# Advanced Gene Mapping Course

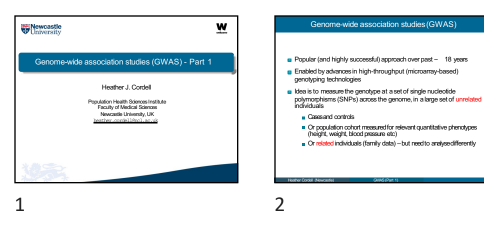
April 22-26, 2024 The Rockefeller University New York, NY

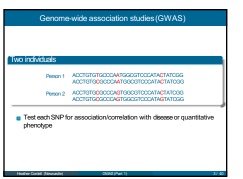
Lectures

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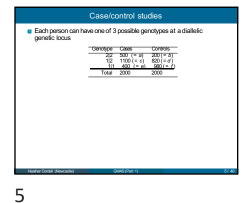
Lectures given by: <sup>1</sup>Heather Cordell, <sup>2</sup>Suzanne Leal, <sup>3</sup>Gao Wang, <sup>4</sup>Jurg Ott; <sup>5</sup>Andrew DeWan, <sup>6</sup>Wayne Patterson; and <sup>7</sup>Shamil Sunyaev

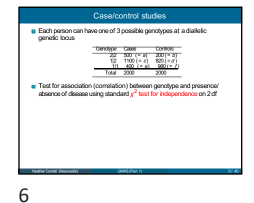


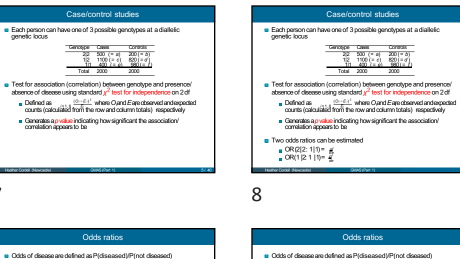


## Association testing: case/control studies Collect sample of affected individuals (cases) and unaffected individuals (controls) Or a else a sample of random "population" controls Most of whom will not have the disease of interest Examine the association (correlation) between alleles present at a genetic locus and presence/absence of disease By comparing the distribution of genotypes in affected individuals with that seen in controls

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- Odds of usease are defined as "(useased)" (fith useased)
   Odds ratio OR (2):2.11) represents the factor by which your odds of
   deseasemust bemultiplied, if you have genotype 2)/2asopposed to 1|1
   i.e. the 'effect' of genotype 2|2
  - Similarly, we can define the OR for 1|2vs 1|1
     As the factor by which your odds of disease must be multiplied, if you have gendype 1|2as opposed to 1|1
     i.e. the 'effect' of gendype 1|2

GWAS (Part

 Odds ratio OR (2|2: 1|1) repesents the factor by which your odds of disease must be multiplied, if you have genotype 2|2as opposed to 1|1

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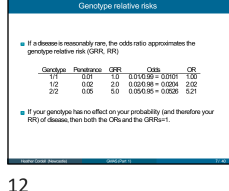
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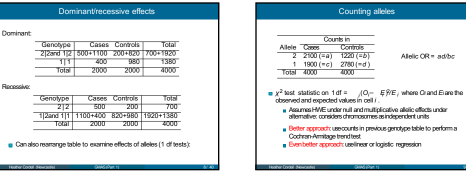
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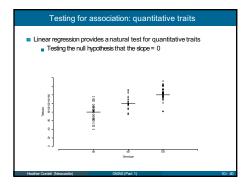
### Odds ratios

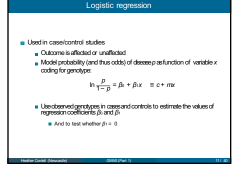
- Odds of disease are defined as P(diseased)/P(not diseased)
   Odds ratio OR (2|2: 1|)repsents the factor by which your odds of disease multiplied, if you have genotype 2|2as opposed to 1|1
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   As the factor by which your odds of dsease must be multiplied, if you have genotype 1|2as opposed to 1|1
  - i.e. the 'effect' of genotype 1|2
- ORs are closely related (often ≈) genotype relative risks
   The factor by which your probability of disease must be multiplied, if you have genotype 1|2asopposed to 1|1(say)
- If your genotype has no effect on your probability (and therefore on your odds) of disease, then the ORs=1.
  - So the association test can be thought of as a test of the null hypothess that the ORs=1

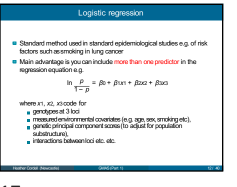


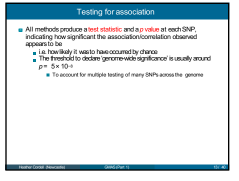


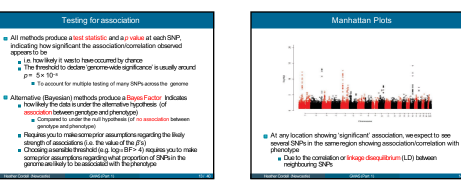


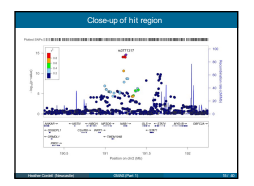




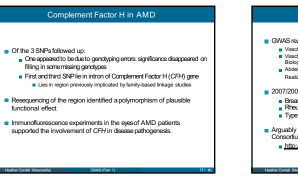














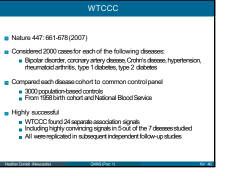
Historical Perspective: Complement Factor H in AMD

- First (?) GWAS was by Klein et al. (2005) Science 308:385-389
- Typed 116,204 SNPs in 96 cases (with age-related macular degeneration, AMD) and 50 controls
- Very small sample size—they were very lucky to find anything! Luck
   wesdue to the fact the polymorphism has a very large effect (recessive OR=7.4)
- Klein et al. followed up on two SNPs passing threshold (p < 4.8×10<sup>-7</sup>)
- Plus a third SNP that just failed to pass significance threshold, but lay in same region as first SNP

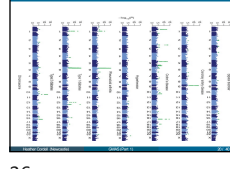
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# GWAS GWAS really got going in around 2007 Visscher et al. (2012) AIHG 907-24 'Five Years of GWAS Discovery' Visscher et al. (2012) AIHG 1015-22 '10 Years of GWAS Discovery: Biology, Function and Translation' Addellaudie et al. (2023) AIHG 1015-194 '15 Years of GWAS Discovery: Realizing the promise' 2007/2008 saw as lew of high-profile GWAS publications Proceed concer (Easbin et al. 2007) Rheumaloid Arthritis (Plenge et al. 2007) Rheumaloid Arthritis (Plenge et al. 2007; Zeggini et al. 2008) Agueably the most influential was the Wellcome Trust Case Control Consortium (WTCCC) study of 7 different diseases http://www.wtocc.org.uk/





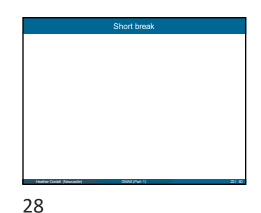


Manhattan plots for 7 diseases

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### Lessons from WTCCC (and others)

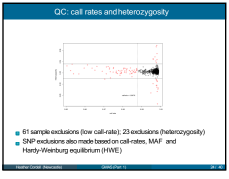
- Typically used rather standard statistical/epidemiological methods (X<sup>2</sup> tests, t tests, logistic regression etc.)
- Success largely due to:
  - An appreciation of the importance of large sample size ( > 2000 cases, similar or greater number of controls)
  - Stringent quality control procedures for discarding low-quality SNPs and/or samples
  - Stringent significance thresholds (p = 5×10-8) to account for multiple testing and/or low prior prob of true effect
  - Importance of replication in an independent data set

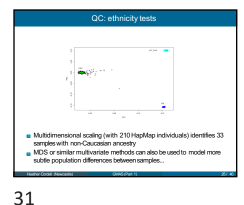


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# Quality Control Stringent QC checks are required for GWAS data Discard samples (people) deemed unreliable Low genchype call rates, excess beterozygosity etc. X chromosomer markers useful for checking relationships and ethnicity Males should "appear homozygous at all X markers Obscard data from SNPs deemed unreliable Nakes for genchype call rates, Mendelian misinheritances, Hardy-Weinberg disequilibrium Stacke SNPs with how more dise frequency (MAF) Sectutorials at: https://pubmed.ncbi.nlm.nih.gov/2048/1742/ Meter Coll (Mexet) Marker Sneed (Mexet)







Multivariate Analysis

### Multivariate Analysis

- Several related multivariate analysis techniques have been proposed for detecting population structure in genome-wide association studies Principal components analysis (PCA)
  - Principal coordinates analysis (PCoA)
     Multidimensional scaling (MDS)
- If population differences can be detected (and adjusted for) in association analysis, this offers a way to deal with the problem of
- population stratification Population sampled actually consists of several 'sub-populations' that
- do not really intermix Can lead to spurious false positives (type 1 errors) in case/control studies

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   Principal components analysis (PCA) Principal coordinates analysis (PCoA)
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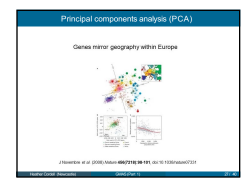
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- If population differences can be detected (and adjusted for) in association analysis, this offers a way to deal with the problem of population stratification
  - Population sampled actually consists of several 'sub-populations' that do not really intermix
  - Can lead to spurious false positives (type 1 errors) in case/control studies

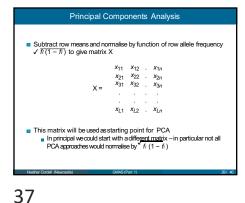
These techniques can also be used in quality control (QC) procedures, to check for (and discard) gross population outliers

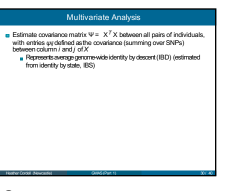
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Principal Components Analysis Price et al. (2006) Nature Genetics 38:904-909; Patterson et al. (2006) PLoS Genetics 2(12):e190 Based on popn genetics ideas from Cavalli-Sforza (1978) Idea is to form a large matrix M of SNP counts (0,1,2) corresponding to the genotype at a L loci (=rows) for n individuals (=columns) g11 g12 · g1n g<sub>21</sub> g<sub>22</sub> . g<sub>2n</sub> g<sub>31</sub> g<sub>32</sub> . g<sub>3n</sub> M = gL1 gL2 · gLn





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### Multivariate Analysis

- Estimate covariance matrix Ψ = X<sup>T</sup>X between all pairs of individuals, with entries  $\psi_{ij}$  defined as the covariance (summing over SNPs) between column *i* and *j* of X
  - Represents average genome-wide identity by descent (IBD) (estimated from identity by state, IBS) ■ Compute the eigenvectors -vand eigenvalues λ/ of matrix Ψ ■ Co-ordinate / of the kth eigenvector represents the ancestry of individual j along 'axis' k

### Multivariate Analysis

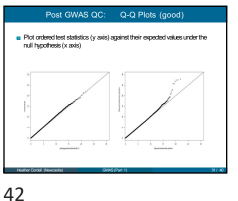
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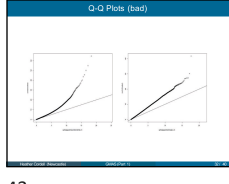
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### Multivariate Analysis

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     Co-ordinate i of the kth eigenvector represents the ancestry of individual j along 'axis' k
- For technical details, see McVean (2009) PLoS Genetics 5;10:e1000686
- Many genetics packages e.g. (PLINK) will allow you to calculate the
- top 10 (or more) PCs Different geographic populations can often be well separated by just the first two or three PCs
- Useful for outlier detection For more subtle differences, you may need to calculate more PCs
- And include them as covariates in the regression equation Post-GWAS QC can determine whether you have included 'enough'







### Population stratification

- A QQ plot showing constant inflation (straight line with slope > 1) can indicate population stratification/population substructure
- Simple solution: Genomic Control (Devlin and Roeder1999)
   Useyour observed test statistics to estimate the slope (=inflation factor λ)
   Divide each test statistic by λ to get an adjusted (deflated) test
- statistic
- More complicated solution: use PCA/MDS or similar Even
   more complicated solution: use linear mixed models

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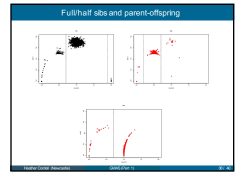


- Using 'thinned' subset of markers with high minor allele frequency (MAF) and in approximate linkage equilibrium
- Simple relationships (PO, FS, MZ/duplicates) can identified with only a few hundred markers
- More complicated relationships require 10,000-50,000 SNPs
- Various software packages, including PLINK, KING and TRUFFLE

ssuming no inbreeding, the IBD s	state pr	obabilitie	s are:
	Number of alleles shared IBD		
Relationship	2	1	0
MZ twins	1	0	0
Parent–Offspring	0	1	0
Full siblings	1/4	1/2	1/4
Half siblings	0	1/2	1/2
Grandchild-grandparent	0	1/2	1/2
Uncle/aunt-nephew/niece	0	1/2	1/2
First cousins	0	1/4	3/4
Second cousins	0	1/16	15/16
Double 1st cousins	1/16	6/16	9/16

(as estimated across the genome) Or kinship coefficient { P(IBD=2)+ P(IBD=1) } againstP(IBD=0)

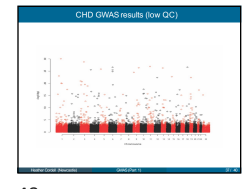
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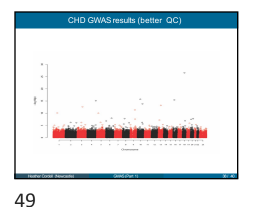


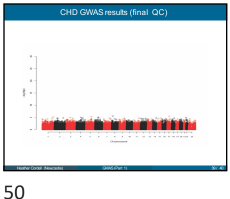


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### Genome-wide meta-analysis

Puts together data (or results) from a number of different studies
 Could analyse asone big study
 But preferable to analyse using meta-analytic techniques
 At each SNP construct an overall test based on the results
 (log ORs and standard errors) from the individual studies

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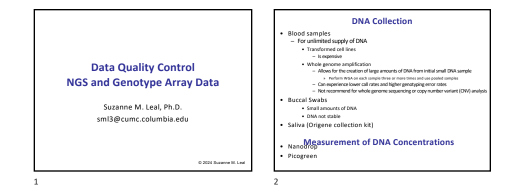
 Meta-analysis is often made easier by using imputation

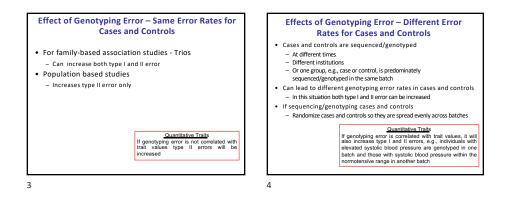
 Inferring (probabilistically) the genotypes at SNPs which have not actually bein genotyped
 On the basis of their incover correlations with nearby SNPs that have been genotyped
 Using artiference panel of people (e.g. 1000 Genomes) who have been genotyped to the SNPs

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### Genotype SNPs (~20-96) before Exome or Whole Genome Sequencing

- · Genotype markers which can be used as DNA fingerprint
- · Allows for Assessment of DNA quality
- Aids in determining the the genetic sex of study subjects
   To aid in identification of potential sample swaps
- Detects cryptic duplicates
- For family data

Aids in determining close familial relationships
 Non-paternity

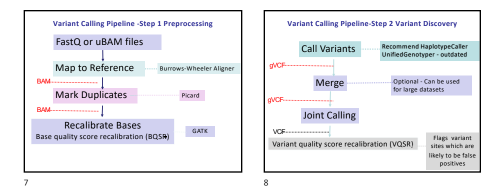
Sample swaps
Cryptic relationships

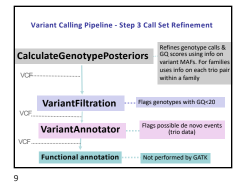
### Detecting Genotyping Errors

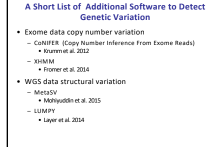
- Duplicate samples genotyped using arrays to detect inconsistencies
- Can use duplicate samples that are inconsistent to adjust clusters to improve allele calls
   Will not detect systematic errors
- will not detect systematic errors
- Usually generated only for genotype array data

   Due to expense, duplicate samples are usually not generated for exome or whole genome sequencing studies

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### Variant Calling

- BAM files are large and take considerable resources
- Storage is expensive
- One 30x whole genome is ~80-90 gigabytes
- A small study of 1,000 samples will consume 80 terabytes of disk space
- The cost of cloud computing to call variants
- (Souilmi et al. 2015)
- \$5 per exome

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- \$50 per genome
- For 1,000 samples

   \$5,000 exome
- \$50,000 genome
- \$50,000 gend

### Working with gVCF Files

Instead of obtaining VCF files

- Can obtain gVCF files to perform joint calling and complete the GATK pipeline
- A whole genome gVCF
  - ~1 Gigabyte

     1/100<sup>th</sup> the size of a BAM file for one individual

### **Influences on Sequence Quality**

- DNA quality
- Age of sample
- Extraction method
- Source of sample • e.g., blood, skin punch, buccal
- Sequencing machines (read length)
- Median sequencing depth
- Alignment
- Variant calling method used
- Single nucleotide variants and insertion/deletions
- Structural variants

### NGS Data Quality Control

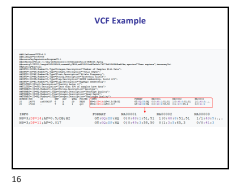
- · Extremely important to perform before data analysis
- Poor data quality can increase type I and II errors - Due to inclusion of false positive variant sites or incorrect
- genotype calls · Protocols for data QC are still in their infancy
- No set protocols for QC QC is data specific
- Dependent on read depth
- Batch effects
- Availability of duplicate samples
- etc.

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- NGS Data Quality Removal of Genotype Calls and Samples Sequence depth of coverage DP\_variant High DP could be an indication of copy number variants Which can introduce false positive variant calls
   » Due to down sampling in GATK maximum DP is 250 DP\_genotype Concerned if depth is too low or too high Low insufficient reads to call a variant site
   Remove genotypes with low read depth, e.g., DP<8</li> Genotype quality (GQ) score Removal genotypes with a low genotype quality core, e.g., GO< 20</li> Bcftools
- Can be used to remove variants sites and genotypes which do not meet quality control criteria

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### Variants with more than 2 Alleles · Genetic analysis tools are usually developed to analyze

- variant sites that are diallelic
- Some sites may have >2 alleles
- · The alleles at these sites need to be split
- New loci are made each multi-allelic site each with only 2 alleles bcftools

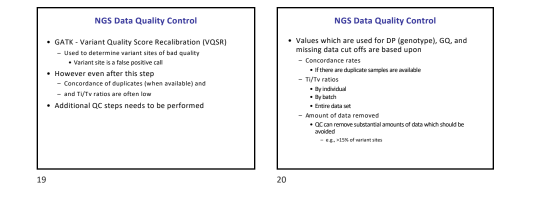
• Multiallelic sites can have higher error rates compared to diallelic sites

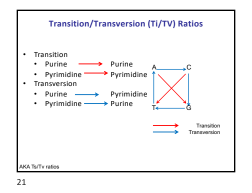


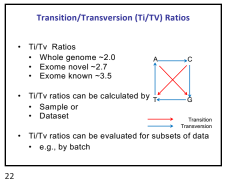
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### NGS Data Quality – Removal of Genotype Calls and Samples · Removal of sites with missing data

- e.g., missing > 10% of genotypes
- Removal of "novel" variant sites which only occur in one batch and the alternative allele is observed multiple times or the minor allele frequency (MAF) is
- high in overall sample · Removal of sites that deviate from Hardy-Weinberg Equilibrium (HWE)
- Must be performed by population, e.g., African American and European American
- Related individuals should be removed from the sample before testing for deviations from HWE







### Sequence Data QC Overview

- · Variant and genotype call level
- Evaluation of batch effects
- Genotype call level Removal of genotype calls
- Low or high depth of coverage DP< 8</li>
- Low genotype quality score GQ< 20</li>
  Removal of individual samples
- >20% missing data
- After taking the intersect of capture arrays
- Samples without phenotype information

### Sequence Data QC Overview

- Variant level removal of variant sites
- Low call rate
- i.e., missing call rate > 10%
- "Novel" variant sites observed≥2 only in a single batch
   Deviation from Hardy-Weinberg-Equilibrium
- Population specific
- Unrelated individuals

   e.g., p<5 x 10<sup>8</sup>, p<5x10<sup>-15</sup>

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### QC – Assessing Sex Chromosomes

- When data is collected on study subjects they are asked about their gender/sex and not their genetic sex - Differences in gender/sex and genetic sex can be due to
  - Sample swaps Study subjects who are not cisgender
- · Some study subjects may have neither a XX nor XY karyotype
  - Turner syndrome X0
  - Klinefelter syndrome XXY

### QC – Assessing Sex Chromosomes

- · Study subjects labeled as females with an excess of homozygous genotypes on the X chromosome can denote
- That their genetic sex is male

- Turner Syndrome

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### QC – Assessing Sex Chromosomes

- · Study subjects labeled as males with an excess of heterozygous SNPs\* on the X chromosome can
- denote
- That their genetic sex is female
- Klinefelter syndrome
- Note: Individuals who are XY will also be heterozygous for markers in the pseudoautosomal regions
- Availability of Y chromosome data
- Can greatly aid in determining genetic sex and if an individual has Turner or Klinefelter syndrome \*Both genetic males and females have two alleles for each locus on the X chromosome in the datafile, although genetic males are hemizygous

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### Data Clean – Assessing Sex Chromosomes

- Individuals whose labeled gender/sex does not match their genetic sex are removed from the analysis
- This observation may be due to a sample swap
- When samples are swapped Phenotype data will be incorrect
  - e.g., may be a case when labeled as a control

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### **Checking for Duplicate and Related Individuals**

- Duplicate samples are sometimes included in a study as part of quality control to detect inconsistencies - Will not detect systematic errors
- Usually not included in exome and whole genome sequencing studies - Intentional duplicates can easily be removed before data quality control
- Cryptic duplicates (unintentional)
- DNA sample aliquoted more than once
- Individual ascertained more than once for a study
- e.g. The same individual undergoes the same operation more than once and is ascertained each time
- · Individuals who are related to each other may participate in the same study
- Unknown to the investigator
- Or be part of the study design

### Duplicate and Related Individuals Need to be Identified

### For duplicate samples Only one can be retained

- For related individuals
- PCA is performed first with unrelated individuals and related individuals are then projected onto the PCs of unrelated individuals
- Mixed-models need to be used to analyze the data if related individuals are included\* Case-Control
- Generalized linear mixed models (GLMM)
- Quantitative traits
- Linear mixed models (LMM)
- · If related individuals are ignored in the analysis type I error rates can be inflated

"If only a few related individuals in sample, may wish to remove them or use LMM/GLMM to control type I errors. Must use LMM/GLMM if related individuals are included in the dataset. If possible, opt for LMM/GLMM since it can help to control type I error due to other types of structure in the data, even when no closely related individuals are included

### Identifying Duplicate and Related Individuals

- Duplicate and related individuals can be detected
   By examining <u>identity-by-State</u> (IBS) adjusted for allele frequencies (p-hat) between all pairs of individuals within a sample
- Identify-by-descent (IBD) sharing can be estimated

Identity by Descent (IBD)/Identity-by-State (IBS) П  $\cap$ 1/3 1/2 1/3 1/2 1/2 1/3 Ó 1/3 1/2 1/1 1/3 1/2 1/2 IBD=0 IBD=1 IBD=2 IBS=2 IBS=1 IBS=1

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### IBD Sharing Estimated Pairwise for all Individuals in a Samples

- PLINK (Purcell et al. 2007)
- Uses sequence (or genotype array) data to check IBD
   Prune markers to remove those in linkage disequilibrium
   (LD)
  - e.g., r<sup>2</sup><0.1</li>
- P-hat is calculated using the "population" allele frequency
  Used to approximates IBD sharing
- IBD is the number of alleles of alleles which are shared between
- a pair of individuals
   Can either share 0, 1, and 2 alleles

### Identifying Duplicate and Related Individuals

- Monozygote twins and duplicate samples will share 100% of their alleles IBD
- IBD=2 is 1.0 (can be lower due to genotyping error)
- Siblings and child-parent pairs will share 50% of their alleles IBD
- For parent-child IBD=1 is 1.0 (IBD=0 is 0 & IBD=2 is 0)
- For sibs IBD=1 is ~0.50 (IBD=0 is ~0.25 & IBD=2 is ~0.25)
   For more distantly related individuals the IBD measure will be lower

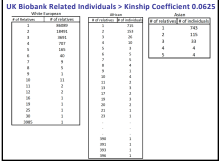
33

### Identifying Duplicate and Related Individuals

- KING [Kinship-based INference for Gwas (Manichaikul et al. 2010)] can also be used to identify
- duplicate and related individuals
- KING is more robust to population substructure and admixture
  - Prune markers for LD (e.g., r<sup>2</sup><0.1)
- Provides kinship coefficients
- Duplicate samples
- Kinship coefficient equals 0.5
- Siblings

   Kinship coefficient equals 0.25

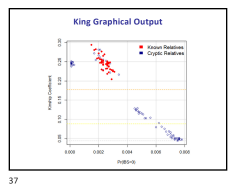




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36

32



### Multiple Individuals Observed That are Distantly "Related"

- If individuals in sample come from different populations e.g., individuals from the same population within the sample will have inflated p-hat values due to incorrect allele frequencies Incorrectly appear to be related to each other
- "Relatedness" amongst many individuals can also be observed when batches are combined if they have different error rates - Individuals from the same batch appear to be related
- DNA contamination can cause "relatedness" between multiple individuals

Principal Components Analysis (PCA) / Multidimensional Scaling (MDS)

- Can be used to identify outliers
- Population substructure
- Individuals from different ancestry e.g., African American samples included in samples of European
- Americans
- Batch effects
- Use a subset of markers which have been LD pruned - Only very low levels of LD between marker loci

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- e.g., r<sup>2</sup><0.1
- MAF cutoff dependent on sample size • e.g MAF> 0.01
- Can use lower MAF for large sample sizes

### Principal Components Analysis (PCA) / Multidimensional Scaling (MDS)

- Unrelated individuals are used to generate PC plots
- Related individuals are projected onto to the PC plots
- Plot 1<sup>st</sup> component vs. 2<sup>nd</sup> component - Additional PCs should also be plotted
- e.g.. PCs 1-10
- Mahalanobis distance can be used to determine outliers – e.g., <1

40

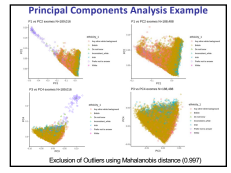
38

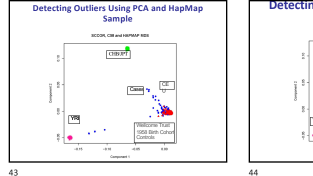
### PCA/MDS Can be Used to Identify Outliers

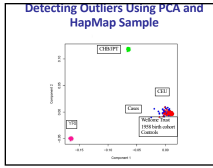
- Individuals of different ancestry
- e.g., African American samples included with European Americans samples
- Can use samples from HapMap/1000 genomes to help to determine the ancestry for samples that are outliers Should not include HapMap/1000 genomes samples when calculating components to control for population substructure/admixture

Batch effects

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### **Detecting Genotyping Error – Examining HWE**

- Testing for deviations from HWE not very powerful to detect genotyping errors
- The power to detect deviations from HWE dependent on: Error rates
- Underlying error model
- Random
- Heterozygous genotypes -> homozygous genotypes
- Homozygous genotypes ->Heterozygous genotype
- Minor allele frequencies (MAF)

### **Detecting Genotyping Error – Examining HWE**

- Controls and Cases are evaluated separately
- Deviation found only in cases can be due to an association Test for deviation from HWE only in samples of the same
- ancestry - Population substructure can introduce deviations from HWE
- Do not include related individuals when testing for

deviations from HWE

- Can cause deviations from HWE

45

46

### **Detecting Genotyping Error – Examining HWE**

- What criterion is used to remove variants due to a
- deviation from HWE
- GWAS studies have used 5.0 x  $10^{\text{--}7}$  to 5.0 x  $10^{\text{--}15}$
- Quantitative Traits
- Caution should be used removing markers which deviate from HWE may be due to an association Remove markers with extreme deviations from HWE and Flag markers
  - with less extreme deviations from HWE
- When performing imputation need to be more stringent in removing variants which deviate from HWE

### Sequence Data QC Overview

- Remove variant sites that fail VQSR
- Remove genotypes with low DP, GQ scores, etc.
- · Remove variant sites with large percent of missing data
- Remove samples with missing large percent of missing data
- Evaluate genetic sex of individuals based upon X and Y chromosomal data
- Sample mix-ups
- Individuals with Turner or Klinefelter Syndrome

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### Sequence Data QC Overview

