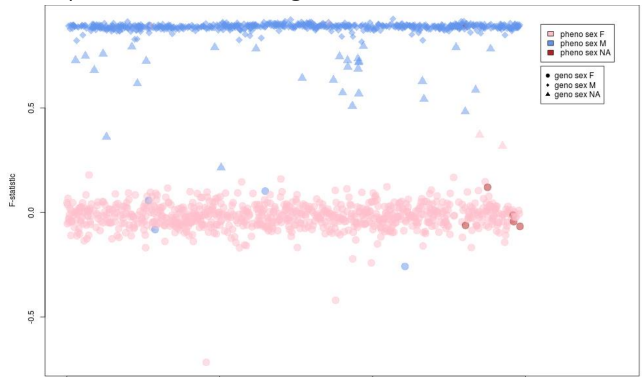




Sample QC: which individuals do we want to remove?

All the different ways in which our samples could be the wrong ones

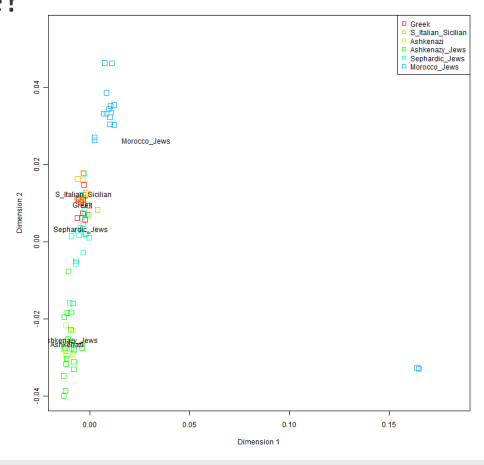
Sex checks





Sample QC: which individuals do we want to remove?

• Ethnicity checks





Sample QC: which individuals do we want to remove?





Sample QC: which individuals do we want to remove?





Sample QC: which individuals do we want to remove?