### • Evaluate samples for cryptically related individuals and

- duplicates
- Use variants which have been pruned for LD • e.g., r²<0.1
- King or Plink algorithm
- Always remove duplicate individuals
- Retaining only one in the sample
- If sample includes related samples use linear mix models (LMM)/Generalized LMM (GLMM) to control for relatedness
- Best to perform even for data without related individuals • If only a few related individuals can retain only one individual of a
- relative group if not using LMM or GLMM
- Removal of outliers (can be determined by Mahalanobis distance) - Inclusion of MDS or PCA components in the association analysis

Sequence Data QC Overview

Use unrelated individuals and then project related individuals

• Due to population substructure/admixture and batch effects

- Perform principal components analysis (PCA) or multidimensional scaling (MDS) to detect outliers

Detection of sample outliers

– e.g., r²<0.1

onto the PCs

· Remove effects by

Additional QC

and\or

50

52

Use variants pruned for LD

49

51

53

### Sequence Data QC Overview Remove/flag variant sites that deviate from HWE in controls

- HWE should be only be tested in unrelated individuals from the same population
- Post Analysis Quantile-Quantile (QQ) plots - To evaluate uncontrolled batch effects and population
  - substructure/admixture

### QQ Plots - Genome Wide Association Diagnosis

- · Thousands of variants/genes are tested simultaneously · The p-values of neutral markers follow the uniform distribution
- If there are systematic biases, e.g., population substructure, genotyping errors, there will be a deviation from the uniform distribution
- QQ plots offers an intuitive way to visually detect biases
- Observed p-values are ordered from largest to smallest and their -log<sub>10</sub>(p) values are plotted on the y axis and the expected -log<sub>10</sub>(p) values under the null (uniform distribution) on the x axis

QQ Plot of Exome Wide P Values

UK Biobank 200K

λ = 1.046

Problem hearing

Cases N=65 660

Controls N= 96,601

with background noise

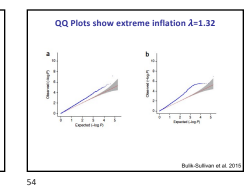


λ = 0.942

Hearing aid users

Controls N= 96,601

Case N= 6.436

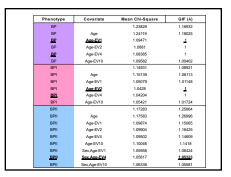


### Genomic Inflation Factor to Evaluate Inflation of the Test Statistic

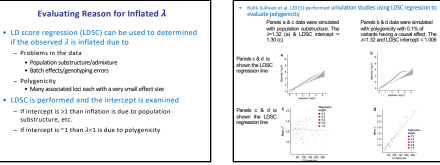
- Genomic Inflation Factor (GIF): ratio of the median of the test statistics to expected median and is usually represented as  $\lambda$
- No inflation of the test statistic  $\lambda\text{=}1$
- Inflation λ>1
- Deflation λ<1</li>
- Can be observed when a study is underpowered
- Problematic to examine the mean of the test statistic - Can be large if many variants are associated Particularly if they have very small p-values

Should not be used



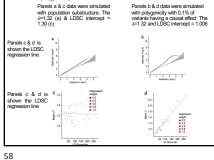


56





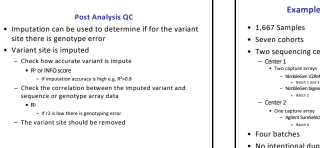
59



Post Analysis QC Observe in Manhattan plots individual associated variants with no surrounding associated variants

### Post Analysis QC

- · Most variants are in LD with neighboring SNPs
- · Genotyping error can cause a variant site not to be in LD with any of its neighbors
- · Genotyping error can also cause a spurious associations
- A lone associated variant site can be due to genotyping error



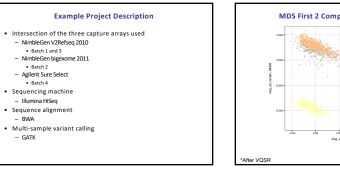


• R<sup>2</sup>

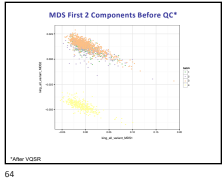


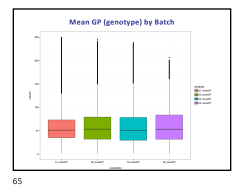


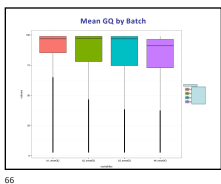


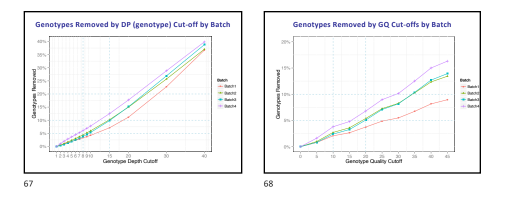


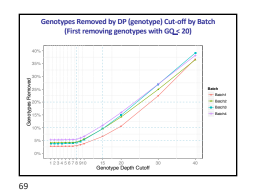


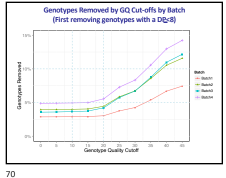


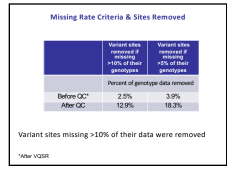


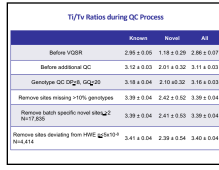


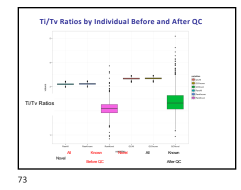


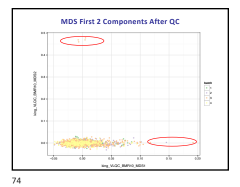












Sequence Data QC

- Batch effects can sometimes be removed with additional QC
- Extreme outliers should be removed
- Additionally, MDS\PCA components can be included in the analysis to control for population substructure\admixture and batch effects

Unless correlated with the outcome (phenotype)

- The MDS or PCA components should be recalculated after QC only including those samples included in the analysis
- Batch (dummy coding) may be included as a covariate in the analysis
  - Unless correlated with the outcome (phenotype)

75

## Convenience Controls Can reduce the cost of a study

- Genotype data
- Type I error can be increased
- Ascertainment from different population
   Differential genotyping error
- Even if performed at the same facility
- Proper QC can reduce or remove biases

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### Convenience Controls–Sequence Data

- Obtain BAM files and recall cases and control together
   Can still have differential errors between cases and controls
- Check variant frequency by variant types in cases and control
- Synonymous variants should have the same frequencies
   Would not expect large differences in numbers of variants between cases and controls
- For single variants can compare difference in frequencies with gnomAD but is problematic
- Differences in frequencies can be due to differences in ancestry and/or sequencing errors
- Cannot adjust for confounders
- e.g., sex, population substructure/admixture
- Don't perform an aggregate test using frequency information obtained from databases, e.g., gnomAD, TOPMed Bravo

### Genotype Array Data

- Genotype Data QC Population Based Studies Initially remove DNA samples from individuals who are missing
- Initially remove DNA samples from individuals who are missing >10% or their genotype data
- For variant sites with a minor allele frequency (MAF)>0.05

   Remove variants sites missing >5% of their genotype data
- For variant sites with a MAF<5%</li>
   Remove variant sites missing > 1% of their genotype data
- The genotypes for variant sites with missing data may have higher genotype error rates



### Order of Data Cleaning-Genotype Array Data

- Remove samples missing >10% genotype data
- Remove SNPs with missing genotype data

   If minor allele frequency >5%
   Remove markers with >5% missing genotypes
- If minor allele frequency <5%
   Remove markers with >1% missing genotypes
- Remove samples missing >3% genotype calls
- Check genetic sex of individuals based on X-chromosome markers & Y chromosome marker data (if available)
- Remove individual whose reported gender/sex is inconsistent with genetic data
- Could be due to a sample mix-up
- Check for cryptic duplicates and related individuals
   Used "trimmed data set of markers which are not in LD
  - e.g. r2<0.1</li>

Remove duplicate samples

### 79

### Order of Data Cleaning-Genotype Array

- Perform PCA or MDS to check for outliers
- Use trimmed data set of markers which are not in LD
   e.g., r2<0.1</li>
- First with unrelated individuals and then project related individuals on the components
- Remove outliers from data
   e.g., Mahalanobis distance
- Check for deviations from HWE
- Separately in cases and controls
- Only unrelated individuals
- If more than one ancestry group
   Separately for each ancestry group

   As determined via PCA or MDS

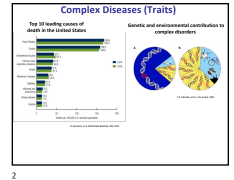
80

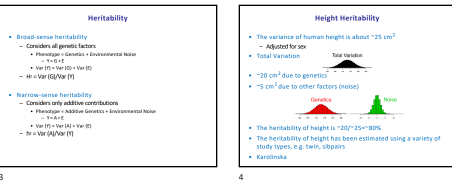
### Order of Data Cleaning-Genotype Array

- Examine QQ plots
- e.g., not controlling adequately for population admixture
   Inflated test statistics Deflated p-values
- Examine Manhattan to detect associated variants which are not in LD with other variants
- Genotyping errors causing spurious associations

## Complex Trait Association Analysis of Rare Variants Obtained from Sequence Data

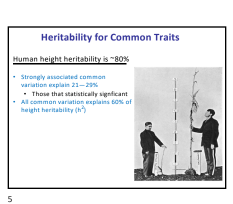
Suzanne M. Leal, Ph.D. Sergievsky Family Professor of Neurological Sciences Director of the Center for Statistical Genetics Columbia University sml3@Columbia.edu

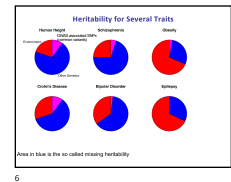


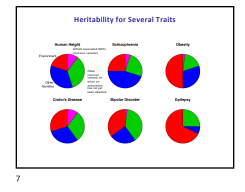


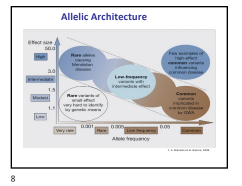
© 2024 Suzanne M. Leal

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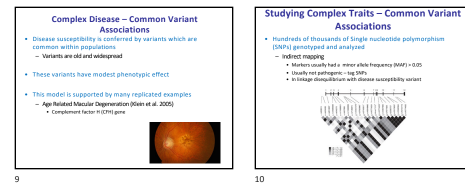


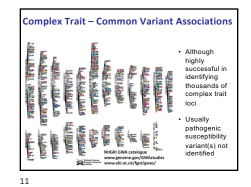


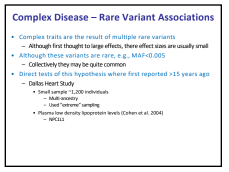


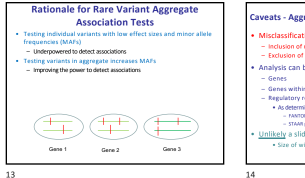


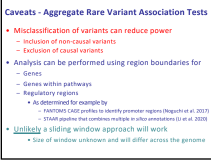
Associations

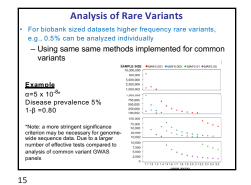


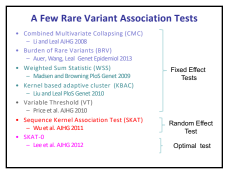


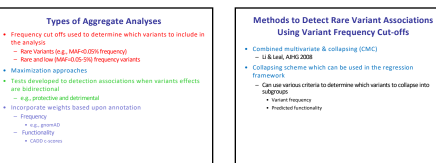












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the analysis

Maximization approaches

e.g., protective and detrimental

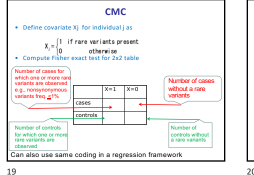
are bidirectional

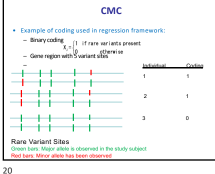
- Frequency • e.g., gnomAD Functionality CADD c-scores

Rare Variants (e.g., MAF<0.05% frequency)</li>

- Rare and low (MAF=0.05-5%) frequency variants

Incorporate weights based upon annotation





GRANVIL

-1

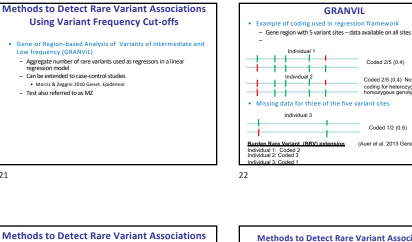
Coded 2/5 (0.4)

Coded 2/5 (0.4) Note same

coding for heterozygous and homozygous genotypes

(Auer et al. 2013 Genet Epidemiol)

Coded 1/2 (0.5)



### Weighted Approaches

- Group-wise association test for rare variants using the Weighted Sum Statistic (WSS)
- Variants are weighted inversely by their frequency in controls (rare variants are up-weighted)
- Madsen & Browning, PLoS Genet 2009 Kernel based adaptive cluster (KBAC)
- Adaptive weighting based on multilocus genotype
   Liu & Leal, PLoS Genet 2010

### Methods to Detect Rare Variant Associations **Maximization Approaches**

### • Variable Threshold (VT) method

- Uses variable allele frequency thresholds and maximizes the test statistic - Can also incorporate weighting based on functional information
- Price et al. AJHG 2010 RareCover
- Maximizes the test statistic over all variants with a region using a greedy heuristic algorithm

Bhatia et al. 2010 PLoS Computational Biology

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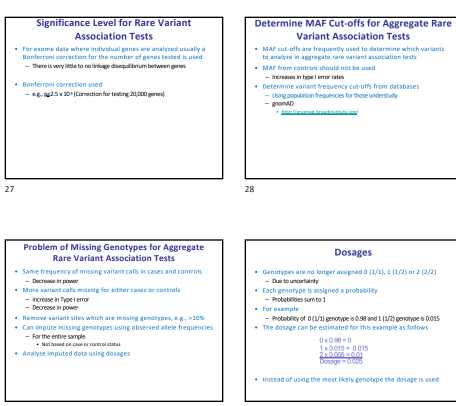
### **Optimal Test**

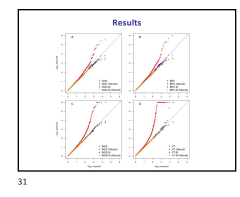
SKAT-O

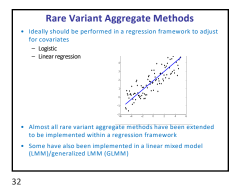
26

 Maximizes power by adaptively using the data to combine a burden test and the sequence kernel association tests
 Lee et al. 2012 AIHG

25





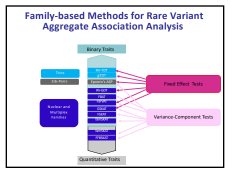




- For data that deviates from normality

   Quantile-quantile normalization
- For data that includes outliers
   Winsorize
- Don't winsorize and then normalize
- Instead of analyzing quantitative trait values
   Residual can be generated
   Adjusting for confounders

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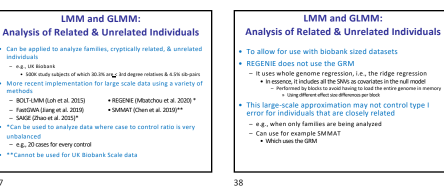


inear Mixed Model (LMM) & generalized LMM (GLMM) Analysis of Related & Unrelated Individuals

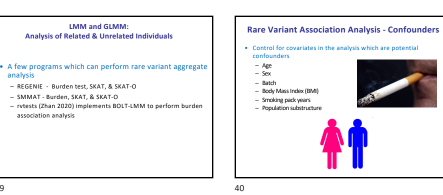
- LMM is an extension of the linear model to allow for both fixed & random effects and also allows for nonindependence of samples
- Early implementations calculated the kinship matrix  $\Phi$  on the basis of known relationships
- Amin et al. (2007) proposed to estimate kinships based on genome-wide variant data

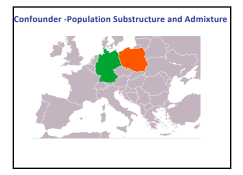
 The generalized relationship matrix (GRM) can be estimated for all individuals using for example identical-by-descent (IBD) sharing

• Extended to binary (case-control) traits - GLMM









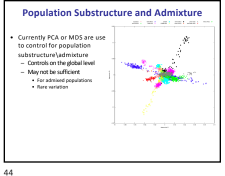


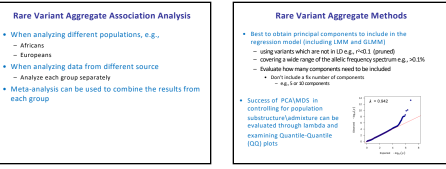
### **Population Substructure and Admixture**

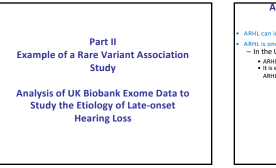
- If proportion of cases and controls sampled from each population is different
- Can occur due to
- Disease frequency is different between populations
  Sloppy sampling
- Population substructure\admixture can cause detection of differences in variant frequencies within a gene which is due to sampling and not disease status

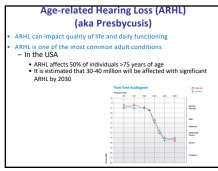
   False positive findings can be increased

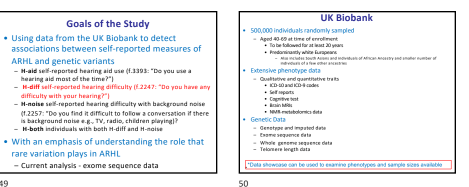




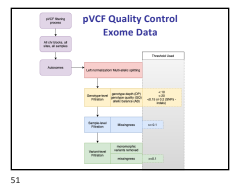


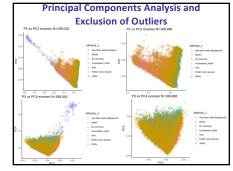












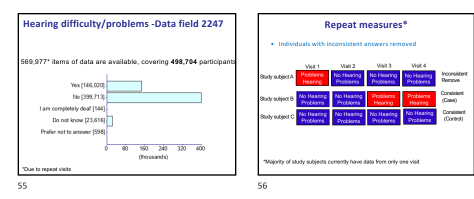
### Exclusion Criteria Obtained from ICD10, ICD9, & Self Report • Deafness • Early-onset hearing impairment • Otosclerosis • Meniere's • Labyrinthitis • Disorders of acoustic nerve • Bell's palsy • History of chronic suppurative and nonsuppurative otitis media • Meningitis

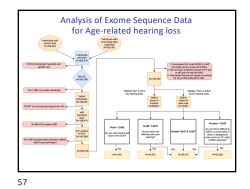
- Encephalitis, myelitis, and encephalomyelitis
- Etc.

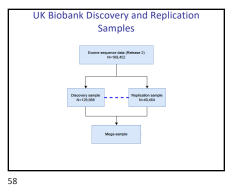
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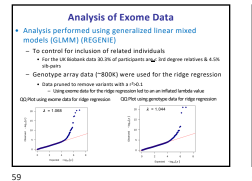
### **Defining Cases and Controls**

- Based on answers obtained from a touch screen
- Cases self-reported hearing difficulty
- f.2247: "Do you have any difficulty with your hearing?"
- Controls did <u>not</u> have any self-reported HL or ID10/9 HL codes









### Analysis of Exome Data

- Analysis limited to individuals of White European
   Ancestry
- Sex, age, and two PCAs included as covariates
- Age
- cases first report of hearing difficulty
  Controls age at last visit
- The PCAs where recalculated for only individuals included in
- the analysis
- Using linkage disequilibrium (LD) pruned genotypes array data (r2<0.1)</li>

### Analysis of Exome data – Single Variant

## • Variants with four or more alternative alleles observed in the sample analyzed

 A very low MAF was used since it was hypothesized some of the variants may have a large effect sizes

### Significance Levels-Single Variant Analysis

### • Discovery sample

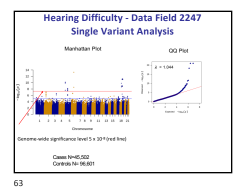
 A genome-wide significance level (single variant analysis) was used to reject the null hypothesis of no association
 pg5.0x10<sup>8</sup>

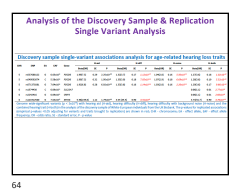
### Replication sample

 Permutation was used to obtain empirical p-values
 Adjusting for the phenotypes and variants brought to replication
 ps0.05

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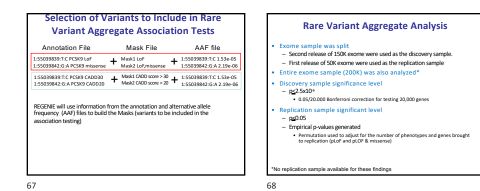
# Rare Variant Aggregate Analysis Genes with at least two variants were analyzed, e.g., pLoF variants

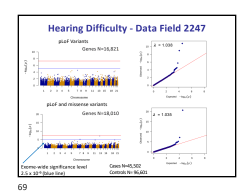
- Max coding was used
- Two masks were used
- Mask 1 pLoF variants
- Mask 2 pLoF and missense variants

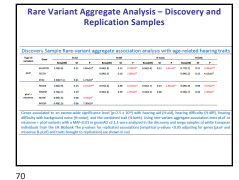
### Minor allele frequency cut-off of <0.01 was used</li>

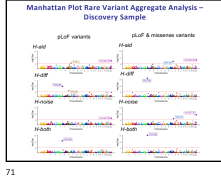
 The frequencies for each variant site were obtained from gnomAD (non-Finnish Europeans)

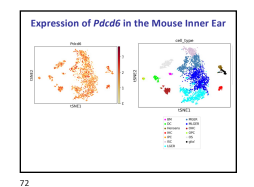
### **REGENIE Rare Variant Aggregate Analysis** Three different codes can be used • Max • Sum Comphet · This term is not correct because the phase is unknown · Variants may be on the same haplotype Note: At the time of the study REGENIE could only perform fixed effect tests it now implements SKAT and SKAT-O Single variant sites max comphet sum 0000000000000 → 0 0 0 $00000100010000 \rightarrow$ 1 2 2 00201011010100 $\rightarrow$ 2 7 2













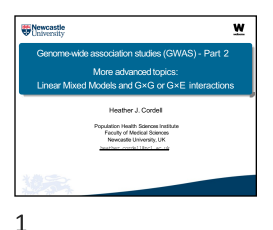
### Overview

- Replicated some previously reported ARHL genes
- Some which had not been previously replicated
- e.g., BAIAP2L2, CRIP3, KLHDC7B, MAST2, and SLC22A7 Identified and replicated a new HL gene, PDCD6 which has not been
- previously reported
- Inner ear expression in humans and mice supports the involvement of gene in HL etiology
- PDCD6 is a cytoplasmic Ca2+ binding protein with an important role in apoptotic cell death
- Rare-variant aggregate analysis demonstrated the important contribution of Mendelian HL genes, i.e. MYO6, TECTA, and EYA4 the genetics of ARHL
- Rare variants for ARHL tend to have larger effect sizes than those for common variants
- Rare variants should play an important role in risk prediction by increasing accuracy
- For additional information see Cornejo-Sanchez et al. (2023) Eur J Hum Genet PMID: 36788145

# **Overview/Future Direction**

- The entire exome sequence data set of White Europeans has been analyzed
- Reveling many additional known Mendelian nonsyndromic HL genes Mendelian genes (although not necessarily the same variants)
- play an important role in ARHL Performing Mendelian Randomization and testing for pleiotropy
- (vertical & horizontal) to evaluate associations between ARHL and comorbidities - e.g., dementia, depression
- Analysis of UK Biobank and All of Us WGS data including structural variants and performing rare variant aggregate tests outside of the coding regions

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Linear Mixed Models (LMMs) Linear Mixed Models have been used for many years in the plant and animal breeding communities In the mid 1990s they became popular in the human genetics field, mostly for performing linkage analysis and estimating heritability Using family (pedigree) data i.e. related individuals 2

# Linear Mixed Models (LMMs)

- Linear Mixed Models have been used for many years in the plant and animal breeding communities
- In the mid 1990s they became popular in the human genetics field, mostly for performing linkage analysis and estimating heritability Using family (pedigree) data i.e. related individuals
- In recent years they have become popular in the genetic association studies field for:
  - Testing for association while accounting for varying degrees of relatedness
    - Releances
       Cose family relationships
       Distant relationships and population stratification/substructure

# 3

# Linear Mixed Models (LMMs)

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  Distant relationships and population stratification/substruct Estimating the heritability accounted for various partitions of SNPs: All SNPs typed on a GWASpanel
   All SNPs typed on a GWASpanel
   All typed SNPs and others in LD with them
   Partitions of SNPs in various functional categories
  - Investigating genetic correlations between different traits

5



Linear Mixed Models (LMMs)

Linear Mixed Models have been used for many years in the plant and

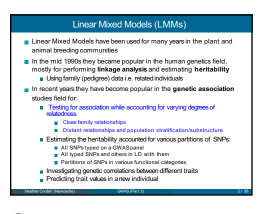
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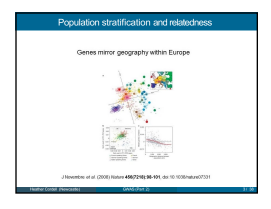
In recent years they have become popular in the genetic association

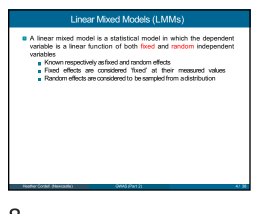
animal breeding communities

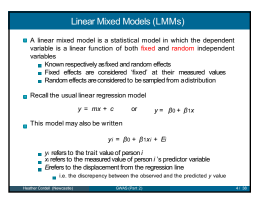
studies field for:

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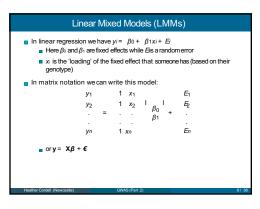


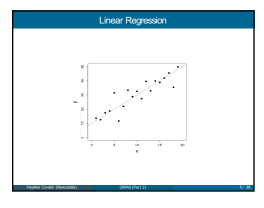




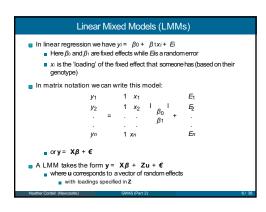






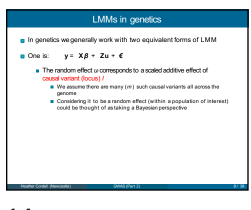




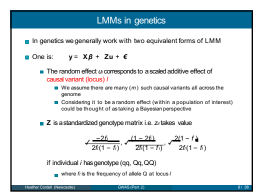


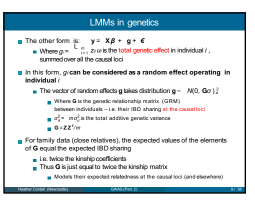


	Linear Mixed Models (LMMs)								
random en	<ul> <li>E.g. suppose 2 fixed effects β1 and β2, and 3 random effects (plus n random errors)</li> <li>Then y = Xβ + Zu + € corresponds to:</li> </ul>								
	X11	$\begin{vmatrix} & & \\ & & & \\ & & & & \\ & & & & & \\ & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & & \\ & & & & & & \\ & & & & & & & \\ & & & & & & \\ & & & & & & $	z <sub>11</sub> z <sub>21</sub>	z <sub>22</sub>	z <sub>23</sub>	и1 и <sub>2</sub> и3	+	El E2 En	
or $y_1 = \beta_1 x_{11} + \beta_2 x_{12} + u_1 z_{11} + u_2 z_{12} + u_3 z_{13} + E$									
Heather Cordell (Newcastle) GWAS (Part 2) 71 3									



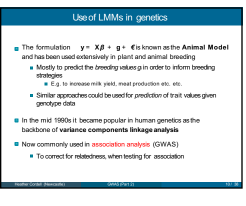
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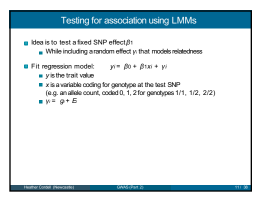






# LMMs in genetics The other form is: y = Xβ + g + € Where g<sub>i</sub> = L <sub>μ</sub>, zw is the total genetic effect in individual i, summed over all the causal loci In this form, g<sub>i</sub> can be considered as a random effect operating in individual i The vector of random effects g takes distribution g ~ N(0, Gσ)<sub>a</sub><sup>2</sup> Where G is the genetic relationship matrix (GRM) between individuals – i.e. their IBD sharing at the causal loci σ<sub>a</sub><sup>2</sup> = mσ<sub>a</sub><sup>2</sup> is the total additive genetic variance G = zz<sup>1</sup>/m





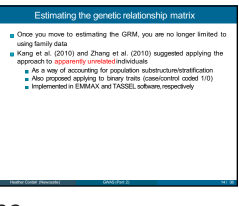
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# Testing for association using LMMs

- LMMs were first (?) applied in human genetics by Boerwinkle et al. (1986) and Abney et al. (2002)
- Chen and Abecasis (2007) implemented them via the "FAmily based Score Test Approximation" (FASTA) in the MERLIN software package
  - Closely related to earlier QTDT method (Abecasis et al. 2000a;b) which implements a slightly more general/complex model
  - FASTA was also implemented in GenABEL, along with a similar test called GRAMMAR (Aulchenko et al. 2007)

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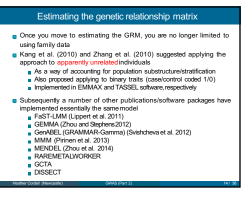


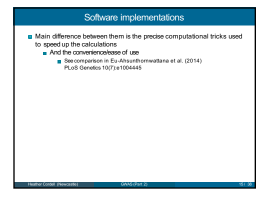


# Estimating the genetic relationship matrix

- These early implementations calculated the kinship matrix Φ on the basis of known (theoretical) kinships constructed from known pedigree relationships
- Amin et al. (2007) proposed instead estimating the kinships based on genome-wide SNP data
  - Ideally we want to use G=ZZ//m, the genetic relationship matrix (GRM) between individuals at the causal loci
  - Since we don't know the causal loci, we approximate G by A, the overall GRM between individuals
     Various different ways to estimate this, usually based on scaled (by aliele frequency) matrix of *identity-by-state* (IBS) sharing

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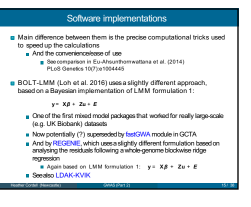
# Binary traits

- For binary traits, coding cases and controls as a 1/0 quantitative trait is not optimal
- Though in practice it seems to work reasonably well LTMLM (Hayeck et al. 2015) and LEAP (Weissbrod et al. 2015)
- instead use an underlying liability model to improve power Assuming known disease prevalence

# 27

# Binary traits SAIGE software (Zhou et al. 2018, AJHG 50(9):1335-1341) implements a mixed model test that deals with large case-control imbalance, as you might see (for example) in UK Biobank REGENIE also implements this same saddle point approximation (SPA) test Along with an approximate Firth penalized likelihood-ratio test See also LDAK-KVIK

# 29



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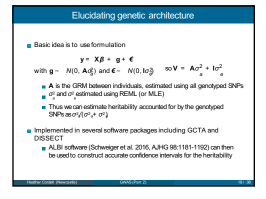
# Binary traits

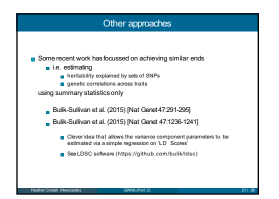
- For binary traits, coding cases and controls as a 1/0 quantitative trait is not optimal
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- LTMLM (Hayeck et al. 2015) and LEAP (Weissbrod et al. 2015) instead use an underlying liability model to improve powe Assuming known disease prevalence
- Chen et al. (2016) showed that high levels of population stratification can invalidate the analysis, when applied to a case/control sample
   Resulting in a mixture of inflated and deflated test statistics
  - Developed GMMAT software to address this problem See also CARAT software (Jiang et al. 2016, AJHG 98:243-55)

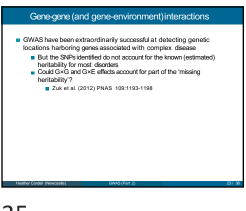
# 28

# Elucidating genetic architecture

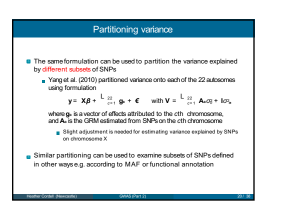
- Seminal paper by Yang et al. (2010) [Nat Genet 42(7):565-9]
- Showed that by framing the relationship between height and genetic factors as an LMM, 45% of variance could be explained by considering
- addition as an Linkin, 40% of Variable Could be explained by considering 294,831 SNPs simultaneously
   So-called 'SNP heritability' or 'chip heritability'
   Demonstrated that modelling effects at all genotyped SNPs explained the 'known' heritability (~ 80%) much better than just the top SNPs from GWAS
- Moreover, if you estimate effects of additional SNPs in LD with the genotyped SNPS, the variance explained goes up to 84% (s.e. 16%), consistent with 'known' value
- Subsequently many papers have shown similar results for a variety of complex traits

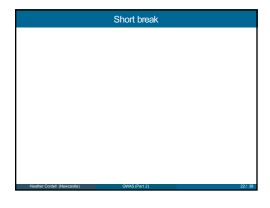


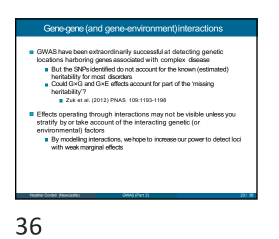


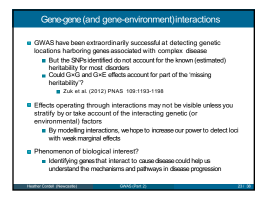


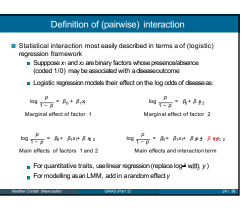


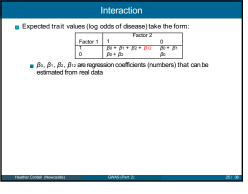




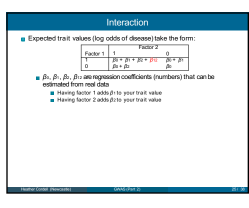




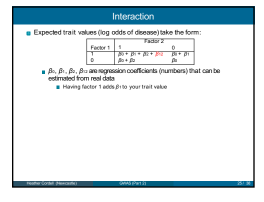




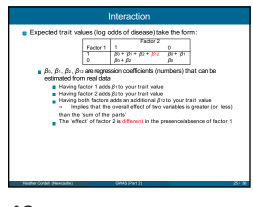






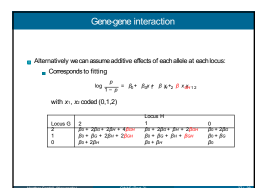


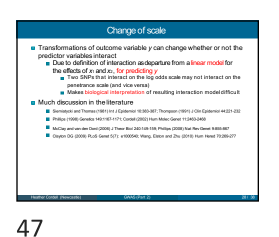


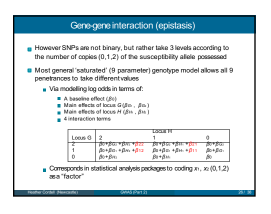


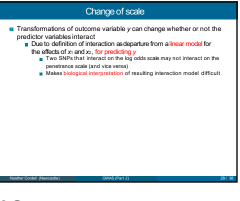


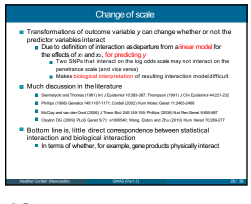
Interaction						
Expected trait values (log odds of disease) take the form:						
Fa 1 0	$\begin{array}{c} \text{Factor 2} \\ \hline & 0 \\ \beta_0 + \beta_1 + \beta_2 + \beta_{12} & \beta_0 + \beta_1 \\ \beta_0 + \beta_2 & \beta_0 \end{array}$					
β₀, β₁, β₂, β₁₂ are regression coefficients (numbers) that can be estimated from real data Having factor 1 adds β₁ to your trait value						
<ul> <li>Having factor 2 adds 21c your trait value</li> <li>Having both factors adds an additional 3/21c your trait value</li> <li>Implies that the overall effect of two variables is greater (or less)</li> <li>than the 'sum of the parts'</li> <li>The 'effect' of factor 2 is different in the presence/absence of factor 1</li> </ul>						
Suppose no main effects ( $\beta_1 = \beta_2 = 0$ )						
	Factor 1         1         0           1         β0 + β12         β0           0         β0 + β0         β0					
Trait value only differs from baseline if both factors present						
Heather Cordell (Newcastle)	GWAS (Part 2) 25/ 38					







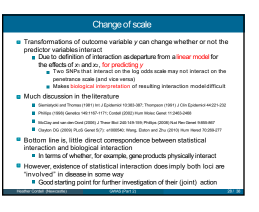




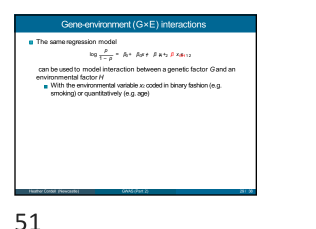
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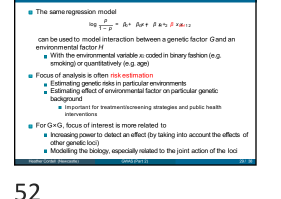
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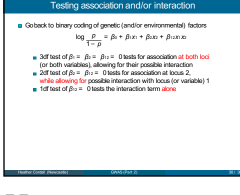


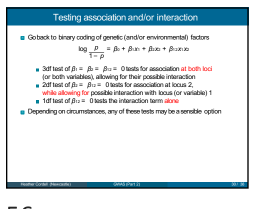


Gene-environment (G×E) interactions

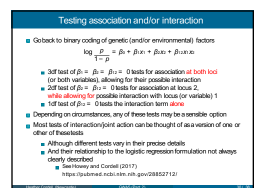
**Testing association and/or interaction** • Goback to binary coding of genetic (and/or environmental) factors  $\log \frac{P}{1-p} = \beta_0 + \beta_{1X1} + \beta_{2X2} + \beta_{12X1X2}$ • 3df test of  $\beta_1 = \beta_2 = \beta_1 = 0$  tests for association at both loci (or both variables), allowing for their possible interaction • 2df test of  $\beta_2 = \beta_1 = 0$  tests for association at locus 2, while allowing for possible interaction with locus (or variable) 1



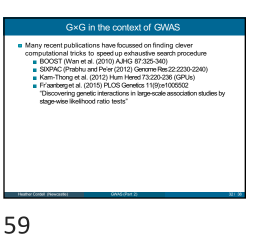




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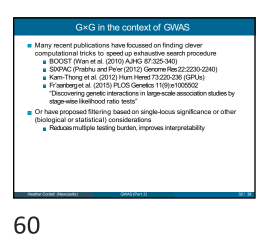


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# G×G versus G×E in the context of GWAS

- Typically GWAS measure thousands if not millions of genetic variants But only a few (tens or at most 100s) of environmental factors
- Feasible to consider all G×E combinations
- All pairwise G×G combinations possible, but much more time consuming
- And leads to greater multiplicity of tests Also, why stop at 2-way interactions? Could look at all 3 way, 4 way etc. combinations Scale of problem quickly gets out of hand Less obvious reason to do this for G×E...



# G×G in the context of GWAS

Many recent publications have focussed on finding clever BOOST (Wan et al. (2010) AJHG 87:325-340)
 SIXPAC (Prabhu and Pe'er (2012) Genome Res 22:2230-2240)

- Kam-Thong et al. (2012) Hum Hered 73:220-236 (GPUs) Fraanberg et al. (2015) PLOS Genetics 11(9):e1005502 "Discovering genetic interactions in large-scale association studies by stage-wise likelihood ratio tests"
- Or have proposed filtering based on single-locus significance or other (biological or statistical) considerations Reduces multiple testing burden, improves interpretability
- Or have proposed testing at the gene level rather than the SNP level
  - Ma et al. (2013) PLoS Genet 9(2): e1003321
     Compared 4 different tests that combine P values from pairwise

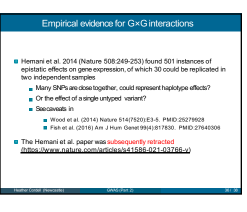
    - (SNP x SNP) interaction tests Showed that the truncated tests did best
    - Presented an application only considering gene pairs known to exhibit protein-protein interactions

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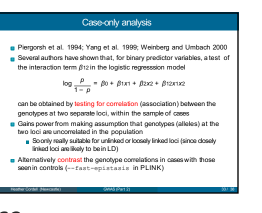
# Testing correlation between loci

- A similar idea is implemented in EPIBLASTER
- (Kam-Thong et al. 2011; EJHG 19:465-571)
- Wu et al. (2010) (PLoS Genet 6:e1001131) also proposed a similar approach though needs adjustment to give correct type I error rates
- See also Joint Effects (JE) statistics (Ueki and Cordell 2012; PLoS Genetics 8(4):e1002625)
- All these methods test whether correlation exists (case-only) or is different in cases and controls (case/control)
  - Via testing a log OR for association between two loci However, the log OR for association (λ) encapsulates a slightly different quantity between the different methods
- All implemented (along with standard logistic and linear regression) in CASSI
  - http://www.staff.ncl.ac.uk/richard.howev/cassi/

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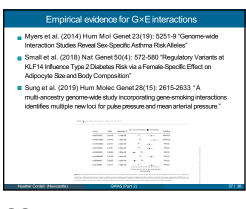


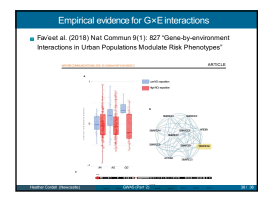
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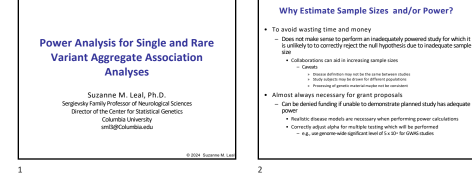
# Empirical evidence for G×G interactions

- Epistasis among HLA-DRB1, HLA-DQA1, and HLA-DQB1 in multiple sclerosis (Lincoln et al. 2009 PNAS 106:7542-7547)
- HLA-C and ERAP1 in psoriasis (Strange et al. 2010)
- HLA-B27 and ERAP1 in ankylosing spondylitis (Evans et al. 2011)
- BANK1 and BLK in SLE (Castillejo-Lopez et al. 2012)
- Gusareva et al. (2014) found a reasonably convincing (partially replicating) interaction between SNPs on chromosome 6 (KHDRBS2) and 13 (CRYL1) in Alzheimer's disease
- Dai et al. (2016) [AJHG 99:352-365] identified 3 loci simultaneously interacting with established risk factors gastresophageal reflux, obesity and tobacco smoking, with respect to risk for Barrett's esophagus

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### Power and Sample Size Estimation for Case-Control Data

- The correct α must be use for sample size estimation/power analysis
- Type I (a) the probability of rejecting the null hypothesis of no association when it is true
- Due to multiple testing a more stringent value than  $\alpha$ =0.05 is used in order to control the Family Wise Error Rate

### Power and Sample Size Estimation for Case-Control Data

- GWAS of common variants where each variant is test separately
   – α=5 X10<sup>s</sup> (Bonferroni Correction for testing 1,000,000 variant sites)
- Shown to be a good approximation for the effective number of tests
   Valid even when more than 1,000,000 variant sites tested
- Effective number of tests is dependent of the linkage disequilibrium (LD) structure
- Single variant tests using whole genome sequence data
   Many more rare variants than common variants
- Lower levels of LD between rare variants than between common variants
   The number of effective tests for rare variants is higher than for analysis

3

4

# An Example of Determining Genome-wide Significance Levels for Common Variants

- Using genotypes from the Wellcome Trust Case-Control Consortium
- Dudbridge and Gusnato, Genet Epidemiol 2008
- Estimated a genome-wide significance threshold for the UK European population
- By sub-sampling genotypes at increasing densities and using permutation to estimate the nominal p-value for a 5% familywise error
- Then extrapolating to infinite density
- The genome wide significance threshold estimate ~7.2X10<sup>-8</sup>
- Estimate is based on LD structure for Europeans
  - Not sufficiently stringent for populations of African Ancestry

### Power and Sample Size Estimation for Aggregate Rare Variant Tests

- For gene-based rare variant aggregate methods a Bonferroni correction for the number of genes/regions tested is used
- $-\,$  e.g., 20,000 genes significance level  $\,\alpha\text{=}2.5\,x\,10^{\,6}$
- Can use a less stringent criteria

   Not all genes have two or more variants
- » Divide 0.05 by number of genes
   If units other than genes are used
- If units other than genes are used
   A more stringent criteria may be necessary
- For rare variants very low levels of LD between variants in
- separate genes
- Therefore, a Bonferroni correction is not overly stringent
   The number of tests ≅ effective number tests
  - This would not be the case for variants in LD

### Power and Sample Size Estimation for Replication Studies

- For replication studies can base the significance level ( $\boldsymbol{\alpha})$
- · On the number of genes/variants being brought from the
- discovery (stage I) study • To replication (stage II)

7

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For example, if it is hypothesized that 20 genes and 80 independent variants will be brought to stage II (replication)

 A Bonferroni correct can be made for performing 100 tests
 An a = 5.0 x 10<sup>2</sup> and be used for a family wise error rate of 0.05

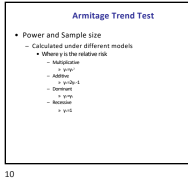
### Estimating Power/Sample Sizes For Single Variant Tests

- Can be obtained analytically
- Information necessary
- Prevalence
- Risk allele frequency
- Effect size (odds ratio-for case control data)
   Genetic model for the susceptibility variant
- Recessive (γ1=1)
- Dominant (γ2=γ1)
  Additive (γ2=2γ1-1)
- Multiplicative (y2=y1<sup>2</sup>)

# Estimating Power/Sample Sizes For Individual Variants

- Usually, information on disease prevalence is known from
- epidemiological data
- A range of risk allele allele frequencies and effect sizes are used
  A variety of genetic models can also used

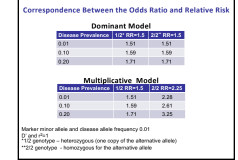
Dominant
 Additive
 Multiplicative



8

# Gamma is the Relative Risk not the Odd Ratio

- Most software for power calculations/sample size estimation use the relative risk  $(\gamma)$  and not the odds ratio
- The relative risk only approximates the odds ratio when disease is rare (Prevalence  ${\sim}<0.1\%)$
- The relative risk is not appropriate for common traits when a case-control design is used



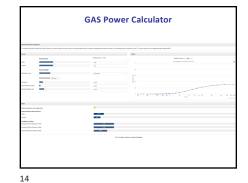
11

# Genetic Association Study (GAS) Power Calculator

- <u>http://csg.sph.umich.edu/abecasis/cats/gas\_power\_calculator</u>/i ndex.html
- A one-stage study power calculator
   Which was derived from CaTs
- Which is to perform two-stage genome wide association studies
   Skol et al. 2006
   Cochran Armitage Trend Test
- Displays graphs of the results
- · Displays graphs of the result

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**Genetic Power Calculator** 

User-defined type I error rate : 0.00000005 (0.00000001 - 0.5) User-defined power: determine N : 0.80 (0 - 1) (1 - type II error rate)

: 10000 (0 - 10000000) : 1 ( >0 ) ( 1 = equal number of cases and controls)

Unselected controls? (\* see below)

D-prime : 1 (0 - 1) Narker allele frequency (B) : 0.01 (0 - 1)

Case - control for discrete traits

Number of cases Control : case ratio

Created by Shaun Purcell 24.Oct.2008

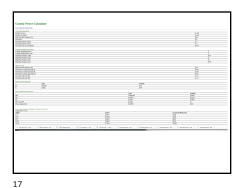
Process Reset

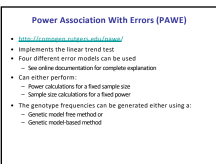


# Genetic Power Calculator

- <u>http://zzz.bwh.harvard.edu/gpc</u>/
- S Purcell & P Sham
- Uses the methods described in Sham PC et al. (2000) Am J Hum Genet 66:1616-1630
- VC QTL linkage for sibships
- VC QTL association for sibships
- VC QTL linkage for sibships conditional on the trait
- TDT for discrete traits
- Case-Control for discrete traits
- TDT for quantitative traits
   Case-Control quantitative traits
- Although input is the relative risk
- Displays odds ratios









### Quanto

- · Provides sample size and power calculations for
- Genetic and environmental main effects
- Interactions
   Gene x gene
- Gene x gene
  Gene x environment
- Sample & power calculations can be carried for:
   Case-control
  - Unmatched
- Matched
   Case-sibling
- Case-parent (trios)
- Quantitative
   Qualitative
- Independent sample of individuals
- Quantitative traits
   Assumption sampled from a random population
- Can only be run under windows
- https://pphs.usc.edu/download-quanto/
- 19

# Linkage Disequilibrium (LD)

- Power will be reduced if causal variant is not in perfect LD ( $r^{2}=1$ ) with the tag SNP
- Can adjust sample size when r<sup>2</sup><1 to increase power to the same level as when r<sup>2</sup>=1
- Can estimate sample size when  $r^2 \! \neq \! 1$
- N/r<sup>2</sup>=N'
   Valid only for multiplicative model
- (Pritchard and Przeworski, 2001)
- Power calculation almost always assume that r<sup>2</sup>=1
- For whole genome sequence data this should be the case since usually the causal variant would be included in the data

### Power Analysis for Rare Variant Aggregate Association Tests

- Many unknown parameters must be modeled
- Allelic architecture within a genetic region
   Varied across genes and populations
- Effects of variants within a region
   Fixed or varied effect sizes of causal variants
- Bidirectional effect of variants
- Proportion of non-causal variants
- Power estimated empirically
- Simplified assumptions can be made to obtain analytical estimates
- All variants have the same effect size
- No non-causal variants within a region that is analyzed in aggregate

# 21

## Simplistic Analytical Power Calculation for Rarevariant Aggregate Association Analysis

Assumption

20

- All rare variants are causal and have the same effect sizeAlthough usual not be correct
- Provides a gestalt of the power for a given samples or sample size for a given power
- Use aggregate of allele frequencies
- For example, assume a cumulative allele frequency of 0.025
   Use an exome-wide significant level e.g., 2.5x10<sup>-6</sup>
- Provide disease prevalence and penetrance model
- Perform calculations in the same manner as was described for single variants

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### **Empirical Power Calculations**

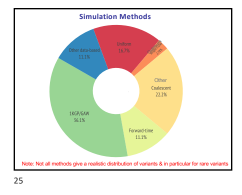
- A variety of methods can be used to generate variant data to empirically estimate power
- · Variant data is generated
- Based upon a penetrance model samples of cases and controls are generated
- Or a quantitative trait is generated based upon the genetic variance
- Multiple replicates are generated and analyzed
- To determine the power

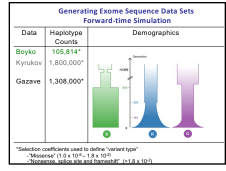
# **Empirical Power Calculations**

### Examples

- 5,000 replicates are generated each with 20,000 cases and 20,000 controls
- The power is the proportion of replicates with p-value less than the specified threshold, e.g.,  $5x10\,^{\rm s}$
- For rare-variant aggregate tests all autosomal genes are generated and those genes with more than two rare variants (e.g., predicted loss of function) are analyzed
- The power is the proportion of genes that were tested with p-value which is below a specified threshold, e.g.,  $2.5 \times 10^{\,6}$

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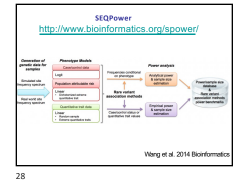


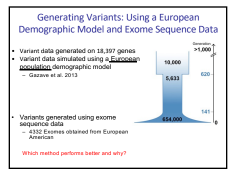


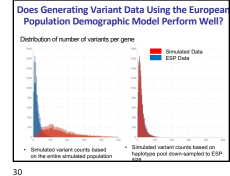
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- Provides a haplotype matrix
- 10,000 haplotypes over 200kb region
- Simulated using a calibrated coalescent model (cosi)
- Mimicking linkage disequilibrium structure of European ancestry
- User can also provide haplotype data
- Power and sample size calculations for binary and quantitative traits

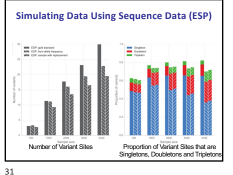
User specify proportion of variants that increase or lower risk



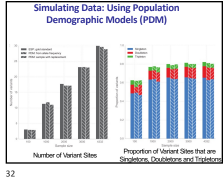


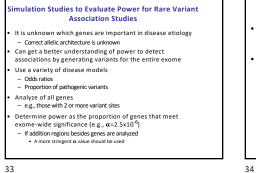










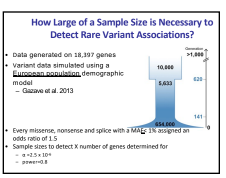


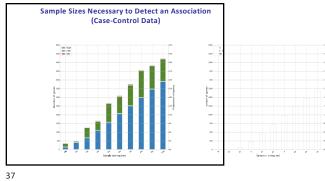
# **Power Analysis**

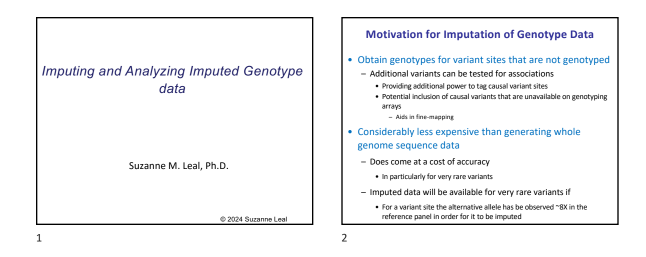
- For tests of individual variants - Power depended on sample size, disease prevalence, minor
- allele frequency, genetic model and variant effect size For rare variants (aggregate association tests)
- Also dependent on the allelic architecture
- Cumulative variant frequency within analyzed region
- Proportion of causal variants How much contamination from non-causal variants
- · Effect sizes the same the same or different across gene regions Effects of variants in the same or different directions
  - » Protective and detrimental for binary traits
    » Increase and decrease quantitative trait values

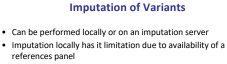
## **Power Analysis Rare Variants** (Aggregate Association Tests)

- · Power will not only vary between traits greatly
- The power to detect an association will also vary drastically between genes for the same complex trait
- For some causal genes even with hundreds of thousands of samples power will be low
- While for other causal genes a few thousand samples may be sufficient





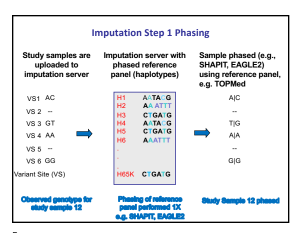


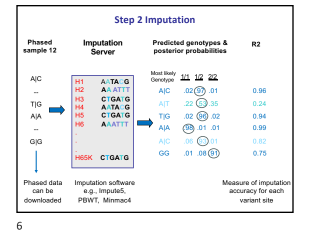


- Internal data
- 1000 genomes
- Haplotype reference consortium (HRC)
  - Only part of this dataset is made publicly available
- Smaller imputation panels will impact the ability to impute lower frequency and rare variants
  - Additionally, regardless variant MAF a decrease in the size and diversity
    of imputation panel will lead to a decrease in the imputation accuracy

Phasing and performing imputation using an Imputation Server

3





5

# Measures of Imputation Accuracy

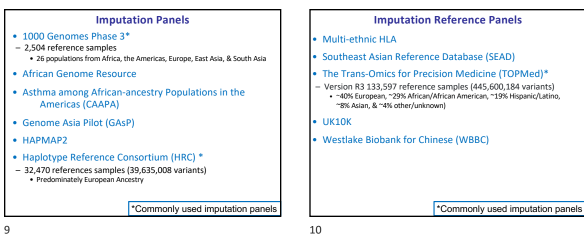
R<sup>2</sup>/INFO

- Measures of imputation accuracy Most programs report R2 Impute provides INFO scores
- r<sup>2</sup> is the correlation between the dosage and genotype obtained from sequence or genotype array data
  - Must have imputed data and sequence or genotype array data for the same person to estimate r<sup>2</sup>.

7

# Step 3 Analysis of Imputed Data • Variants are filtered according to R<sup>2</sup> values e.g., analyze variants with an R<sup>2</sup>>0.8 Most likely genotypes are <u>not</u> analyzed instead dosages are analyzed The dosage can be estimated as follows for variant site 1 sample 12: A|C with prior probabilities 1/1= 0.02, 1/2=0.97, & 2/2=0.01 (R<sup>2</sup>=0.96) Genotype 1/1 0 x 0.02 = 0.0 Genotype 1/2 1 x 0.97 = 0.97 <u>Genotype 2/2 2 x 0.01 = 0.02</u> Dosage 0.99 The dosage for variant site 2 sample 12: A|T with prior probabilities 1/1=0.22, 1/2=0.53, & 2/2=0.35 (R<sup>2</sup>=0.23) • Genotype 1/1 0 x 0.22 = 0.0 Genotype 1/2 1 x 0.53 = 0.53 Genotype 2/2 2 x 0.35 = 0.70 Dosage 1.23

8



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# **Imputation Servers**

- Michigan (US)
  - Reference panels include, HRC, 1,000 Genomes, etc.
  - Phasing EAGLE2
  - Imputation Minmax4
  - https://imputationserver.sph.umich.edu/index.html#!
- NHLBI (US)
  - Reference panel TOPMed
  - Phasing EAGLE2
  - Imputation Minmax4
- https://imputation.biodatacatalyst.nhlbi.nih.gov/#!

- Phasing SHAPEIT or EAGLE2 - Imputation PBWT

• Sanger (UK)

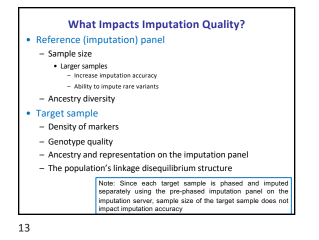
- https://www.sanger.ac.uk/tool/sanger-imputation-service/
- Westlake (People's Republic of China)
  - Reference panels include 1000 Genomes, GAsP, SEAD, & WBBC

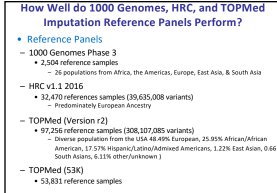
**Imputation Servers** 

- Reference panels include HRC, 1,000 Genomes, etc.

- Phasing SHAPEIT2
- Imputation Minmax4
- https://imputationserver.westlake.edu.cn/index.html

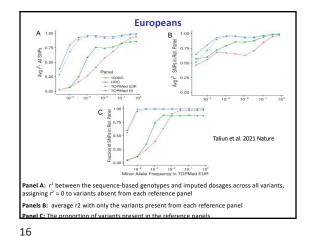




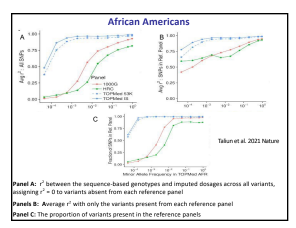


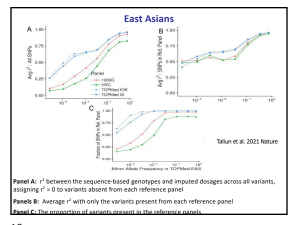
How Well do 1000 Genomes, HRC, and TOPMed Imputation Reference Panels Perform? • Target Sample - 100 ancestry specific samples, • e.g. Europeans, African-Americans, & South Asians

- Obtained from BioMe
  - Samples are not included in any of the reference panels

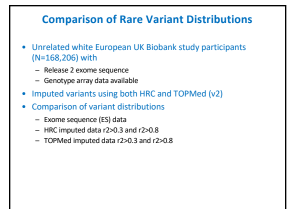


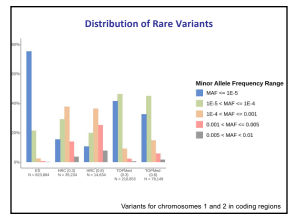
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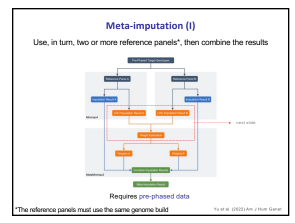












Imputation of Variants without using an

Imputation Server • Imputation locally has it limitation due to availability of

Only part of this dataset is made publicly available to download to use locally

• Due to data sharing limitation in particular within the European

It may not be possible to use imputation servers which are located in the

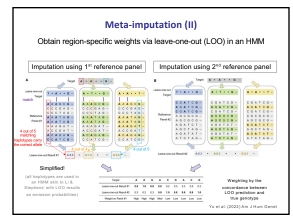
· Can be computationally intensive to phase and impute

• All haplotype phasing and imputation software used on

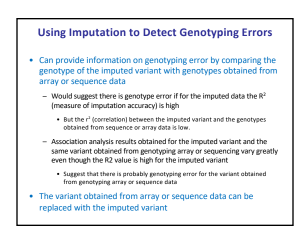
imputation servers are publicly available

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Union

references panels

genotypes locally

US, UK or China

Internal data
1000 genomes

- HRC



# Combining data obtained from different genotyping arrays

- Variants that don't overlap between arrays can be imputed
  - As well as variants not available on any of the arrays
- Caution should be used because the imputation quality can vary between datasets
  - Influenced by different error rates between datasets
  - Principal components analysis (PCA) can be used to determine if the potential problems
    - If additional quality control is necessary
- If there are more cases or controls for a particular dataset – Type I errors can be increased
  - Type Terrors can be increa

# Linkage disequilibrium in genetic association studies

# Gao Wang, Ph.D.

Advanced Gene Mapping Course, May 2024 The Gertrude H. Sergievsky Center and Department of Neurology Columbia University Vagelos College of Physicians and Surgeons

# Genetic association studies (recap)

### Identify genetic variants associated with complex traits

- · Association does not imply causality
- Disease, quantitative traits, molecular phenotypes

# in order to

2

4

- Understand biological mechanism
- Identify potential drug targets
- Identify individuals with high disease risk

1

# Sources of association signals

### Causal association — meaningful

- · Tested genetic variations influence traits directly
- Linkage disequilibrium (LD) —useful
- Tested genetic variations associated with other nearby variations that influence traits
- Meaningful or misleading, in different contexts

### Population stratification - misleading

- Tested genetic variations is unrelated to traits, but is
   associated due to sampling differences
- eg, minor allele frequency, disease prevalence

# Sources of association signals: analysis tools

- Causal association meaningful
- Fine-mapping, colocalization, Mendelian randomization
- Linkage disequilibrium (LD) —useful
- Meaningful: LD scores regression, polygenic risk scores (PRS), transcriptome-wide association studies (TWAS)
- Misleading: fine-mapping, LD pruning / dumping
- Population stratification misleading
- Principle component analysis, linear (mixed) models

# 3

# Linkage disequilibrium (LD)

## LD: the sharing of certain combinations of variants

- · Formally, equivalents to Haplotype structure
- There are several measures of LD but largely irrelevant to
   our learning objectives
- In gene-mapping, let's simply understand LD as Pearson's correlation between variants

# Linkage disequilibrium (LD)

Levels of LD is a result of chromosomal "shuffling"

# Segregation and Recombination



# Each row is a variant site

Shuffle within rows does not change marginal MAF.
Multi-loci MAF, *i.e.*, *haplotype frequency*, will change.

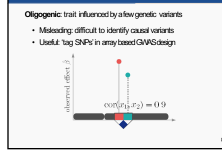
# 5

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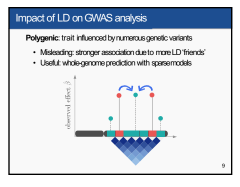
# Why do we care about LD?

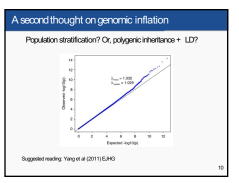
- When obviously LD is an issue
- Many variants will look "similar" by genotype but have different biological function — mapping "causal" variants is challenging
- When LD is useful
- Can leverage non-causal genetic variables to predict phenotypes when causal variant is not observed in data
- Can leverage variants that are LD to infer each other's
- genotype to complete missing genotype data
  - · also, association study summary statistics

# Impact of LD on GWAS analysis



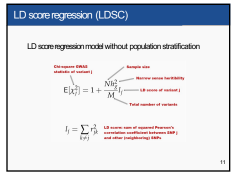
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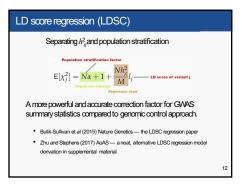


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# LDSC application: heritability estimation

# Narrow sense heritibility

 Proportion of phenotypic variation explained by additive genetic factors

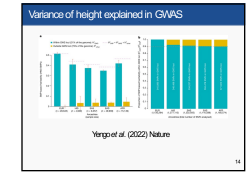
# Estimation strategy

- Pedigree design: genetic covariance and IBD sharing
- Population design: linear mixed models

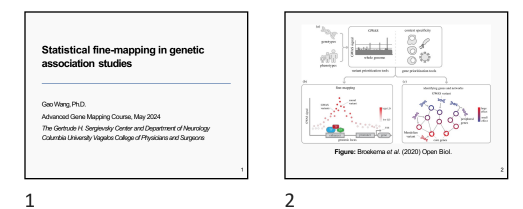
# Population design, summary statistics

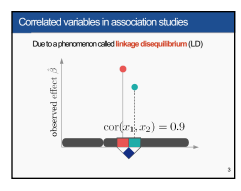
LDSC to estimate SNP-basedheritability
 Stratified LDSC (S-LDSC) to partition heritability by
functional annotations

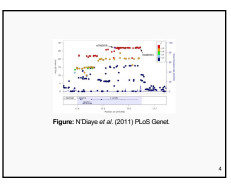


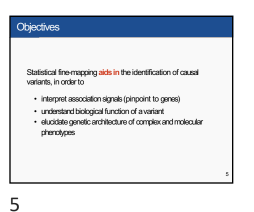


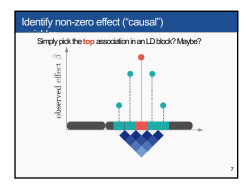
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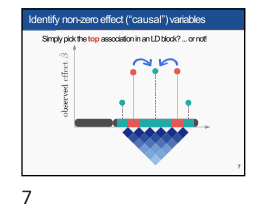


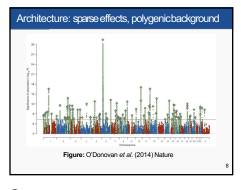


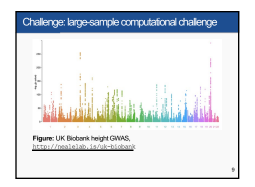


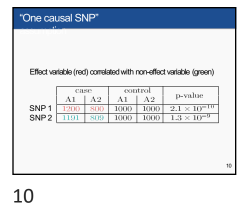


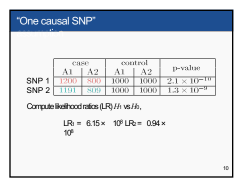


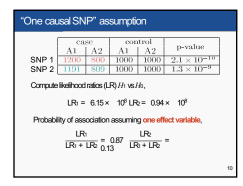


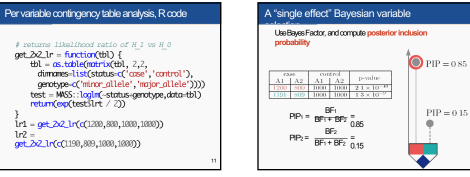




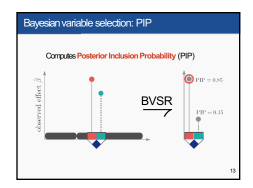


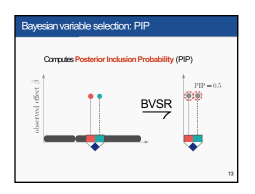




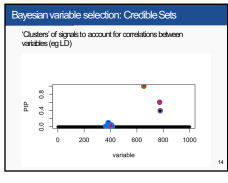






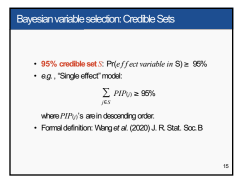


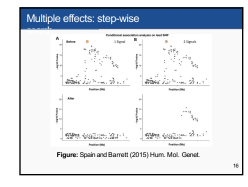












# A simple frequentist conditional analysis Forward selection algorithm 1. For each SNP fit a simple linear regression model 2. Select the SNP / that has the largest model likelihood 3. Form residuals y' := y- X<sub>i</sub>b<sub>i</sub>, and repeat

# 20

# A simple frequentist conditional analysis Forward selection algorithm 1. For each SNP fit a simple linear regression model 2. Select the SNP/ that has the largest model likelihood 3. Form residuals $y' := y - Xb'_{n}$ , and repeat A greedy algorithm to choose the "best" SNPs, but is incapable of capturing multiple SNPs in LD

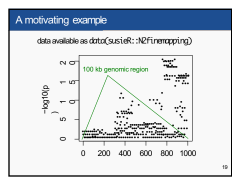
# To quantify uncertainty

# Bayesian forward selection algorithm

- For each SNP j, fit a simple Bayesian linear regression model to get Bayes Factor BFj
- 2. Form weight for each SNP,  $w_j \propto BF_j$
- 3. Form residuals  $\mathbf{y}' := \mathbf{y} \sum_{j \ w_j} \mathbf{X}_{jb} \hat{j}$ , and repeat

# 21

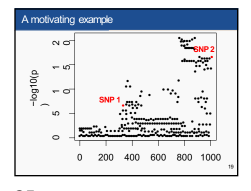
# To quantify uncertainty Bayesian forward selection algorithm For each SNP*j*, fit a simple Bayesian linear regression model to get Bayes Factor BF<sub>7</sub> Form weight for each SNP, w<sub>1</sub> ≈ BF<sub>7</sub> Form residuals y := y - ∑ w<sub>2</sub> X<sub>2</sub>b<sub>1</sub>, and repeat What if a "bad decision" is made early on?

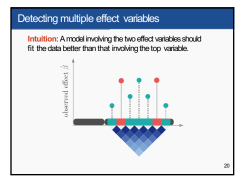


24

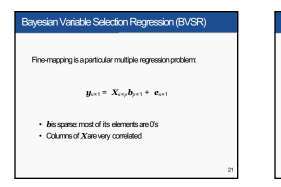
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18

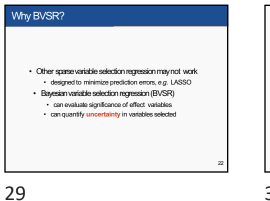


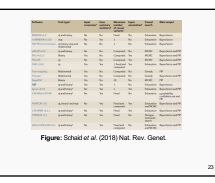


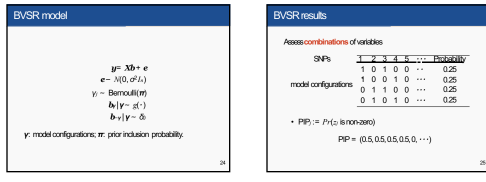


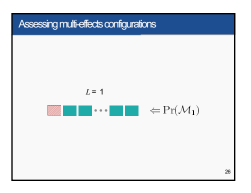


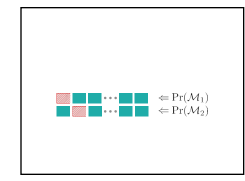




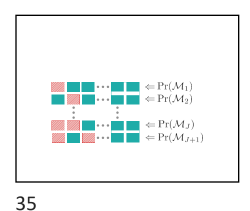


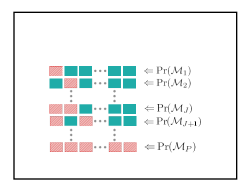


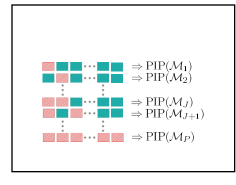




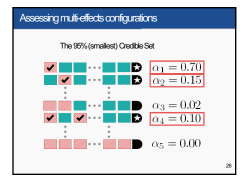


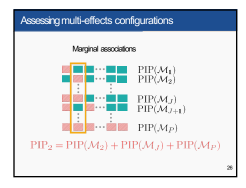


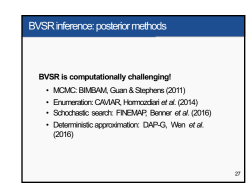




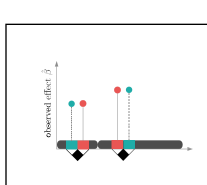








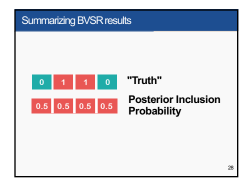


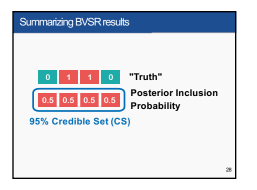


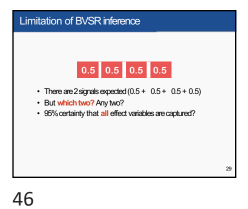


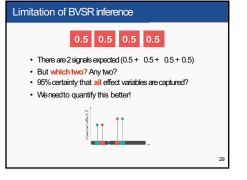


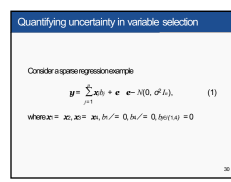


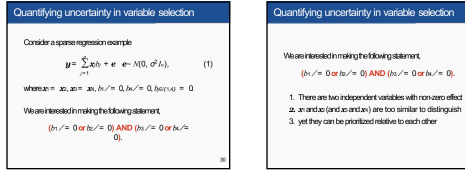






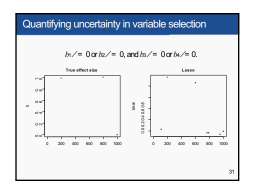


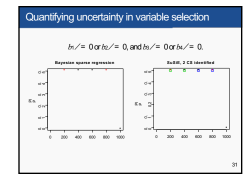


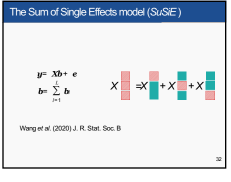




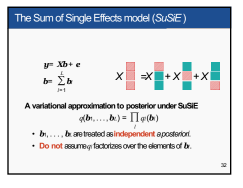


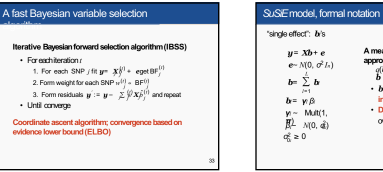






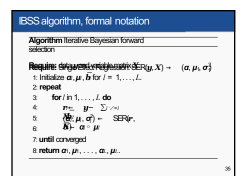


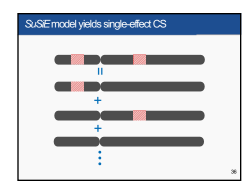












A mean-field

approximation

 $q(\mathbf{b}_1,\ldots, L) = \prod_l q_l(\mathbf{b}_l)$ 

•  $\boldsymbol{b}_1,\ldots,\boldsymbol{b}_{\!\scriptscriptstyle L}$  are treated as

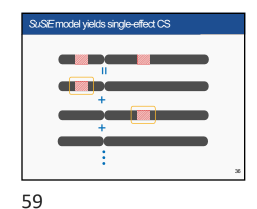
independent a posteriori.

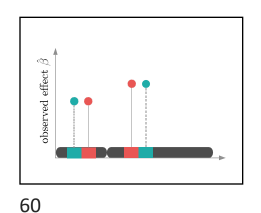
over the elements of  $b_{\ell}$ .

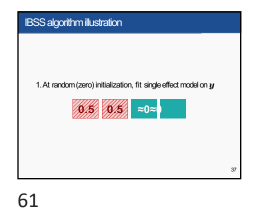
• Do not assume ql factorizes

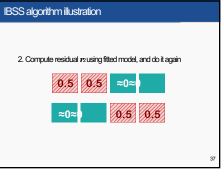


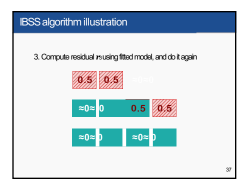
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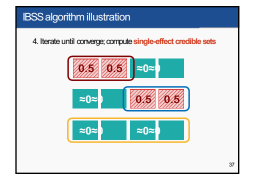


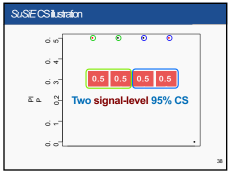


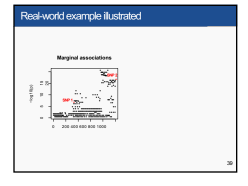


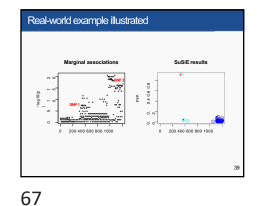


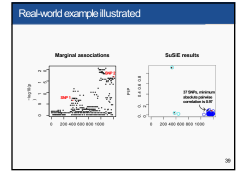


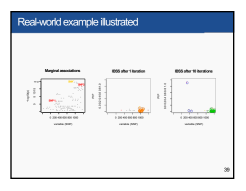


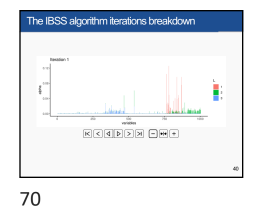




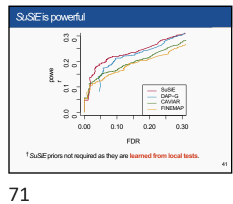




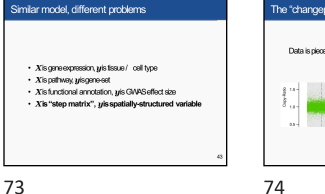


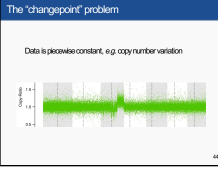


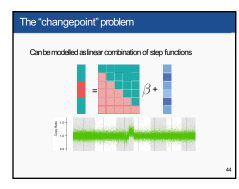


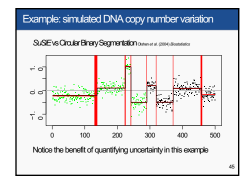


<i>SuSiE</i> is	fast										
Speed comparison (3 causal variables; unit: sec.)											
	Method	Avg.	Min.	Max.							
	SuSiE <sup>†</sup>	0.64	0.34	2.28							
	DAP-G	2.87	2.23	8.87							
	FINEMAP	23.01	10.99	48.16							
	CAVIAR	2907.51	2637.34	3018.52							
<sup>†</sup> An Rimplementation of SuSIE. Others are implemented in C++.											
						42					

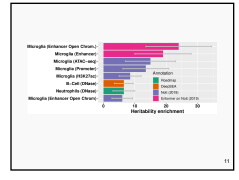








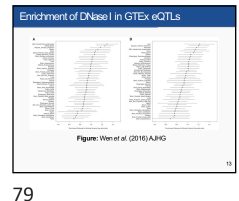
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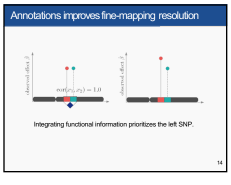




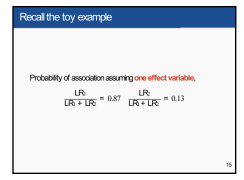
76

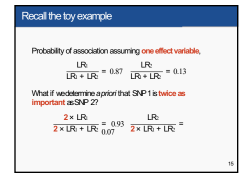
A sparse model (a somewhat oligogenic Generalized linear model for SNP effects given K annotations  $\beta_j = (1 - \pi_j)\delta_0 + \pi_j g(\Theta)$  $\pi_j := \Pr(\gamma_j = 1 | \boldsymbol{\alpha}, \boldsymbol{d})$  $\log \left(\frac{\pi_j}{1-\pi_j}\right) = \alpha + \sum_{k=1}^{K} \alpha_k d_k$  $\alpha$  are log fold enrichment of functional genomic features Suggested reading: Wen (2016) AoAS 12



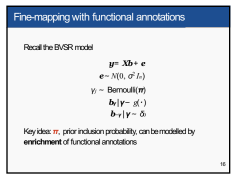








81

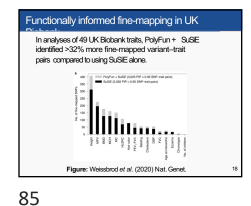


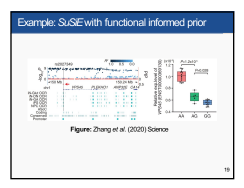
### Genome-wide approach with S-LDSC

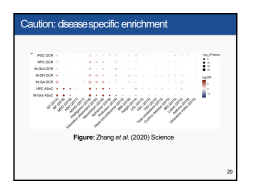
- A single locus may not have enough power to leverage annotation enrichment
- Genome-wide evaluation of thousands of annotations can increase power of fine-mapping
  - · Lead to new loci to discover
- Functional enrichment can be done under the same framework
- Prioritize genomic features / tissues / cell-types
- · Enrichment coefficient may be transferrable cross population
  - Weissbrod et al. (2022) Nat. Genet.

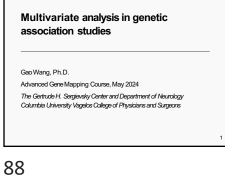
83

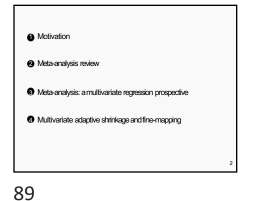
77

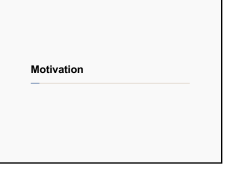












### Beyond per trait per variant association studies

### Statistical fine-mapping (multiple regressors)

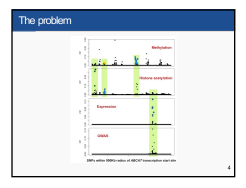
Identify non-zero effect variables by accounting for LD

### Meta-analysis (multiple responses)

Integrate information across multiple conditions / studies

### "Causal" variants across multiple conditions?

Cross-population fine-mapping; colocalization; pleiotropy; mediation; . . .



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# The problem For agenetic variable analyzed in two conditions: P("causal" in trait 1 & 2 | association data for 1 & 2) 5

### The problem

For a genetic variable analyzed in two conditions:

P("causal" in trait 1 & 2 | association data for 1 & 2)

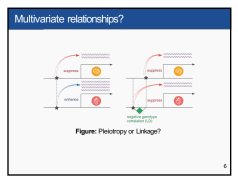
Denote data as  $D_1$  and  $D_2$ , and use indicator variables  $\gamma_1$ ,  $\gamma_2$  for variable having effects in 1 and 2, and hyperparameters  $\Theta$ :

 $P(\gamma_1 = 1, \gamma_2 = 1 | D_1, D_2, \Theta)$ 

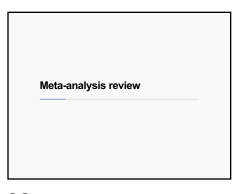
93

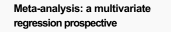
91

# 94







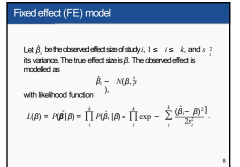


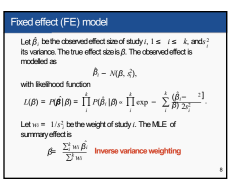
### Fixed effect and random effects models

### Different assumptions on effects across studies

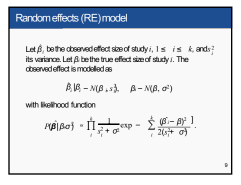
 Fixed effect model: all studies share a common effect size
 Random effects model: effect sizes are random variables from an underlying distribution

97





99



## 101

100

98

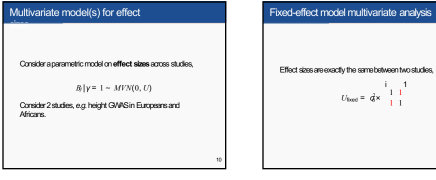
### Random effects (RE) model

Let  $\hat{\beta}_i$  be the observed effect size of study i,  $1 \le i \le k$ , and  $s_i^2$  its variance. Let  $\beta_i$  be the true effect size of study i. The observed effect is modelled as

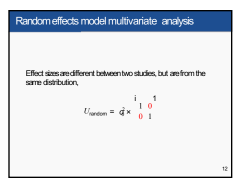
 $\beta_i \mid \beta_i \sim N(\beta_i s_i^3), \quad \beta_i \sim N(\beta, \sigma^2)$ 

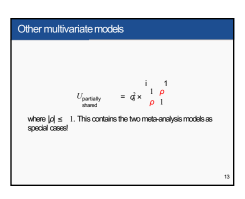
$$P(\hat{\boldsymbol{\beta}}|\boldsymbol{\beta},\boldsymbol{\sigma}^{2}) \propto \prod_{i}^{k} \frac{1}{s_{i}^{2} + \sigma^{2}} \exp - \sum_{i}^{k} \frac{(\boldsymbol{\beta}_{i} - \boldsymbol{\beta})^{2}}{2(s_{i}^{2} + \sigma^{2})}.$$

RE has weight  $w_i^* = 1/(s_i^2 + \sigma^2)$ ; summary effect  $\beta$ can be similarly computed as FE, replacing  $w_i$  with  $w_i^*$ .  $\sigma^2$  can be estimated (e.g., MLE).



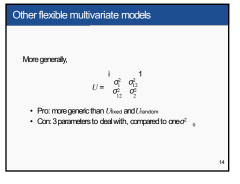






 $U_{\text{fixed}} = q_0^2 \times \frac{\begin{vmatrix} i & 1 \\ 1 & 1 \end{vmatrix}}{1 1}$ 

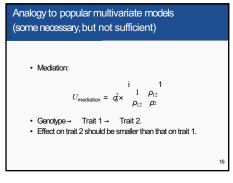
105

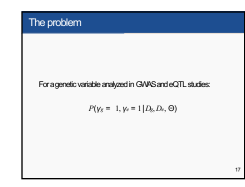


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Analogy to popular multivariate models (some necessary but, not sufficient) · Colocalization correlation matrix:  $^{i}_{1} \rho^{1}$ ρ1 Condition specific correlation matrix: 15





# 110



coloc [Giambartolomei et al. (2014) PLoS Genet.]

- On X: "one causal" assumption
- On Y: the null + 4 combinations given "one causal"
- 1. In 1 but not 2
- 2. In 2 but not 1
- In 1 and 2 but not the same variable
   In 1 and 2 and the same variable (colocalization)
- 5. No association in both data 1 and 2

# Colocalization method: eCAVIAR

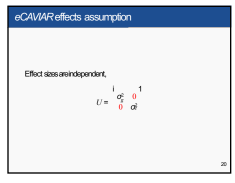
eCAVIAR [Hormozdiari et al. (2016) Am. J. Hum. Genet.]

- On X: multiple effect variables
- On Y: each effect variable can be

1. In 1 but not 2 2. In 2 but not 1

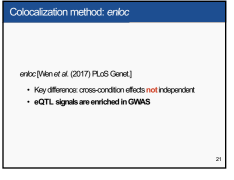
- 3. In both 1 and 2
- 4. No association in both data 1 and 2

# 111

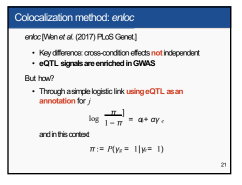












# 116

enloc two step procedure

# Connections between colocalization methods

- eCAVIAR is a special case of enloc with  $\alpha = 0$ .
- coloc is a special case of "one causal" fine-mapping based enloc with fixed, high(!) α value by default.
- Recent coloc extension: coloc version 5, aka SuSiE-coloc removed the "one causal" assumption.
  - Wallace (2021) PLoS Genetics
     https://chrlswallace.github.io/coloc/

### Connections between colocalization methods

1. Obtain  $P(\gamma_g = 1)$  and  $P(\gamma_e = 1)$  using fine-mapping

22

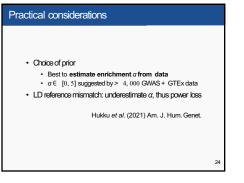
23

2. Fit the enrichment model via multiple imputation

- eCAVIAR is a special case of enloc with α = 0.
   coloc is a special case of "one causal" fine-mapping based enloc with fixed, high (!) α value by default.
- Recent coloc extension: coloc version 5, aka SuSiE-coloc removed the "one causa" assumption.
- Wallace (2021) PLoS Genetics
- https://chrlswallace.github.io/coloc/

Summary: pattern and scale of effect size correlations, represented as different prior models.

# 117



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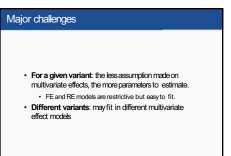
# 118

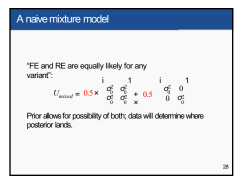
23

Multi-trait colocalization										
		Hypothesis	Number of hypotheses		Example configuration	Number of configurations				
	No 1	No association with any of in trails	1		Truit 1 (00000-0) Truit 2 (00000-0) Truit m (00000-0)	1				
	N1 -	One that has a CV in the region			• (1000-0 0000-0 1111111	ng .				
	Na -	Two traits have a shared CV	(7)		<ul> <li>(10000-9) (10000-9) (1111151) (00000-A)</li> </ul>	( <sup>10</sup> <sub>2</sub> )e				
	N <sub>210</sub> -	Two traits have distinct CVs	(7)		• (10000-4) (11111\1 (0000-A) 0 S(11)	${m \choose 2} (Q(Q-1))$				
					(10000-0)	1.1				
	K <sub>[4-1,1]</sub> :	m - 2 traits share a CV two balts have distinct CVS	()		C(300-9 00:00-9 00:00-9 00:00-9 00:00-9	() x00-0 x00-0				
	K98-5.0 '	m - 1 traits share a CV one trait has a CV elsewhere	-		(10000-€) (0000-€) (11111<) (0000-£) € X <sub>(00-610</sub>	#002-10				
	x., -	- Falls have a shared CV	1		$\begin{pmatrix} 18000-9\\ 18000-0\\ 11111 \\ 18000-0\\ 18000-0 \end{pmatrix} \in S_n$	0				
		C	$\operatorname{Pell}(m+1)$		$indicator = \begin{cases} i, causal variant \\ 0, otherwise \end{cases}$	$(Q+1)^{\rm m}$				
Figure: HyPrColoc, Foley et al. (2021) Nat. Comm.										
Assuming a single causal variant in the loci.										



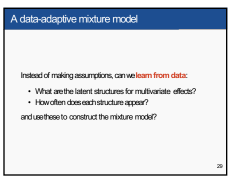
# 



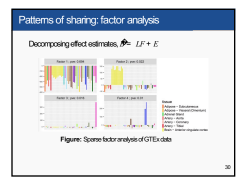


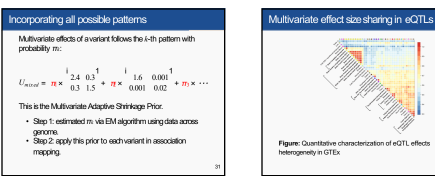
or, ...

# 

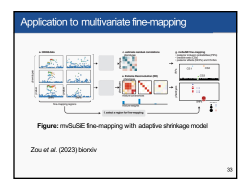


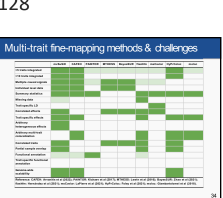


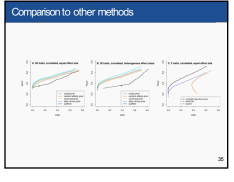




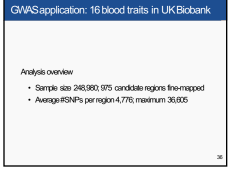




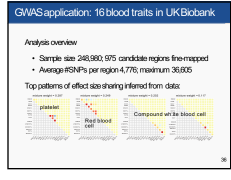


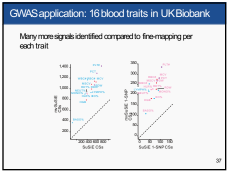










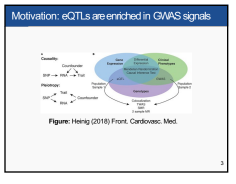




Complex phenotype prediction and transcriptome-wide association studies Gao Wang, Ph.D. Advanced Gene Mapping Course, May 2024 The Gertrude H. Sergievsky Center and Department of Neurology Columbia University Vagelos College of Physicians and Surgeons

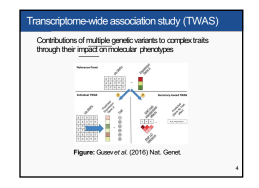
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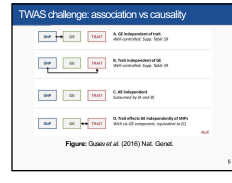
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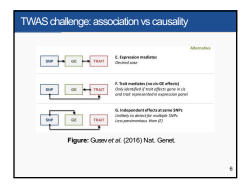


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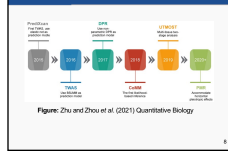
# 140 TWAS methods overview

### Ideal TWAS setup

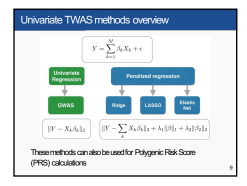
- Homogenous population
- Tissue and cell-type specific
- Training data-set is large and complete (N > 200)
   But in reality

TWAS challenge: technical considerations

- Cross population TWAS aplications
- Multiple tissue and cell-types
- Availability of individual level data vs summary statistics

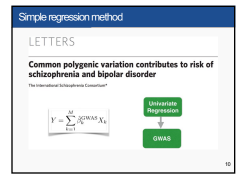


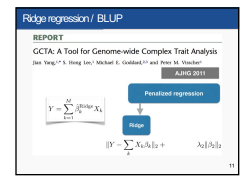
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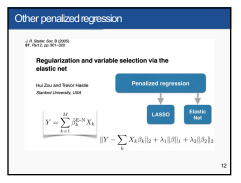


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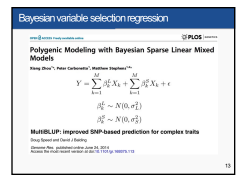
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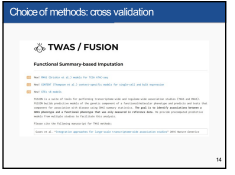




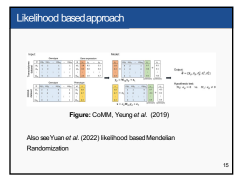


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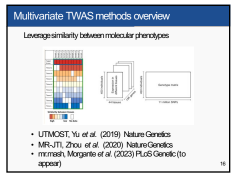


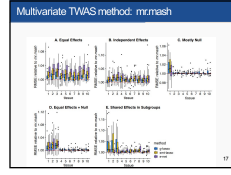


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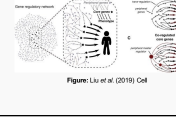




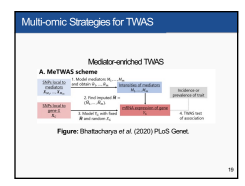


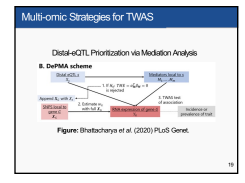


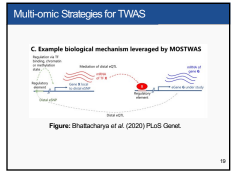




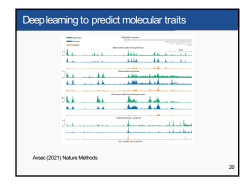
An omnigenic view of genetic regulations

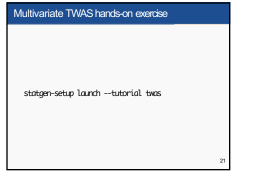






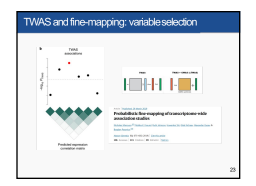


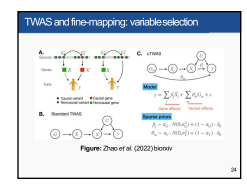




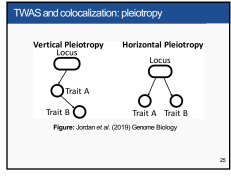
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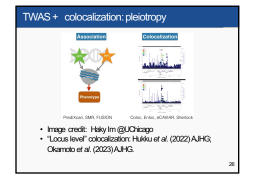


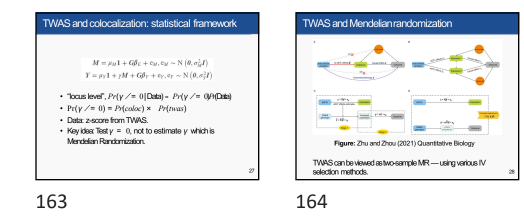












### Fine-mapping with summary statistics: current methods and practical considerations

### Gao Wang, Ph.D.

Advanced Gene Mapping Course, May 2024 The Gertrude H. Sergievsky Center and Department of Neurology Columbia University Vagelos College of Physicians and Surgeons

## Association analysis summary statistics =scores from univariate association studies: $z_{j}^{2}:=\beta_{j}^{2}/s_{j}$ , where $\beta_{j}^{2}:=(\mathbf{x}_{j}^{T}\mathbf{x})^{-1}\mathbf{x}^{T}\mathbf{y}$ $s_{j}^{2}:=-\overline{\sigma_{j}^{2}(\mathbf{x}_{j}|\mathbf{x})^{-1}}$ • Sufficient statistics: $\mathbf{x}^{T}\mathbf{x}, \mathbf{x}^{T}\mathbf{y}, \sigma_{j}^{2}$ • "Summary" statistics: • =scores: $\hat{z}$ • Genotypic correlation: $\mathbf{\hat{K}}$

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### Reasons to work with summary statistics

Advantage over full data (genotypes and phenotypes):

- Easier to obtain and share with others
- Convenient to use: QC and data wrestling barely needed
- Computationally suitable for large-sample problems
   O(p<sup>2</sup>) (summary statistics) ≪ O(np) (full data)
  - when sample size n ≫ variants in fine-mapped region p

Suggested reading: Pasaniuc and Price (2017) Nat. Rev. Genet.

# Regression with Summary Statistics (RSS) $\hat{z} \sim N(\hat{Rz}, \hat{R})$ Assumptions: 1. Heritability of any single SNP is small 2 $\hat{R}$ is sample genotypic correlation for the same study 3. Genotypes used to computed *z* and $\hat{R}$ are accurate

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### Properties of per SNP *z* scores • *z*-score for a SNP depends on effects of both itself and other correlated SNPs: $E(z)R^{*}) = \sum_{i=1}^{p} r_{ij}z_{i}.$

GWAS marginal effects are biased due to LD!

z-scores are correlated,

 $Cor(\hat{z_j}, \hat{z_k}) = r_{jk}, \forall j, k$ 

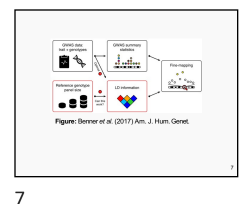
Recall the previously discussed connection with LDSC

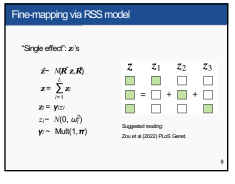
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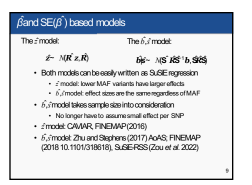
Summary of summary statistics

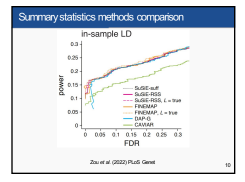
- X, genotype matrix
- Y, phenotype matrix, can be multiple traits
- $X^T Y$ , association results effect size estimate
- $X^{\chi}$ , LD matrix
- XX<sup>T</sup>, genomic relatedness matrix, reflects kinship
   Y<sup>Y</sup>, trait correlation, relevant in multi-trait analysis and integration

6

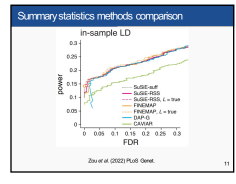








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# Impact of allele flips

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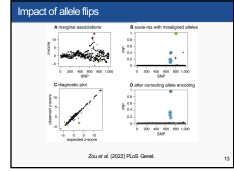
### What is allele flip?

Different allele encoding between GWAS and LD reference
 e.g. AA=0, AC=1, CC=2 in GWAS; AA=2, AC=1,
 CC=0 in LD reference genotype

12

 A challenging problem coupled with strand flip, when merging sequence data from different platforms

11



### Addressing the allele flip challenge

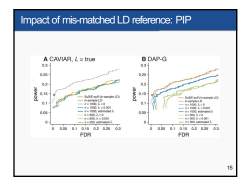
- susieR::susie\_rss() function implements a diagnosis bigsnpr::snp\_match() function implements a basic allele matching for two sets of summary statistics
- Other resources

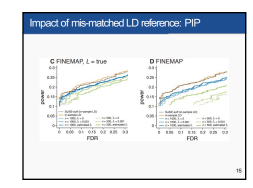
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 Allele flip illustration: <u>https://stataen.us/</u> <u>lab-wiki/compbio\_tutorial/allele\_ac</u> A powerful, multi-set data merger (by Yin Huang): https://cumc.github.io/xgtl-pipeline/ pipeline/misc/summary stats merger.html

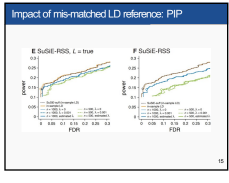
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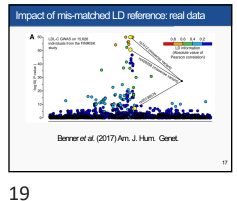


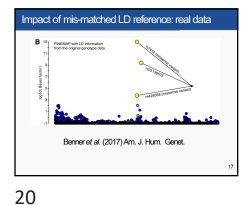
Impact of mis-matched LD reference: credible sets

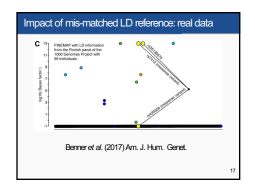


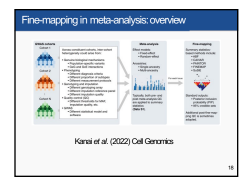


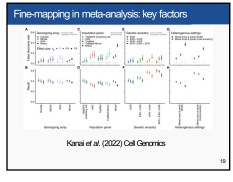




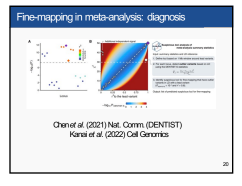


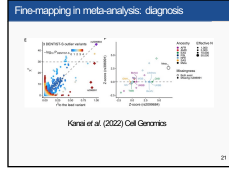














2. Fit model Trait ~ Age+Sex+PCs, compute residual of Trait (remove covariates), and evaluate SNP effect in model Residual Trait ~ SNP

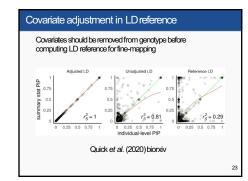
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Are these two analysis equivalent?

More technical details see McCaw et al. (2020) Biometrics

25

# Covariate adjustment in LD reference Consider two GWAS regression analysis: 1. Evaluate SNP effect in Trait ~ SNP+Age+Sex+PCs 2. Fit model Trait ~ Age+Sex+PCs, compute residual of Trait (remove covariates), and evaluate SNP effect in model Residual [Trait ~ SNP More technical details see McCaw et al. (2020) Biometrics



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### Gao Wang, Ph.D.

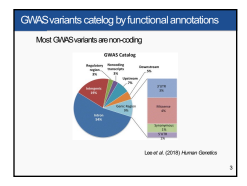
Advanced Gene Mapping Course, May 2024 The Gertrude H. Sergievsky Center and Department of Neurology Columbia University Vagelos College of Physicians and Surgeons

Non-coding functional annotation in GWAS

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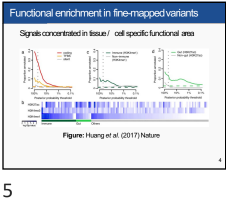
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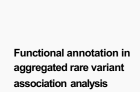
# Which are the causal variants In which cell types do the variants act? SNP enrichment Which genes are regulated by the variants? H. Colocalization

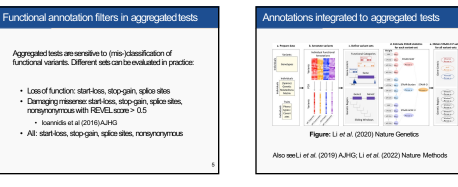


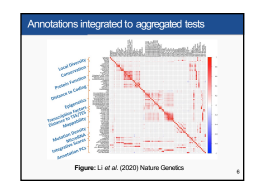
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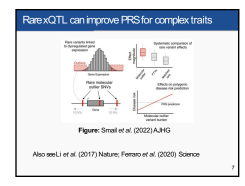
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A polygenic model: stratified LD score regression

Nh

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 $E[\chi_{i}^{2}] = 1 +$ 

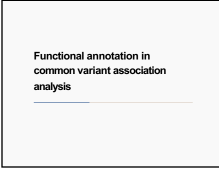
 $l_j = \sum_{k \neq j} r_{jk}^2$ 

LD score of variant j

im of squared Pearson's coefficient between SNP j leighboring) SNPs

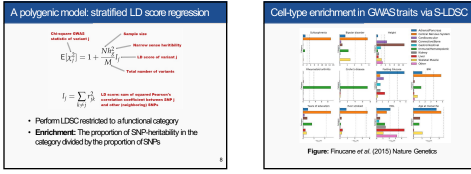
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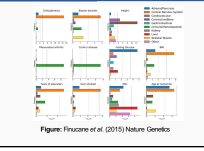
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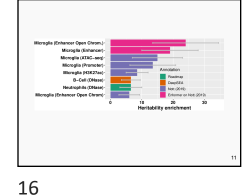


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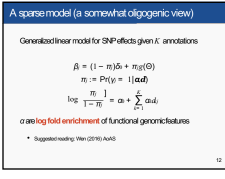
### Integration approaches

- · Integrate directly as range based binary annotations • Finucane et al (2015) Nature Genetics - Stratified LDSC paper
- Extension: variant specific continuous annotations · Gazal et al (2017) Nature Genetics
- · Tissue specific variant level annotations independent of GWAS results

  - Deep Learning methods
    Zhou et al (2015) Nature Genetics, Zhou et al (2018) Nature Genetics, Lai et al. (2022) PLoS Comp Bio
  - · Avsec et al. (2021) Nature Methods

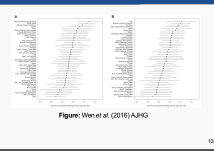


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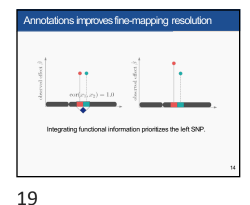


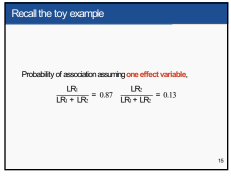
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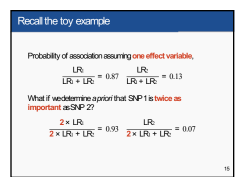
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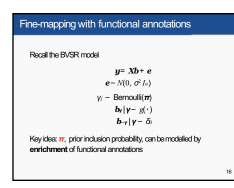
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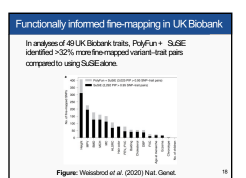


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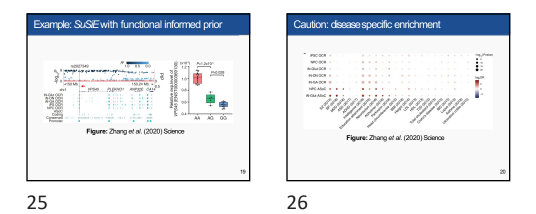
### Genome-wide approach with S-LDSC

- A single locus may not have enough power to leverage annotation enrichment
- Genome-wide evaluation of thousands of annotations can increase power of fine-mapping
  - · Lead to new loci to discover
- Functional enrichment can be done under the same framework
- Prioritize genomic features / tissues / cell-types
- Enrichment coefficient may be transferrable cross population
  - Weissbrod et al. (2022) Nat. Genet.



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# Multivariate analysis in genetic association studies

### Gao Wang, Ph.D.

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2 Meta-analysis review

Meta-analysis: a multivariate regression prospective

Multivariate adaptive shrinkage and fine-mapping

# Motivation

### Beyond per trait per variant association studies

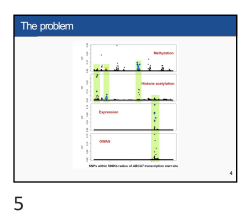
### Statistical fine-mapping (multiple regressors)

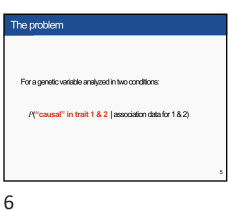
- Identify non-zero effect variables by accounting for LD
- Meta-analysis (multiple responses)
- Integrate information across multiple conditions / studies

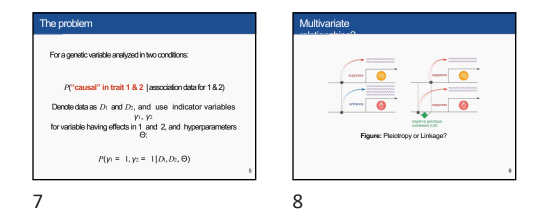
"Causal" variants across multiple conditions?

 Cross-population fine-mapping; colocalization; pleiotropy; mediation; . . .

4



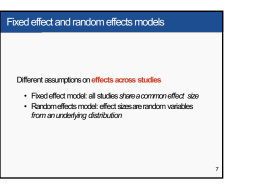




Meta-analysis review

Meta-analysis: a multivariate regression prospective

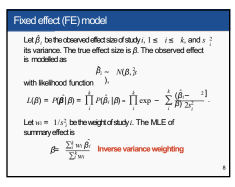
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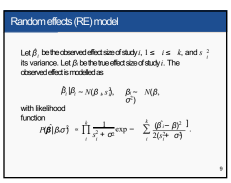


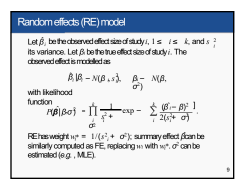
### Fixed effect (FE) model

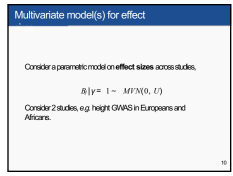
```
Let \hat{\beta}_i be the observed effect size of study i, 1 \le i \le k, and s \stackrel{?}{=} i
its variance. The true effect size is \beta. The observed effect is
modelled as
\hat{\beta}_i \sim N(\beta, \frac{2}{\beta})
with likelihood
function
L(\beta) = P(\hat{\beta}|\beta) = \prod_i^k P(\hat{\beta}_i|\beta) \propto \prod_i^k \exp - \sum_i^k \frac{(\hat{\beta}_i - \beta)^2}{2s_i^2}.
```

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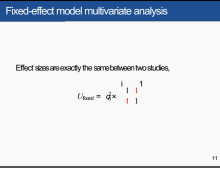






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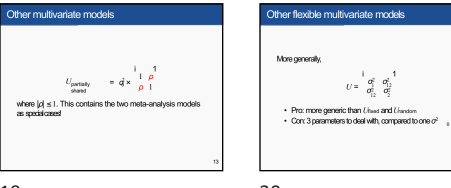
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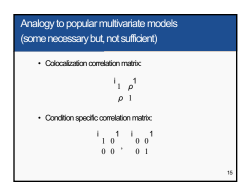


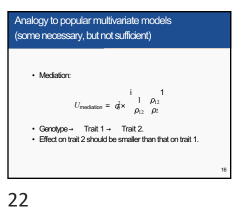


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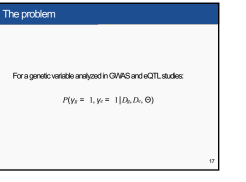
Random effects model multivariate analysis Effect sizes are different between two studies, but are from the same distribution,  $U_{\text{random}} = q_{1}^{2} \times \begin{pmatrix} i & 1 \\ 1 & 0 \\ 0 & 1 \end{pmatrix}$ 







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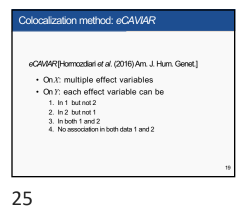


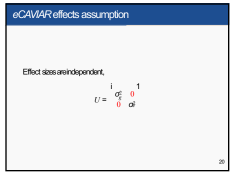
• On Y: the null + 4 combinations given "one causal" 1. In 1 but not 2 2. In 2 but not 1 3. In 1 and 2 but not the same variable 4. In 1 and 2 and the same variable (colocalization) 5. No association in both data 1 and 2

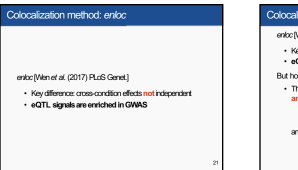
Colocalization method: coloc

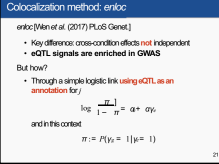
coloc [Giambartolomei et al. (2014) PLoS Genet.] • On X: "one causal" assumption

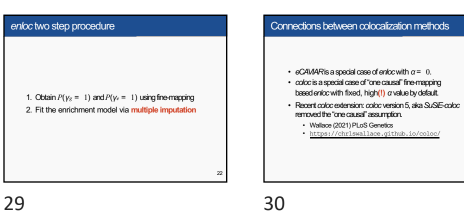
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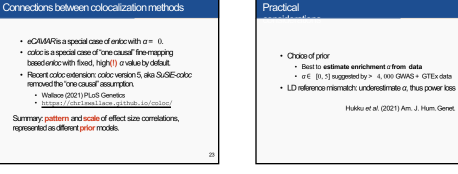






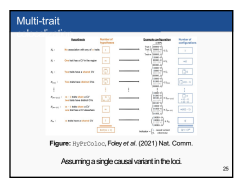


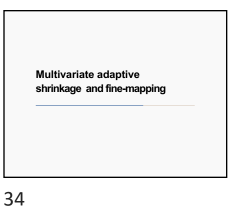


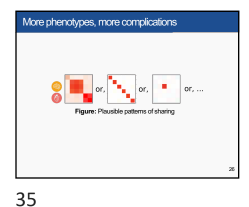


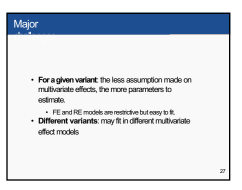


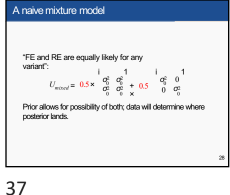


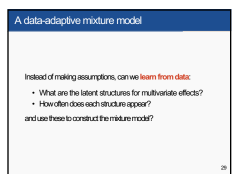


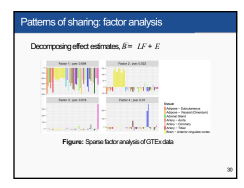


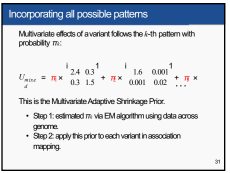


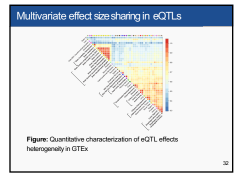


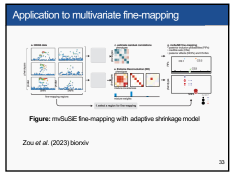




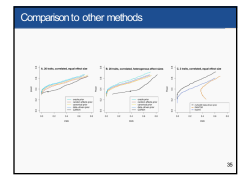


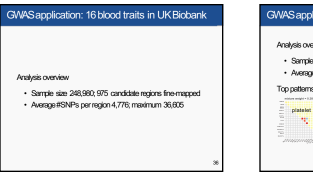


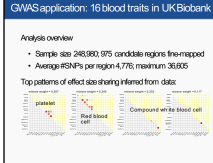


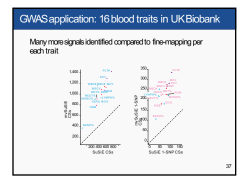












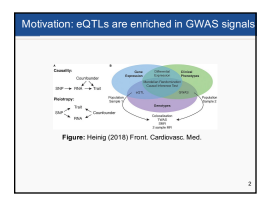


Complex phenotype prediction and transcriptome-wide association studies

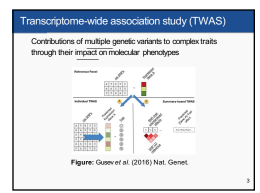
Gao Wang, Ph.D.

Advanced Gene Mapping Course, May 2024 The Gertrude H. Sergievsky Center and Department of Neurology Columbia University Vagelos College of Physicians and Surgeons

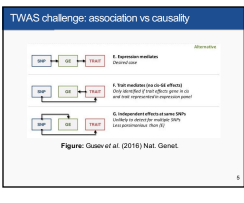
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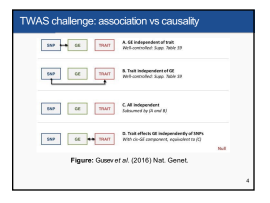
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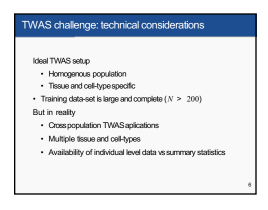
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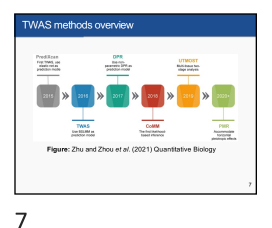




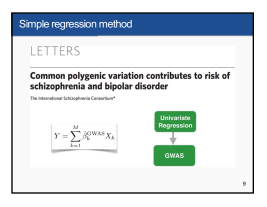


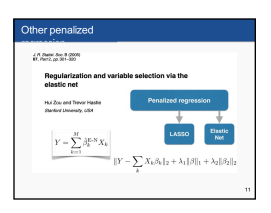
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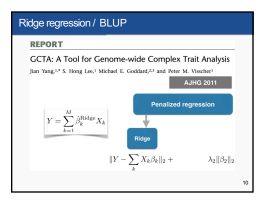


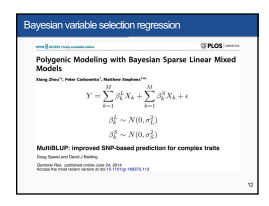


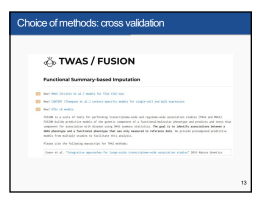
Univariate TWAS methods overview 
$$\begin{split} & Y = \sum_{k=1}^{M} \beta_k X_k + \epsilon \\ & \downarrow \\ &$$

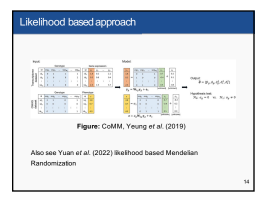


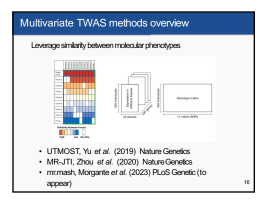


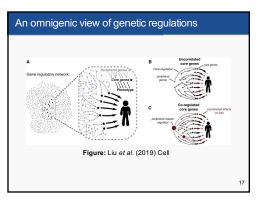


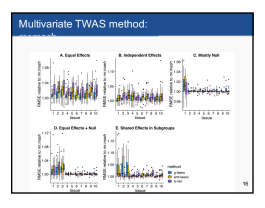




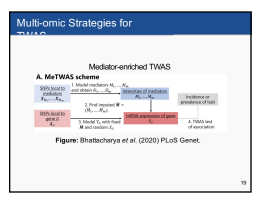




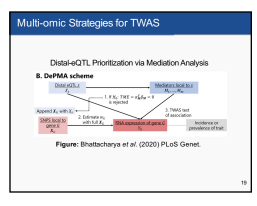


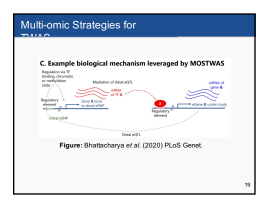


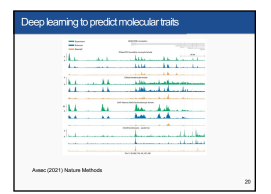




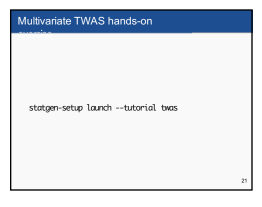


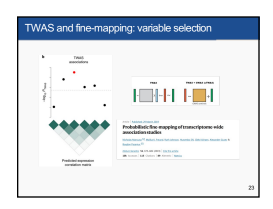




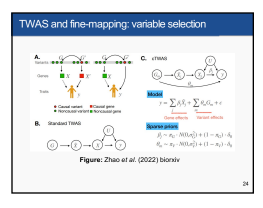


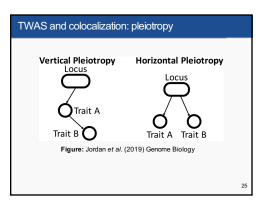


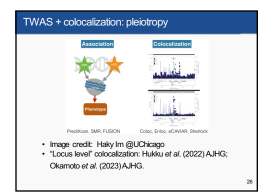


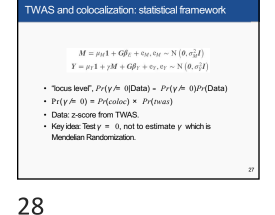


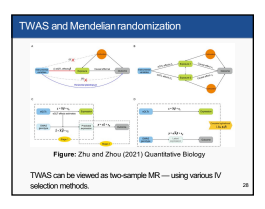




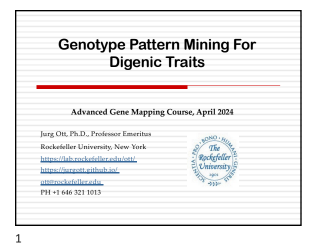




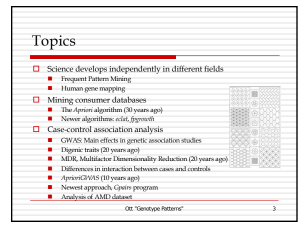


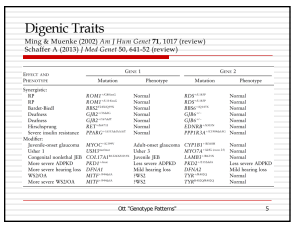


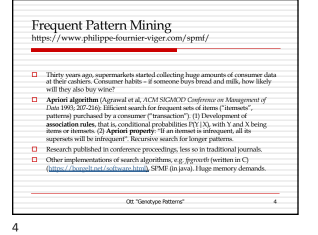


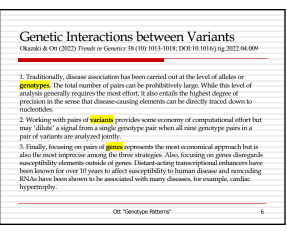


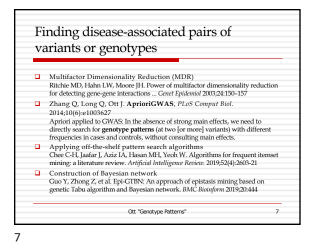


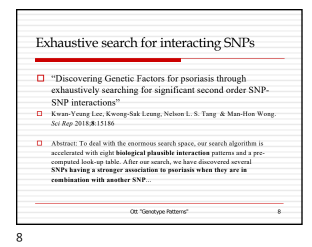




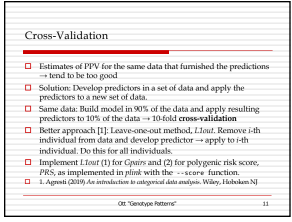




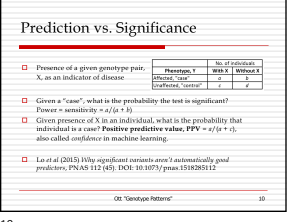




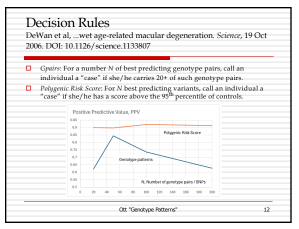
Gpairs program: All pairs of genotypes, schizophrenia data https://github.com/jurgott/gpm\_prog\_ □ Schizophrenia case-control data: 1,044 cases and 2,052 controls genotyped for 892,850 SNPs. Pruned and focused on males. Evaluate all pairs of genotypes for SNPs. For each SNP pair, analyze each of the 9 genotype pairs: 81,972,176,883 genotype pairs tested. Distribute work over many threads (CPUs, up to 192 CPUs in new PCs). For each genotype pair, X, make 2 × 2 table: Min. 20 occurrences of any genotype No. of individuals With X Without X Phenotype, Y pair (support) Affected, "case" Unaffected, "control" Π. Each table analyzed by Fisher test a pBon = min(#tests × pNom, 1) □ 69 genotype pairs significantly more frequent in cases than controls □ Genotypes → variants → genes: Network of 17 genes **Prediction**, classification:  $c = 0 \rightarrow \text{person with } X \text{ must be a case}!$ Ott "Genotype Patterns" ٥



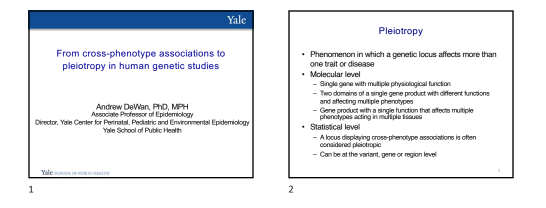


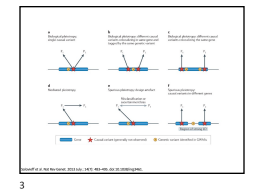


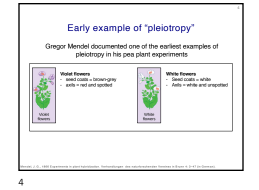












#### Examples in humans

Marfan syndrome

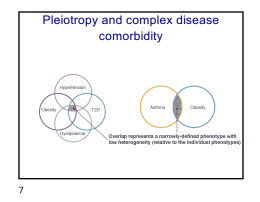
- FBN1 (fibrillin-1)
- thinness, joint hypermobility, limb elongation, lens dislocation, and increased susceptibility to heart disease.
- Holt-Oram syndrome,
- TBX5 (transcription factor)

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- cardiac and limb defects
- Nijmegen breakage syndrome
  - NBS1 (DNA damage repair protein)
  - microcephaly, immunodeficiency, and cancer predisposition

## Pleiotropy and complex disease comorbidity

- Examples of correlated (comorbid) disease
- Obesity, hypertension, dyslipidemia, type 2 diabetes (metabolic disorder)
- Depression, anxiety, personality disorders (psychiatric disorder)
- Asthma, obesity (pro-inflammatory conditions)
- · Why do certain disease occur together
  - Causality
- Shared environmental risk factors
- Shared genetic risk factors



## Pleiotropy and complex disease comorbidity

- · Pleiotropy-informed analyses consider multiple phenotypes together and take into account the correlation between the phenotypes
- Analyzing multiple correlated phenotype (e.g. comorbid diseases) is equivalent to analyzing a single narrowly-defined phenotype with low heterogeneity

#### Pleiotropy and complex disease comorbidity

- · Detecting shared genetics and/or molecular pathways between comorbid diseases can help us understand exactly how the etiology of the diseases overlap
- · Etiologic overlaps:

9

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- · provide opportunities for novel interventions that prevent or treat the comorbidity, rather than preventing/treating each disease separately
- · facilitate drug repurposing (that is, known drugs targeting a pleiotropic locus may be repurposed to treat other diseases controlled by that locus, precluding the need for the development and testing of a brand-new drug)

Abundant Pleiotropy in Human Complex Diseases and Traits Shanya Sivakumaran,<sup>1,6</sup> Felix Agakov,<sup>1,2,6</sup> Evropi Theodoratou,<sup>1,6</sup> James G. Prendergast,<sup>3</sup> Lina Zgaga,<sup>1,7</sup> Teri Manolio,<sup>5</sup> Igor Rudan,<sup>1</sup> Paul McKeigue,<sup>1</sup> James F. Wilson,<sup>1</sup> and Harry Campbell<sup>1,\*</sup> The American Journal of Human Genetics 89, 607–618, November 11, 2011 Table 6. Extent of Pleiotropy in Different Disease Classes Genes Pleiotropic (%) Nonpleiotropic (%) p Value<sup>®</sup> SNPs Pleiotropic (%) Nonplei lisease Class %) p Value\* 
 233 (16.9)
 1147 (83.1)
 77 (4.6)
 1610 (95.4)

 pes
 106 (17.7)
 175 (62.3)
 <0.0001</td>
 31 (8.3)
 343 (91.7)
 (comparison group) e-mediated phenotypes 106 (37.7) 0.0066 
 173 (02.3)
 COMMI
 31 (8.3)
 343 (91.7)

 92 (65.2)
 <0.0001</td>
 8 (4.8)
 158 (95.2)

 198 (71.5)
 <0.0001</td>
 30 (8.4)
 327 (91.6)
 49 (34.8) 0.8456 79 (28.5) 198 (71.5) 0.0056 The statement of the st Restrict free (FRCH) Trainer Biol (FRCH) Construction (FRCH) Construction (FRCH) Brief Bourne (FRCH) Brief Bourne (FRCH) Source Color I (CARI) Patrosofia Secon 2,599 Source regeneral (MAC) Mercelle: age of error) (F Hercelle: age of error) (F Hercelle: Age of error) (F

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#### Pleiotropy in gene mapping

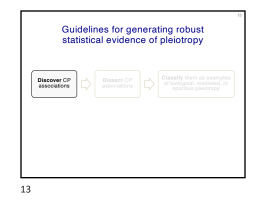
- · Mapping a single genotype to multiple phenotypes has the potential to uncover novel links between traits or diseases
- · It can also offer insights into the mechanistic underpinnings of known comorbidities
- · It can increase power to detect novel associations with one or more phenotypes

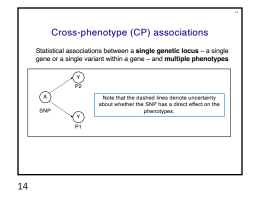
## A practitioners' guide for studying pleiotropy in genetic epi studies

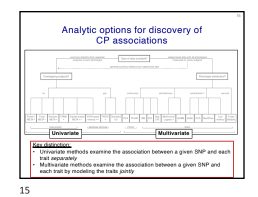
Am J Epidemiol. 2017 Aug 11. doi: 10.1003/aja/wx296. (Epub ahead of print) Statistical Analysis of Multiple Phenotypes in Genetic Epidemiological Studies:From Cross-Phenotype Associations to Pleiotropy.

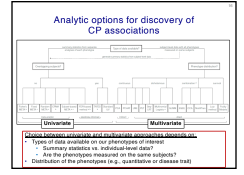
#### Salinas YD, Wang Z, DeWen AT.

Additional technological pointing where is the phenomenon in which a single grantic loss affects must have one built or descent and constraint grantic pointing where the phenomenon in which a single grantic loss affects must have one built or descent and the constraint grantic pointing with the phenomenon in the ph KEYWORDS: genetic epidemiology; mediation analysis; pleiotropy









#### Univariate methods are by far the most commonly used to detect CP associations

- Univariate methods include (but are not limited to) the methods you've discussed in class so far:
- · allelic Chi-Square test
- genotypic Chi-Square test
- · regression-based methods
- · The overall approach is to:
  - · obtain univariate association p-values for each phenotype declare CP associations at genetic loci that are statistically significantly associated with each phenotype



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Hypothetical example: Discovery of CP associations for hypertension and heart disease by using logistic regression

Step 1. Fit two univariate regression models within PLINK  $E[hypertension] = \beta_0 + \beta_1 * SNP$  $E[heart disease] = \beta_0 + \beta_1 * SNP$ 

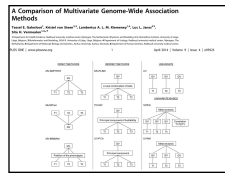
Word of caution: The univariate tests of association should be marginal tests (conducted irrespectively of the second phenotype) NOT conditional tests (conducted on a subset defined based on absence/presence of the second phenotype). In this example, what that means is that the regression for hypertension should be fit on all subjects irrespectively of their heart disease status; and the regression for heart disease should be fit on all subjects irrespectively of their hypertension status. More on this later!

#### Hypothetical example: Discovery of CP associations for hypertension and heart disease by using logistic regression

- Step 1. Fit two univariate regression models within PLINK  $E[hypertension] = \beta_0 + \beta_1 * SNP$
- $E[heart \, disease] = \beta_0 + \beta_1 * SNP$
- Step 2. For a given SNP, examine p-values for β<sub>1</sub> from each model.
  P-value for β<sub>1</sub> in hypertension model = 1.03 x 10<sup>-12</sup>
- P-value for  $\beta_1$  in heart disease model = 6.02 x 10<sup>-9</sup> Step 3. Declare CP associations at a given SNP, if the p-values for  $\beta_1$  in
- acch model surpass the study significance threshold.
  Assuming the standard GWAS significance threshold (alpha=5 x10<sup>-8</sup>), there
- is a statistically significant association with both hypertension and heart disease at this particular SNP. Therefore, we have sufficient statistical evidence to declare a CP association at this SNP.

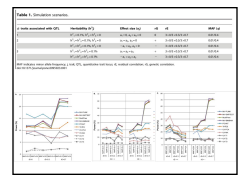
Using multivariate methods to increase the power to detect cross-phenotype associations

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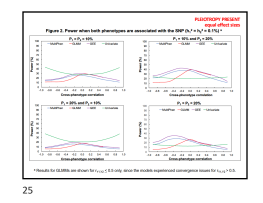
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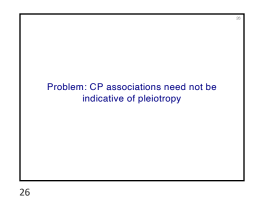


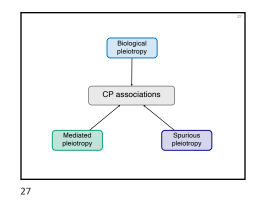
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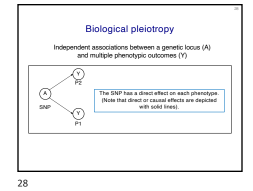
A comparison of univariate and multivariate GWAS methods for analysis of multiple dichotomous phenotypes	# traits
Yasmmyn D. Salinas <sup>1</sup> , Andrew T. DeWan <sup>1</sup> , and Zuoheng Wang <sup>2</sup>	2
<sup>1</sup> Department of Chronic Disease Epidemiology; <sup>2</sup> Department of Biostatistics, Yale School of Public Health, Yale University, 60 College St, New Haven, Connecticut, USA	2
Genet. Epidemiol. 41 (7), 689-689	

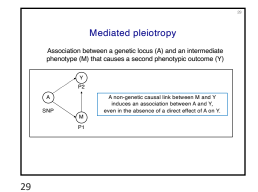
	Simulation s	nulation scenarios					
# traits associated	h <sub>i</sub> ²	r <sub>Y1,Y2</sub>	Pj				
1	h12=0.1%,h22=0%	[-0.9,0.9]	P1 = P2 = 10%				
			P1 = P2 = 20%				
			P1 = 10%, P2 = 20%				
			P1 = 20%, P2 = 10%				
2	h <sub>1</sub> <sup>2</sup> = h <sub>2</sub> <sup>2</sup> = 0.1%	[-0.9,0.9]	P1 = P2 = 10%				
			P1 = P2 = 20%				
			P1 = 10%, P2 = 20%				
			P1 = 20%, P2 = 10%				
2	h <sub>1</sub> <sup>2</sup> = 0.1%,h <sub>2</sub> <sup>2</sup> = 0.05%	[-0.9,0.9]	P1 = P2 = 10%				
			P1 = P2 = 20%				
			P1 = 10%, P2 = 20%				
			P1 = 20%, P2 = 10%				

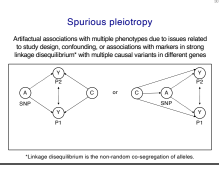


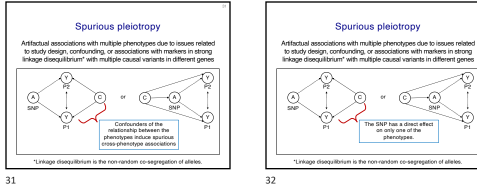








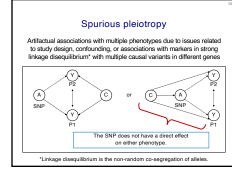




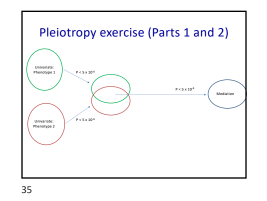
(A)

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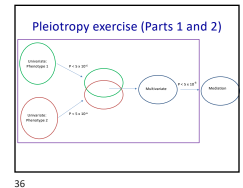


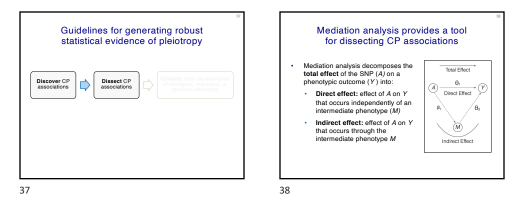
Spurious pleiotropy

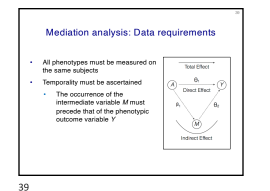
Artifactual associations with multiple phenotypes due to issues related to study design, confounding, or associations with markers in strong linkage disequilibrium\* with multiple causal variants in different genes

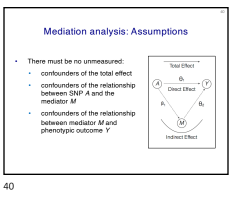
Variables associated with the phenotypes and the SNP induce spurious cross-phenotype associations

\*Linkage disequilibrium is the non-random co-segregation of alleles.

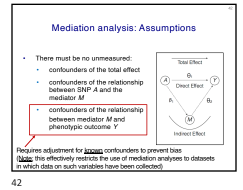


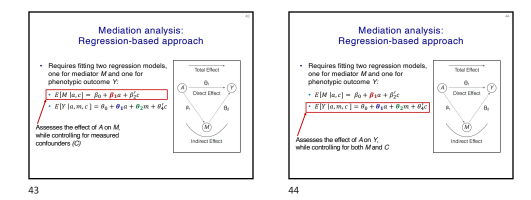


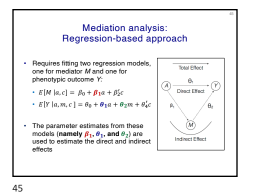


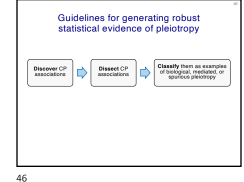


Mediation analysis: Assumptions Typically met in genetic epi studies . There must be no unmeasured: Total Effect confounders of the total effect θ. (A) · (Y) confounders of the relationship Direct Effect between SNP A and the mediator M β, ∕ θ₂ confounders of the relationship M between mediator M and phenotypic outcome Y Indirect Effect



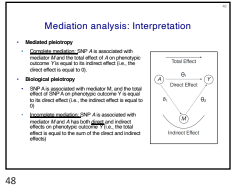


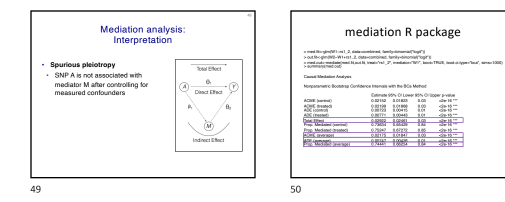


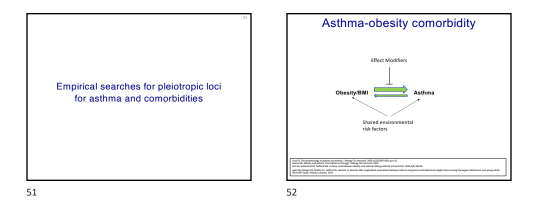


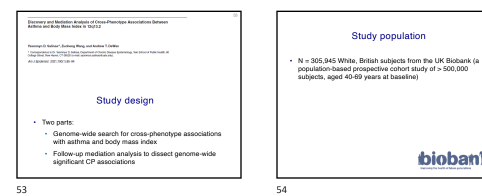
Mediation analysis: Interpretation Mediated pleiotropy <u>Complete mediation</u>: SNP A is associated with mediator M and the total effect of A on phenotypic outcome Y is equal to its indirect effect (i.e., the Total Effect direct effect is equal to 0). θ Incomplete mediation: SNP A is associated with mediator M and A has both direct and indirect (A) -· (Y) Direct Effect effects on phenotypic outcome Y (i.e., the total effect is equal to the sum of the direct and indirect effects) β, θ₂ Biological pleiotropy M SNP A is associated with mediator M, and the total effect of SNP A on phenotypic outcome Y is equal to its direct effect (i.e., the indirect effect is equal to Indirect Effect

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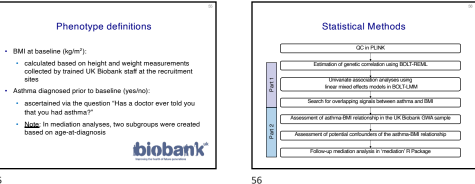


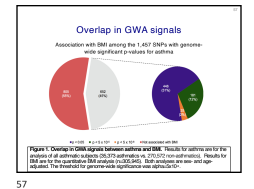


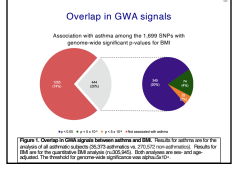


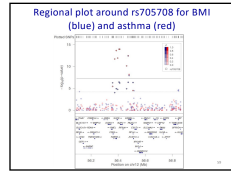


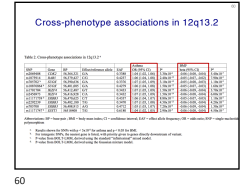
biobank



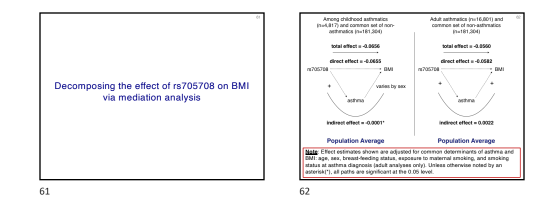


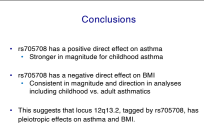












#### Conclusions

- 12g13.2 is multigenic and our CP associations span genes CDK2, RAB5, SUOX, IZK4, RPS26, ERBB3, and ESYT1.
- rs705708 is the top regional BMI signal and resides in ERBB3. The top regional asthma signal, rs2456973, resides in IZKF4.
- · While rs705708 and rs2456973 could be in LD with the same causative variant in either ERBB3 or IKZF4 or another gene in 12q13.2, it is also possible that each variant could tag a distinct, trait-specific causative variant in different genes.
- Therefore, locus 12q13.2 displays pleiotropic effects on asthma and BMI, but this may not be an example of pleiotropy at the gene level (biological pleiotropy).

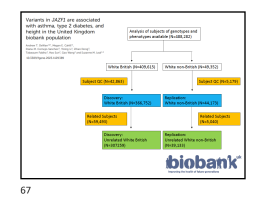
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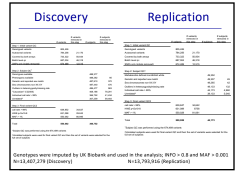
64

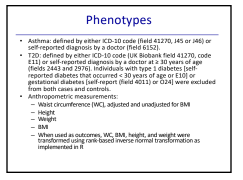
What if we expand this investigation to look at more phenotypes correlated with asthma?

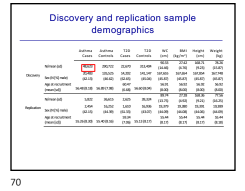
#### Asthma, T2D and anthropometric measures

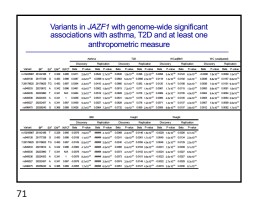
- Obesity is a well-established risk factor for both asthma and T2D.
- While highly correlated, waist circumference (WC) can provide distinct information on adiposity as it is a measure of visceral obesity, specifically WC adjusted for BMI. WC is often used in studies of chronic diseases.
- success of UNION Liseases. In Transmission of UNION as been shown to be an additional risk factor for Ingread astimutes been adjusting for BMI Elevated blood glucose and T2D have been linked to increased risk of asthma in adults, and conversely, asthma has been associated with increased risk of developing T2D in adults.
- adults
- Height is a highly heritable polygenic trait; there is evidence that shorter individuals have an increased risk for developing T2D and individuals with childhood onset asthma have shorter stature as adults compared to non-asthmatics

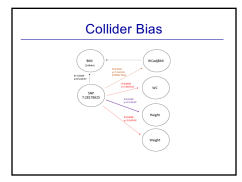


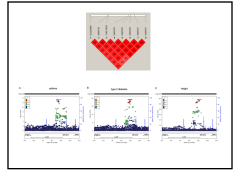






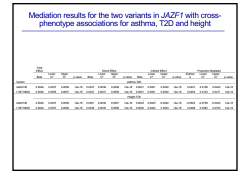








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					ation purel	en Autor	• 12	D Her	,
Variant I	Position A1	AF*	NFO'	Antre	720	Heigh CS	PP CS	PP CS	100
7,27940054	27345054 CT	0.067	0.096	3345-01	6245-01	2.54E-16 None	0.0038 None	0.0034 HEIGHT 1	0.0047
18773503722	27949622 CA	0.089	0.967	0.005-01	3 005-02	0.33E-13 None	0.0034 None	0.0047 HEIGHT_1	0.0248
rs12538298	27982312 C	0.059	0.984	9.475-01	7.405-02	1.12E-17 None	0.0036 None	0.0038 HEIGHT_1	0.0223
157783433	27962633 C	0.099	0.984	9.186-01	6.25E-02	1.08E-17 Nore	0.0034 None	0.0039 HEIGHT_1	0.0233
194722748	27988368 C	0.069	0.983	9.085-01	6.486-02	2.00E-17 None	0.0034 None	0.0039 HEIGHT_1	0.0172
196070113	2799182N C	0.050	0.983	9.145-01	7.685-02	2,21E-17 None	0.0034 None	0.0038 HEIGHT_1	0.0167
rs1557779	27294758 T	0.072	0.964	9565-01	2,255-02	1.102-18 None	0.0034 None	0.0044 HEIGHT_1	0.0585
7.20154215	20154215 GT	0.792	0.979	3/036-13	8.325-05	4.STE-02 ASTHMA_1	0.0452 None	0.0034 None	0.0035
194722758	28156606 C	0.000	0.999	1.075-13	6.666-05	3.57E-02 ASTHMA_1	0.1135 None	0.0034 None	0.0033
156077955	28156887 C	0.800	0.999	1.616-13	6.80E-05	329E-02 ASTHMA_1	0.0779 None	0.0034 None	0.0054
194719822	28158058 C	0.800	1.000	1.426-13	6.448-05	3.58E-02 ASTHMA_1	0.0877 None	0.0034 None	0.0033
7.28150/95	28/60/96 GT	0.190	0.854	2645-01	1.138-02	1.68E-57 None	0.0040 None	0.0033 HEIGHT_2	1.0000
ra9645345	20160113 C	0.000	1.000	1.505-13	6.265-05	3.50E-02 ASTHMA_1	0.0034 None	0.0034 None	0.0033
7.28156745	20160345 GTCTT	0.7%2	0.996	6715-14	2.505-05	1.15E-02 ASTHMA_1	0.1410 None	0.0034 None	0.0028
112189905	28172014 C	0.000	1.000	3015-13	1725-05	3.00E-02 ASTHMA_1	0.0440 None	0.0034 None	0.0028
152180000	28172066 T	0.792	1,000	1,406-13	4.50E-08	1.35E-02 ASTHMA_1	0.0711 None	0.0034 None	0.0054
194722760	28172183 A	0.800	0.999	3076-13	7.106-05	2.89E-02 ASTHMA_1	0.0432 None	0.0034 None	0.0038
19017116	28172586 T	0.792	0.000	1.365-13	4505-06	1.44E-02 ASTHMA_1	0.0729 None	0.0034 None	0.0035
rs702014	28472732 C	0.494	0.997	6.97E-08	6.205-18	1.252-56 None	0.0052 T2D_1	0.0200 None	0.0017
18917116	26472739 T	0.791	0.996	2.325-13	4.305-05	1.20E-02 ASTHMA_1	0.0454 None	0.0034 None	0.0033
1967250450	28174966 T	0.800	0.000	3065-13	6.535-05	3.05E-02 ASTHMA 1	0.0454 None	0.0034 None	0.0029
10017117	28478305 G	0.001	0.996	1945-13	7,055-05	3.67E-02 ASTHMA 1	0.0656 None	0.0034 None	0.0034
rs11492901	28477301 C	0.001	0.996	1,935-13	7.096-05	3.60E-02 ASTHMA_1	0.0660 None	0.0034 None	0.0034
11004745	20100555 T	0.494	1.000	2,896-07	3 836-18	1.725-70 None	0.0048 T2D_1	0.0286 None	0.0021
19840142	26165691 T	0.494	1.000	4756-07	2315-18	7.775-70 None	0.0046 T2D_1	0.0071 None	0.0018
+\$11455069	28186775 A	0.494	0.999	8796-07	3.55E-18	1.12E-69 None	0.0047 T2D_1	0.0013 None	0.0018
1910622246	26167111 A	0.486	0.994	5.965-07	1.065-18	3.08E-68 None	0.0046 T2D_1	0.0850 None	0.0020
ra640133	26/192260 C	0.405	0.999	4.565-07	7,255-19	2.005-71 None	0.0047 T2D_1	0.1330 None	0.0019
1000252	20194397 C	0.490	0.999	5755-07	5775-19	1.40E-71 None	0.0046 T2D_1	0.1545 None	0.0020
11049134	20196222 A	0.496	1.000	7.836-07	1.045-18	2.09E-70 None	0.0045 T2D_1	0.0575 None	0.0017
19549135	28199413 G	0.499	1.000	6.096-07	6245-19	1.81E-70 None	0.0045 T2D_1	0.1522 None	0.0019
rs1708302	28169677 C	0.499	0.999	6076-07	7.036-19	6.775-71 Nore	0.0045 T2D_1	0.1363 None	0.0017
191513272	28200097 C	0.499	0.999	5.925-07	7.365-19	6.60E-71 None	0.0046 T2D_1	0.1308 None	0.0017
ra637124	26200142 C	0.295	0.999	8.595-01	1.405-08	8.110-112 None	0.0055 None	0.0033 HEIGHT_3	0.0009
10552797	28205303 T	0.296	0.999	9265-01	1.395-08	4.502-112 None	0.0056 None	0.0033 HEIGHT_3	0.0400
m543511926	28207300 G	0.999	0.020	6.855-02	5275-01	4.23E-01 None	0.0056 None	0.0033 HEIGHT_3	0.0917
14520191	28210600 T	6.299	1.000	8.875-01	6.605-09	1.915-112 None	0.0052 None	0.0034 HEIGHT_3	0.5365
15508347	28212804 T	0.299	0.999	8.14E-01	1.11E-08	3.535-112 None	0.0050 None	0.0033 HEIGHT_3	0.3277



JAZF1

- · JAZF1 encodes a protein with three zinc fingers and acts as a transcriptional repressor.
- It is member of a chaperone complex that orchestrates acetylation at regulatory regions controlling the expression of many genes involved in ribosome biogenesis.
   Work on the Jazf1 knockout mouse induced pluripotent stem cells suggests JAZF1 is involved in differentiation of β-cells and glucose homeostasis.

- JAZF1 appears to limit inflammation in adipose tissue and mice overexpressing JAZF1 have lower body and fat weight. In mouse airway epithelial cultures, JAZF1 expression was shown to be necessary for multicilisted cell differentiation, which is important for removing contaminants from the airway.
- · These functional studies suggest the plausibility of the role of JAZF1 in asthma and T2D, but do not suggest a genetic link between these phenotypes.

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## Previous associations with JAZF1 · Previous studies have found variants within JAZF1 to be associated

- separately with T2D, obesity phenotypes, as well as, height.
- · These findings include at least one study that reports a significant association with SNPs in JAZF1 with WC adjusted for BMI. Our findings also suggest that previous associations with SNPs in JAZF1 with WC adjusted for BMI are likely due to the same collider bias we observed, and the variants are associated with height, not adiposity.
- · There is evidence JAZF1 is associated with child-onset and possibly adultonset asthma

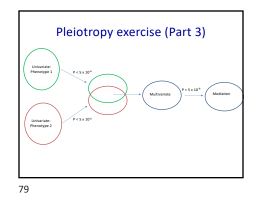
#### Conclusions

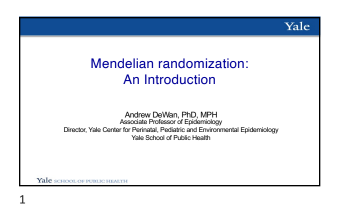
- While previous studies have identified associations with variants in JAZF1 associated with some aspect of all three phenotypes, this is the first time that asthma, T2D, and antirropometric measurements have been analyzed simultaneously in the same dataset and the first attempt at dissecting whether three are overlapping causal variants and/or biological pathways for these phenotypes.
- This study provides the strongest evidence for an association of variants in JAZF1 with asthma compared to previous studies.
- Variants in JAZF1 are associated with asthma, type 2 diabetes and height which provides a promising link between these three phenotypes, but the fine-mapped variant(s) for asthma, type 2 diabetes and height are unique.
- These results are consistent with biological pleiotropy at the gene-level for all three phenotypes.

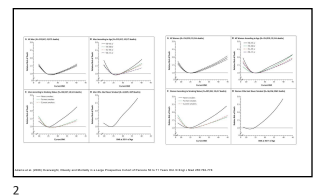
Mounting evidence that pleiotropy is more common at the gene-level (different causal variants) rather than at the variant level (shared causal

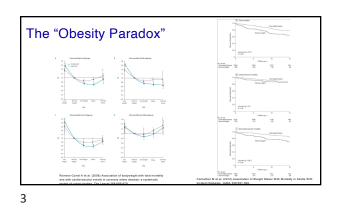
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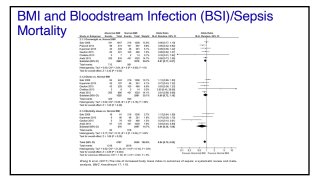
78

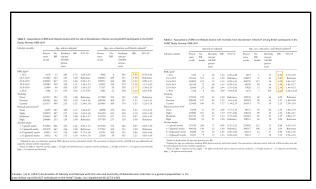


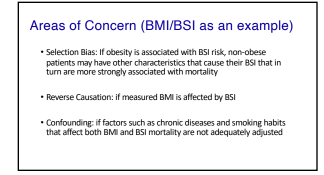








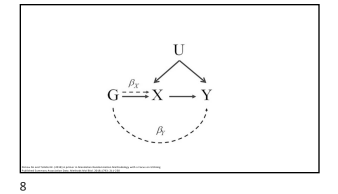




### Mendelian randomization

- Mimic randomized trial using genetic data as instruments for exposures
- Leverages information on genetic variants that segregate randomly at conception
- If an association between the instrument and outcome is detected, a causal relationship for this association is strengthened

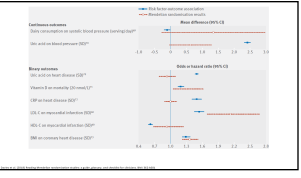
7



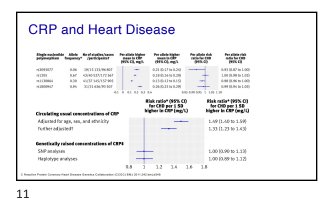
## **MR** Assumptions

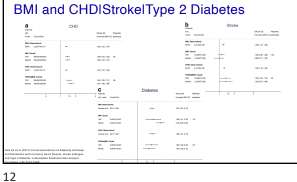
- The genetic instrument (G) is associated with the exposure (X)
- The genetic instrument is not associated with any confounder (U) of the exposure-outcome association
- The genetic instrument is conditionally independent of the outcome (Y) given the exposure and confounders











## One-sample vs. two-sample designs

#### One-sample

- Genotype(s), risk factor and outcome all measured in the same set of study subjects
- Individual level data must be available

 Two-sample
 Genotype(s) and risk factor measured in one set of study subjects and genotype(s) and outcome measured in a separate set of study subjects

 Can use summary statistics or individual level data

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## One-sample vs. two-sample designs

Assumption/Issue	One-sample	Two-sample
Instrument variable related to risk factor	Weak instrument biases towards the confounded regression result	Weak instrument biases towards the null
Confounders	Can (and should) check this for measured confounders	Not often possible when using summary statistics
Pleiotropy	Multiple methods to explore this issue (including MR-Egger)	Multiple methods to explore this issue (including MR-Egger) and may be more powerful with large consortium datasets since methods tend to be statistically inefficient
Subgroup analyses	Possible if large sample sizes and data on relevant risk factors are available	Only possible if individual level data are available
Bias from adjustments made in GWAS	N/A as all adjustments made in the same set of subjects	Summary data may or may not have been adjusted

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## Selecting genetic variants for an instrument

- Single or multiple variants
- Current recommendation is to select variant(s) that are significantly associated with the exposure at the genome-wide level
- Want a strong genetic instrument to avoid weak instrument bias
   A single variant or variants with modest effects in small samples are likely to have low power and can suffer from bias
- If selecting multiple variants these should not be in LD and assumes negligible gene-gene interaction among variants

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## Instrument strength

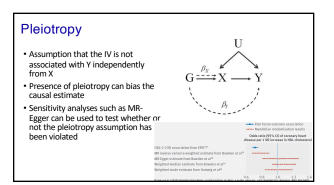
Measured using the F statistic in the regression of the IV on the exposure

$$F = \frac{N-K-1}{K} * \frac{R^2}{1-R^2}$$

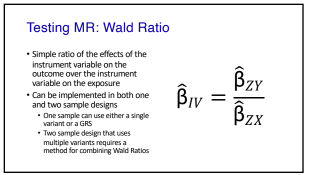
R<sup>2</sup>: proportion of the variance of the exposure explained by IV N: sample size K: number of genetic variants

General Rule: F < 10 is an indication of a weak instrument

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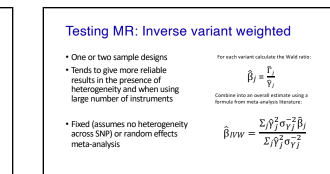
## Testing MR: 2 stage least squares (2SLS)

- Single continuous instrument (GRS)
- Only for one sample methodAssumes a linear relationship
- between exposure and outcome
- Calculate genetically predicted values of X
  Regress Y on genetically

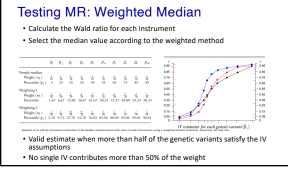
• Regress X on G

- predicted values of X
- Fix the standard errors (e.g. sandwich estimator)

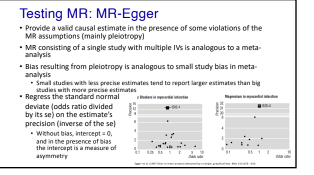
19

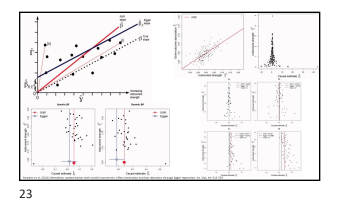


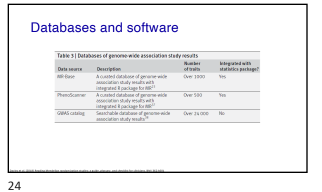
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## Body mass index and risk of dying from a bloodstream infection: A Mendelian randomization study

Tormod Rognes<sup>1,2,3</sup>\*, Erik Solligård<sup>1,3</sup>, Stephen Burgesse<sup>4,8</sup>, Ben M. Brumpton<sup>6,7,8</sup>, Julie Paulsen<sup>9</sup>, Hallie C. Prescottig<sup>1011</sup>, Randi M. Mohusg<sup>1,3</sup>, Lise T. Gustad<sup>0,12</sup>, Arne Mehl<sup>2</sup>, Bjørn O. Asvold<sup>6,13</sup>, Andrew T. DeWan<sup>1/24</sup>, Jan K. Damäsg<sup>1,14,194</sup> PLOS Medionie | <u>https://doi.org/10.1371/journal.gmed.1003413</u> November 16, 2020

Assess the causal association between BMI and risk of and mortality from BSI by overcoming the limitations of previous observational studies by conducting an MR study in a general population of approximately 56,000 participants in Norway with 23 years of follow-up

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# • The Trondelag Health Study (HUNT) is a series of cross-sectional surveys carried out in Nord-

- Trondelag County, Norway
   130,000 inhabitants who are representative of the general Norwegian population in terms of morbidity, mortality, sources of income and age distribution
- Based on HUNT2 survey conducted in 1995-1997 with 65,236 participants, 55,908 of whom had complete data for the analysis

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	Total population (n = 55,908)	BSI incidence (n = 2,547)	BSI death (n = 451)
Age (years) <sup>6</sup>	48.3 (36.5-62.3)	63.6 (52.9-71.4)	67.3 (57.1-74.5)
Male sex"	26,324 (47.1)	1,345 (52.8)	263 (58.3)
BMI (kg/m <sup>2</sup> ) <sup>3</sup>	26.3 (4.1)	27.7 (4.5)	27.9 (4.8)
Median follow-up time (years) <sup>5</sup>	21.1 (17.1-21.8)	13.8 (8.4-18.3)	13.3 (7.7-17.9)
Self-reported cancer*	1,955 (3.7)	144 (6.2)	24 (5.9)
Smoking*			
Never	23,594 (43.0)	876 (35.2)	156 (35.6)
Previous	15,133 (27.6)	893 (35.8)	164 (37.4)
Current	16,117 (29.4)	723 (29.0)	118 (26.9)
Physical activity*			
None	3,821 (7.6)	243 (11.9)	54 (15.4)
Slight	15,662 (31.0)	714 (34.9)	117 (33.3)
Moderate	17,167 (34.0)	693 (33.9)	116 (33.1)
High	13,810 (27.4)	397 (19.4)	64 (18.2)
Education*			
≤9 years	19,033 (35.7)	1,305 (55.8)	240 (58.8)
10-12 years	23,468 (44.0)	762 (32.6)	125 (30.6)
≥13 years	10,832 (20.3)	274 (11.7)	43 (10.5)

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## Outcome

- Linked to all prospectively recorded blood cultures at the two community hospitals in the catchment area (Levanger and Namsos Hospitals) as well as St. Olav's Hospital in Trondheim (tertiary referral center)
- Data on blood cultures were available from January 1, 1995 through the end of 2017
- Date of death and emigration out of Nord-Trondelag County were obtained from the Norwegian population registry
- BSI was defined as a positive blood culture of pathogenic bacteria
  BSI mortality was defined as death within 30 days of BSI diagnosis
- · bai mortaiity was defined as death within 50 days of bai diagnosis

28

## **Genetic Instrument**

- Based on a BMI meta-analysis of ~700,000 individuals
- 939 of 941 SNPs identified as associated with BMI (p<5x10<sup>-8</sup>, two SNPs did not pass imputation quality control)
- Genetic risk score (GRS) was calculated for BMI using the --score command in PLINK (version 1.9) and weighted based on the effect estimates from the meta-analysis
- GRS (939 variants) explained 4.2% of the variation in BMI in the population (F-statistic = 2,461)

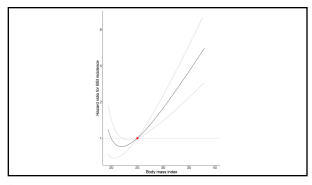
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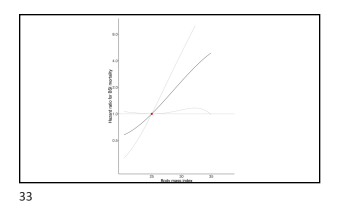
## Analysis Methods

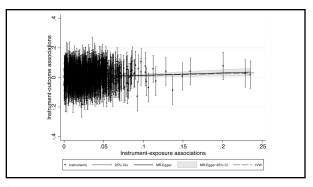
- Fractional polynomial model (suggestion of a nonlinear relationship between BMI and BSI)
- 2-stage least squares (with sandwich estimator) for analyses assuming a linear relationship between exposure and outcome
- Sensitivity analyses
- MR Egger (random effects)
- INW
- Weighted median
   Sample (using Yanga
- 2-sample (using Yengo et al. for SNP-exposure associations)

Characteristic	Total population (u = 55,988)	BSI incidence (n = 2,547)	881 death (w = 451)
Age (years) <sup>6</sup>	48.3 (36.5-62.3)	63.6 (52.9-71.4)	67.3 (57.1-74.5)
Male sex"	26,324 (47.1)	1,345 (52.8)	263 (58.3)
BMI (kg/m <sup>2</sup> )^	26.3 (4.1)	27.7 (4.5)	27.9 (4.8)
Median follow-up time (years)6	21.1 (17.1-21.8)	13.8 (8.4-18.3)	13.3 (7.7-17.9)
Self-reported cancer'	1,955 (3.7)	144 (6.2)	24 (5.9)
Sneking'			
Never	23,594 (43.0)	876 (35.2)	156 (35.6)
Previous	15,133 (27.6)	893 (35.8)	164 (37.4)
Current	16,117 (29.4)	723 (29.0)	118 (26.9)
Physical activity"			
Note	3,821 (7.6)	243 (11.9)	54 (15-4)
Slight	15,662 (31.0)	714 (34.9)	117 (33.3)
Moderate	17,167 (34.0)	493 (33.9)	116 (33.1)
High	13,810 (27,4)	397 (19.4)	64 (18.2)
Education*			
≤9 years	19,033 (35.7)	1,345 (55.8)	240 (58.8)
10-12 years	23,468 (44.0)	762 (32.6)	125 (30.6)
≥13 yaara	10,832 (20.3)	274 (11.7)	43 (10.5)
school, folk high school"), 10–12 years (' college, A levels'), and $\geq\!13$ years ('univ	currence, otherwise, last occurrence is us high school, intermediate school, vecatio entry or other post-secondary education, ght (~3 h light activity/week and no vig	nal school, 1–2 years high school" and less than 4 years" and "aniversity/colli	yars ("primary school 7-10 years, continuation "university qualifying essentiation, jurier get years or reace"). Activity defined as follows activity/week ce <1 h sigonas activity/week"),

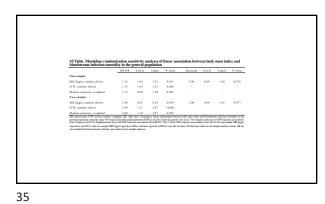




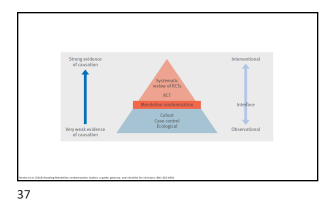


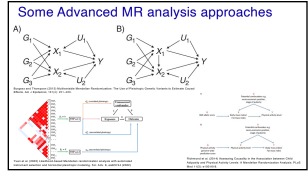


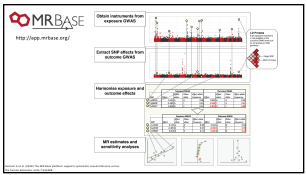


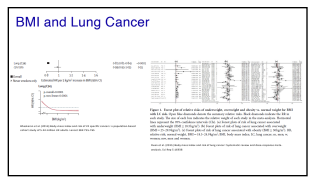


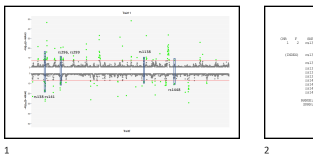


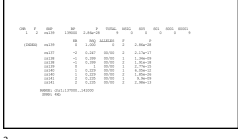


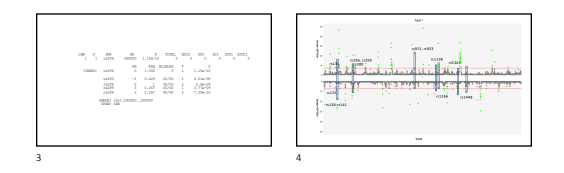


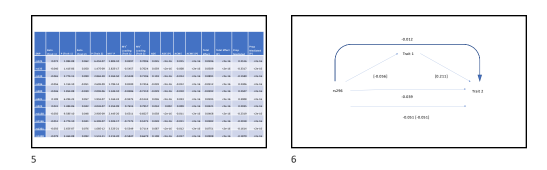


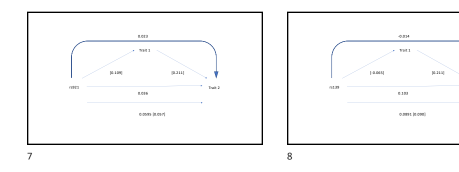
















Trait 2

0.032

0.011\* 0.043 [0.044] [0.211]

Trait 2

#### Discovery and Refinement of Loci Associated with Lipid Levels

A full list of authors and affiliations appears at the end of the article. # These authors contributed equally to this work.

#### Abstract

Abstract
Low-density lipoprotein (LDL) cholesterol, high-density lipoprotein (HDL) cholesterol,
triglycerides, and total cholesterol are heritable, modifiable, risk factors for coronary attery
disease. To is density me low is and refres hown hois influencein the lipolity, eventual 188,578
individuals using genome-wide and coston perosping arrays. We identify and annotate 157 keit
associated with high levels at *P* < 54x0<sup>-1</sup>, including 26 action typersoluty and annotate 157 keit
and African ancescup we narrow association signals hat 12 keit. We find that los associated with high
levels in humans. Using dense genotyping in individuals of European, East Akana, Soath Akana,
and African ancescup we marrow association signals hat 12 keit. We find that los associated with high
levels in humans. Using dense genotyping in individuals of European, East Akana, Soath Akana,
di African ancescula we narrow association signals hat 12 keit. We find that los associated with
genotype 2 diabetes, hode greestere, wate hop massis, and hody mass index. Our result
liberate the value of genotic data from individuals of diarrogenetic meetries and provide insights into the
biological mechanism regulating blood lipids to guide future genetic, hislogical, and therapeutic
research. research.

#### A comprehensive 1000 Genomes-based genome-wide association meta-analysis of coronary artery disease A full list of authors and affiliations appears at the end of the article.

# These authors contributed equally to this work.

#### Abstract

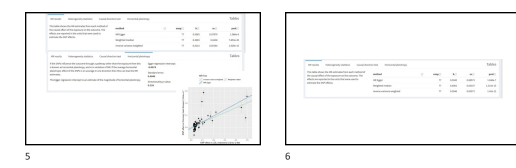
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**District** Existing knowledge of genetic variants affecting risk of coronary antery disease (CAD) is largely based on genome-wide association studies (GWAS) analysis of common SNPs. Leverging phased hapdroyses from He 1000 Genomes Projects ver eyrot a GWAS meta-malysis of 15 bonosand CAD cases and controls, interrogating 6.7 million common GMAPoD53 and well as 2.7 million requeres; (1005-844)-0153 virtualis. Landiton to confirmation of most known CAD) loci, we identified 10 novel (sci. eight additive and two recessive, that contain candidate genes that newly implicate biological processes in vessel value). We observed intro-iona allelic heterogeneity (bH inter evidence of low frequency virtuas with larger effects and ne evidence of symbolic dowing that genetic susceptibility to this common disease is largely dotermined by common SNPs of small effect size.

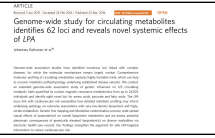
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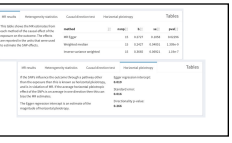


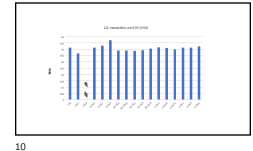




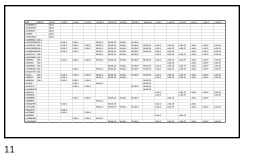


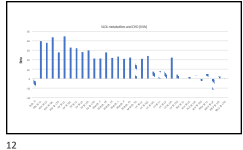












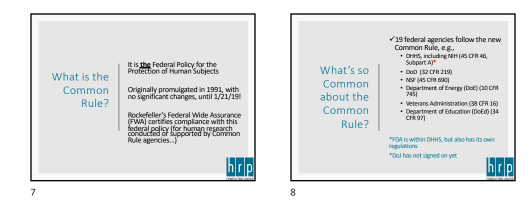


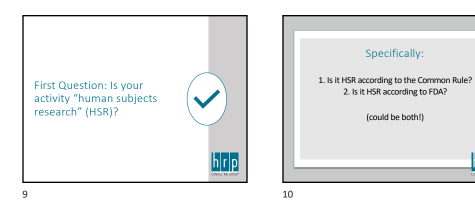


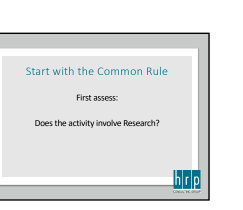








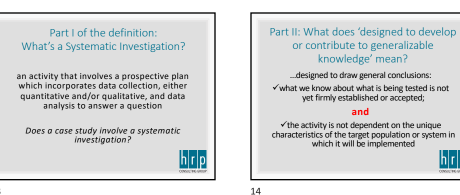






(could be both!)

h r p



### If the activity IS research: Does the research involve human subjects, according to the Common Rule? A living individual about whom an investigator conducting research: Results only to be applied to populations, or inform practice (i) Obtains information or biospecimens through intervention or interaction with the individual, and uses, studies, or within the target population or within the site where the activity analyzes the information or biospecimens; or (ii) Obtains, uses, studies, analyzes, or generates identifiable Implementation and evaluation of an evidence-based practice, private information or identifiable biospecimens. process, or program (is it functioning as intended within the site where the activity is being conducted or with the local target h|r|p h|r|p

15

is being conducted

population

16



An activity is not likely to be

generalizable if the intent is:

The evaluation or improvement of a process, practice, or program at the site where the activity is being conducted

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**FDA** Decisions

h|r|p

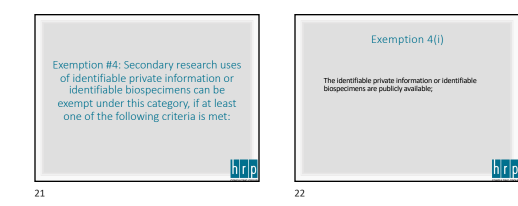
Does the activity evaluate an FDA-regulated test article (i.e., drug, biologic, device)?

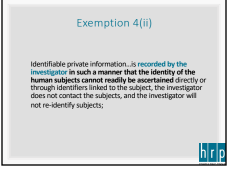
Does the activity involve Human Subjects? An individual who is, or becomes, a participant in research, either as a recipient of the test article or as a control. A subject may be either a healthy human or a patient. Also included in the FDA human subject definition: The use of a biological specimen –even if de-identified-from an individual used to test an investigational device

Does the activity involve research (clinical investigation)? Any experiment that involves a test article and one or more human subjects... h|r|p









## Exemption 4 (iii)

"The research involves only information collection and analysis involving the investigator's use of identifiable health information when that use is regulated under 45 CFR parts 160 AND 164, subparts A and E [HIPAA], for the purposes of "health care operations" or "research" as those terms are defined at 45 CFR 164.501 or "public health activities and purposes" as described under 45 CFR 164.512(b)"

h r p

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## Exemption 4 (iv)

The research is conducted by, or on behalf of, a Federal department or agency using government-generated or government-collected information obtained for non-research activities, if the research generates identifiable private information that is or will be maintained on information technology that is subject to and in compliance with section 208(b) of the E-Government Act of 2002, 441.U.S.C. 3501.une, if all of the identifiable private information collected, used, or generated as part of the activity will be maintained in systems of records subject to the Privacy Act of 1974, <u>5.U.S.C. 552a</u>, and, if applicable, the information used in the research was collected subject to the Paperwork Reduction Act of 1995, <u>44.U.S.C. 3501</u> *et seq.* 

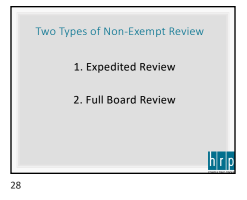


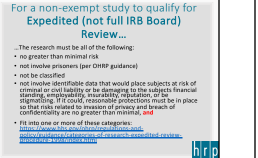
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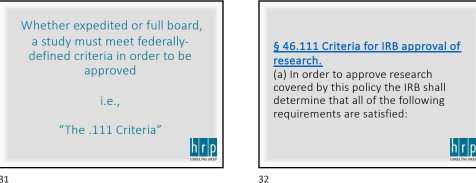












33



2. Risks to subjects are reasonable in relation to anticipated benefits, if any, to subjects, and the importance of the knowledge that may reasonably be expected to result

## 3. Selection of Subjects is Equitable Consider:

- The setting in which the research will be conducted
- Who is included, who is excluded? Does it make scientific sense? Ethical sense?
- If applicable: Are children in a study involving a test article that hasn't first been tested in adults? Pregnant women before non-pregnant women?
- Costs or compensation that may impact 'fairness'
- Screening and recruitment?
- What about non-English speakers?

4. Informed consent will be sought from each prospective subject or the subject's legally authorized representative, in accordance with, and to the extent required by, §46.116

## If not:

Are **ALL** the criteria for waiving informed consent or for altering/excluding specific elements of informed consent met?

h|r|p

h r p

h r p

## 5. Informed consent will be appropriately documented or appropriately waived in accordance with §46.117

### If not:

Does the research meet one of the allowable criteria to waive documentation?



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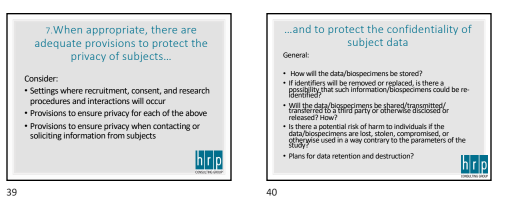
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## 6. When appropriate, the research plan makes adequate provision for monitoring the data collected to ensure the safety of subjects

- What data will be monitored for safety purposes? When? How?
- Who will be responsible for evaluating safety data? Is a DSMB needed?
- Stopping Rules?
- · Communication plan of findings to investigators and IRBs (from the IRB of Record or Sponsor)

h|r|p

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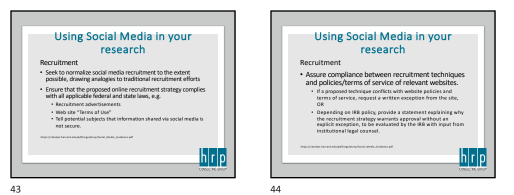
## Transporting or transmitting the data If any research data will be collected on a mobile device, such as an

electronic tablet, cell phone, or wireless activity tracker, details are needed regarding the physical security of the device, electronic security, and how the transfer of data from device to research storage location will be securely accomplished. accompliance. If any research data will be directly entered/sent by subjects over the internet or via email, will a University-provided database application (like REDCap) be used, or is there an encrypted tunnel to the site/application?

## Access to the data How will the investigators ensure only approved research personnel have access to the stored research data? Password-protected files, role-based security, etc.?

 Sharing of the data Will data be transferred or disclosed to or from the University? Is a contract
or data transfer agreement necessary? What (if any) identifiers will be
included? How will the data be securely transferred or disclosed (Universityapproved secure file transfer, etc.)? h r p

42

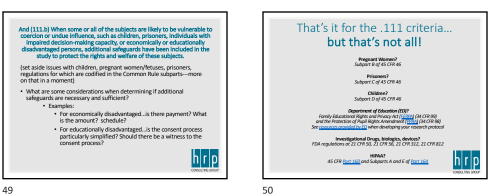






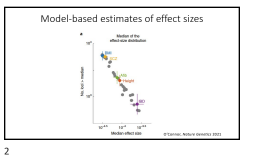
Using Social Media in your research Data source • A key issue in observational research using social media is whether the proposed project meets the criteria as human subjects research, and if so, what type of review is needed • Identifiable/de-identified data • Minimal risk/greater than minimal risk

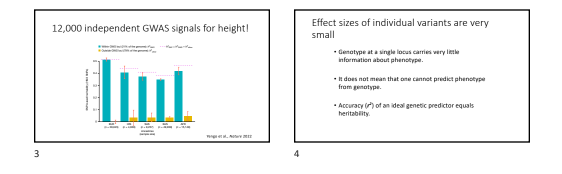






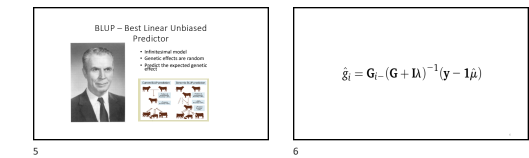


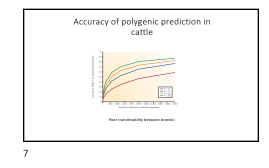


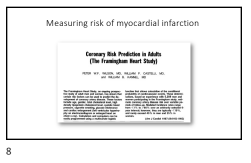


$$R_{GWS-AFR}^2$$
  
 $R_{GWS-AFR}^2$   $R_{HM3}^2$ 

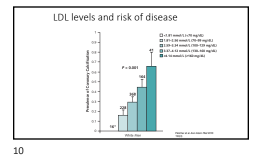
 $R_{\rm HM3}^2$ 

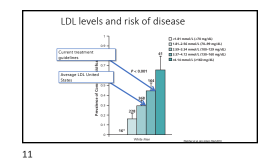




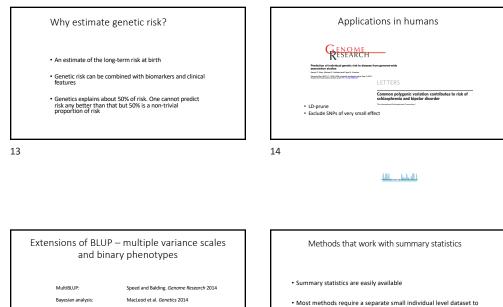














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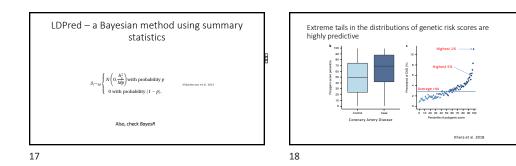
GeRSI:

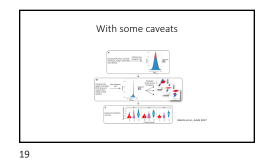
Zhou et al. PLOS Genetics 2013

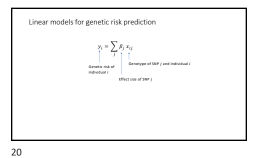
Golan and Rossett. AJHG 2014

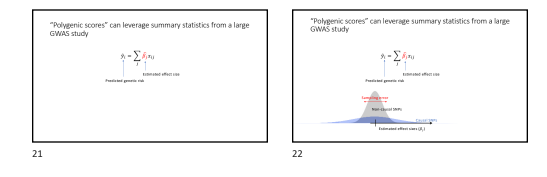


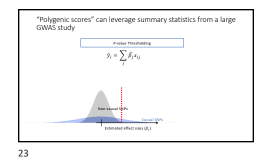


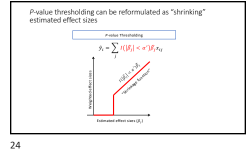


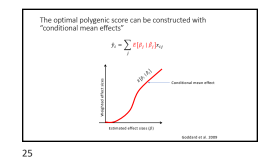


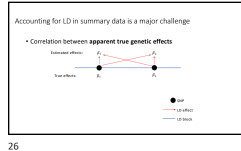


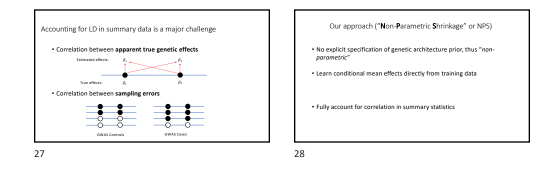


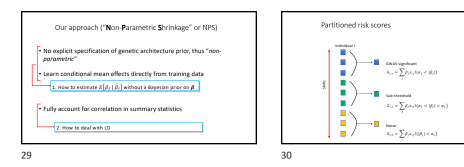


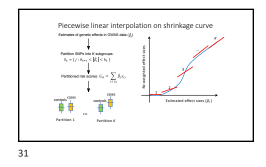


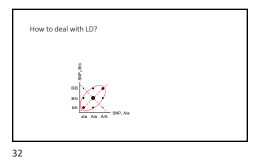




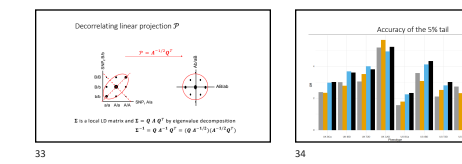


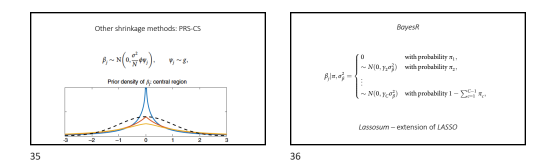


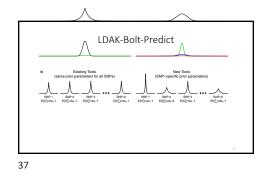


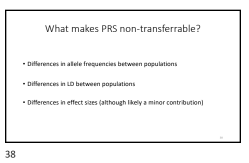


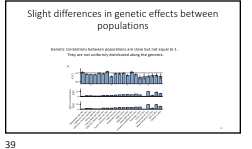
Chun et al. AJHG 202

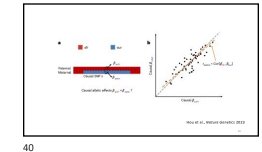


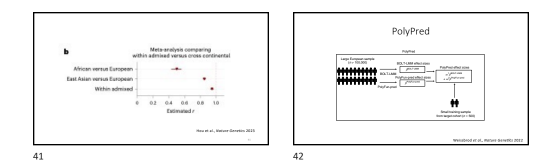






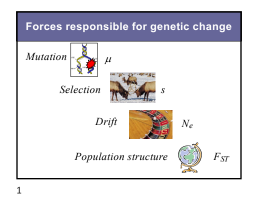


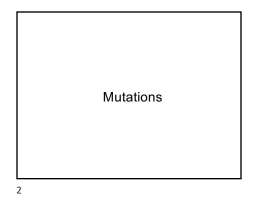


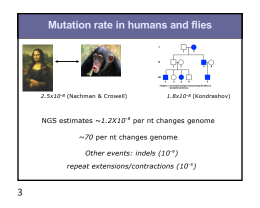


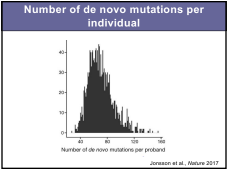


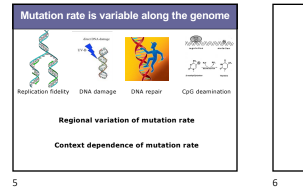


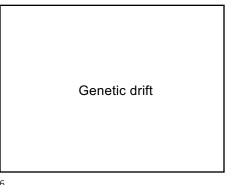


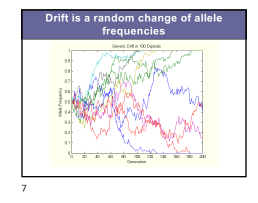


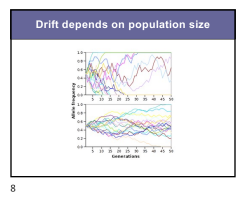






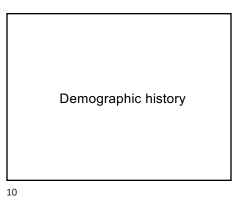


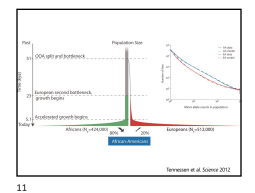


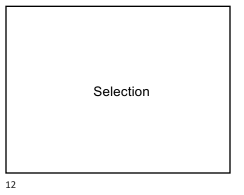


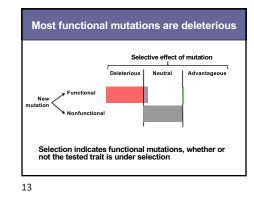
## Effective population size

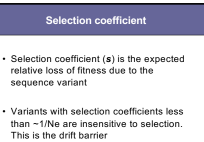
- In an idealized model, the intensity of genetic drift depends on population size (mean squared change in allele frequency is proportional to 1/Ne)
- In more realistic situations, effective population size (Ne) is a parameter characterizing intensity of drift

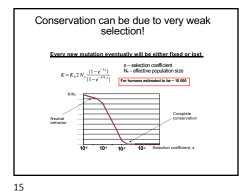






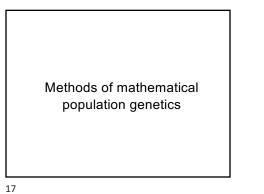


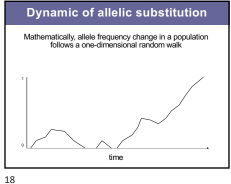


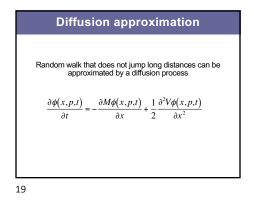


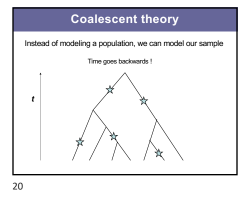
## Basic facts about human genetic variation

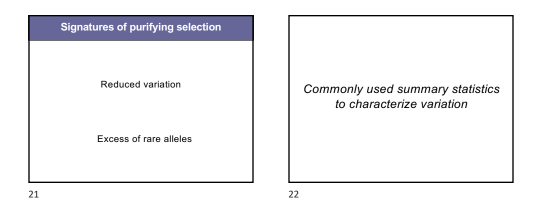
- Nucleotide diversity (density of nucleotide differences between two randomly chosen chromosomes) is about 0.001
- Most common SNPs are very old (~300-400K years old)
- Protein coding regions are showing clear signs of selection (reduced diversity and excess of rare alleles)

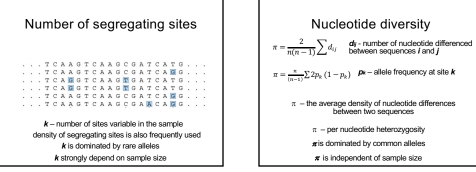


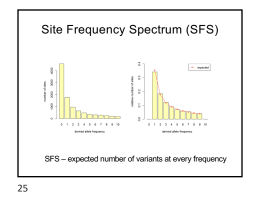


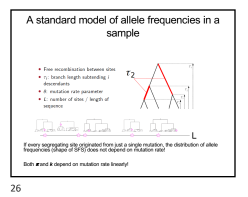


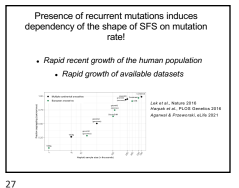




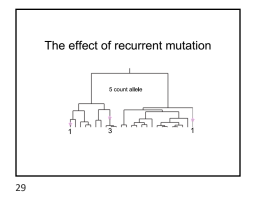


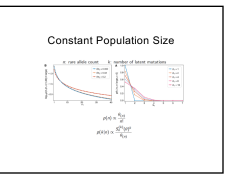


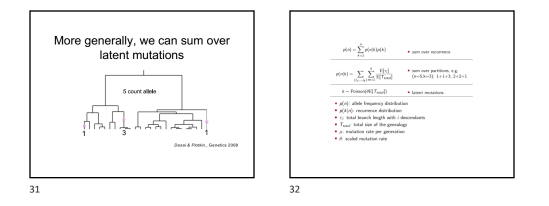


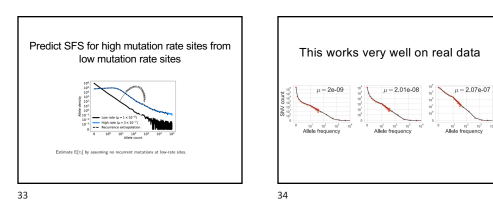




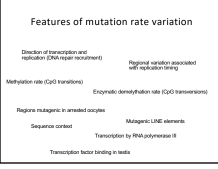












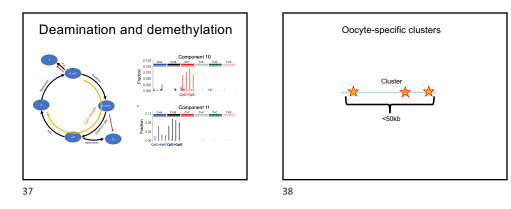
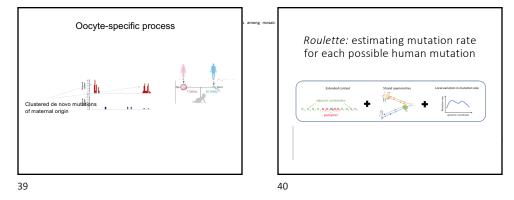
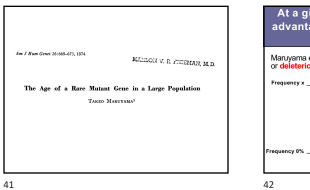
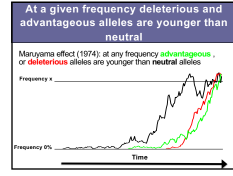
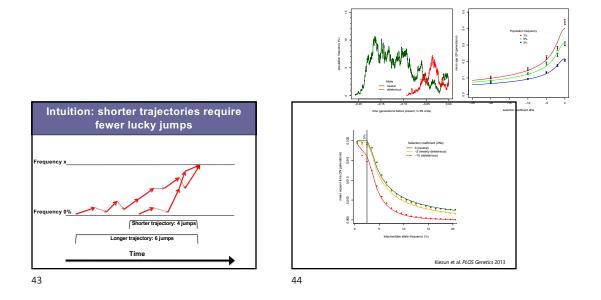


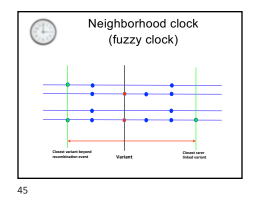
Fig. 4. Cytosine deamination and cytosine demethylation. (A and C) Spectra of components 10 and 11. (B, D) The intensity

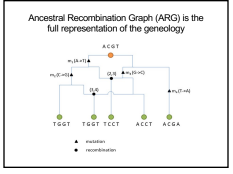


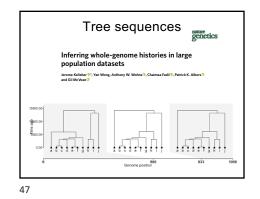


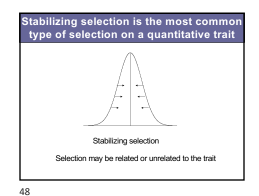


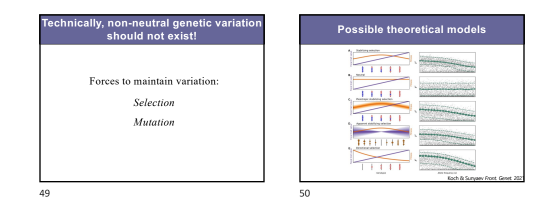


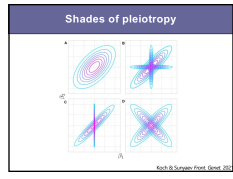




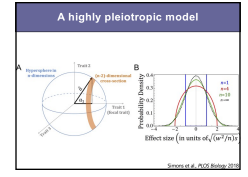












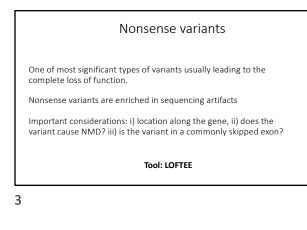
# Functional annotation of genes and variants

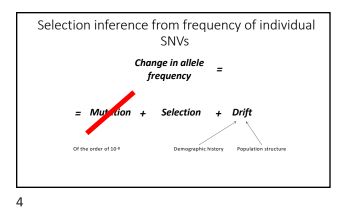
Map variants onto genomic annotation

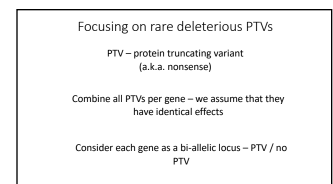
Watch for multiple transcripts!

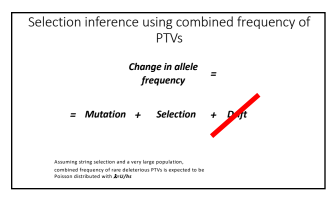
Watch for conflicting annotations!

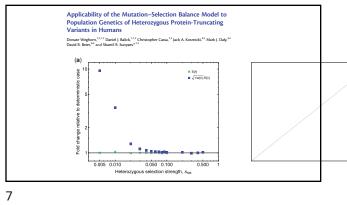
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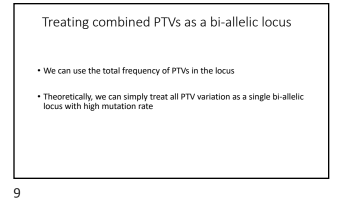


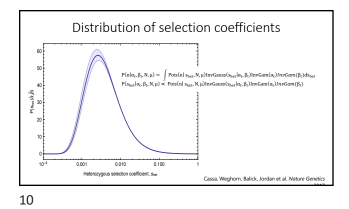


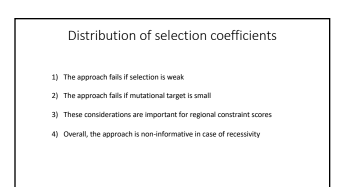


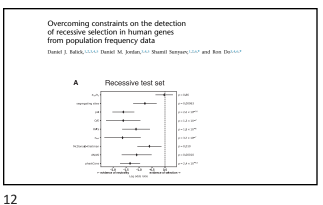
## Loss-of-function observed/expected upper bound fraction (LOEUF)

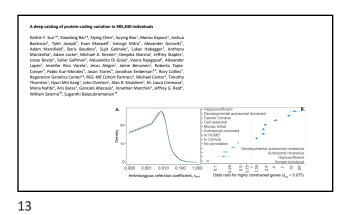
- LOEUF is based on the number if segregating sites as the statistic
- LEOUF assumes Poisson distribution for the number of segregating sites. It computes the expectation. The constraint metric is based on the Poisson likelihood ratio upper bound.

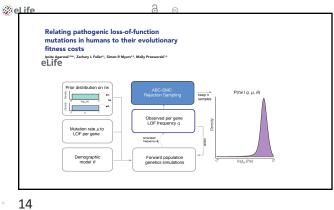


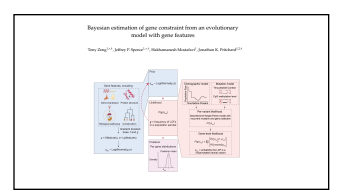


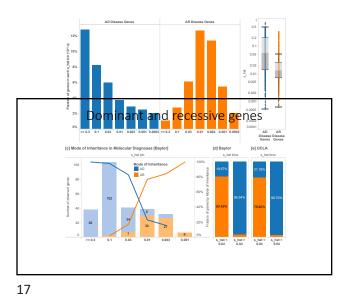


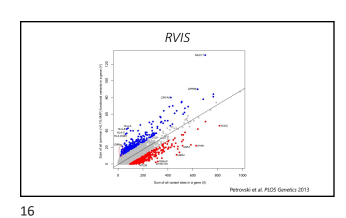


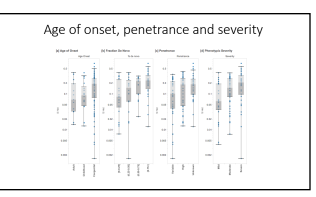


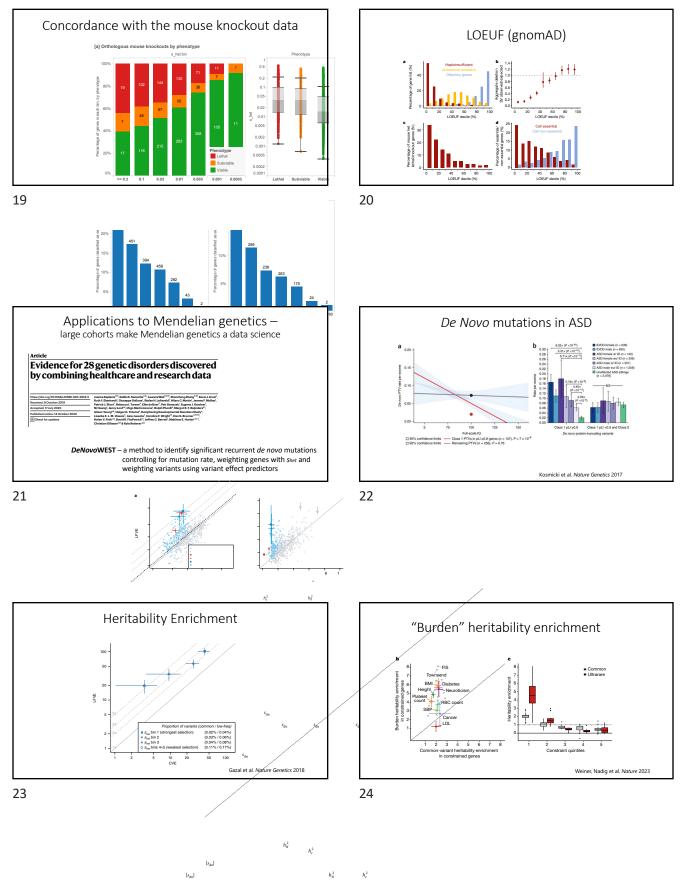


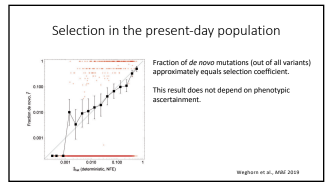




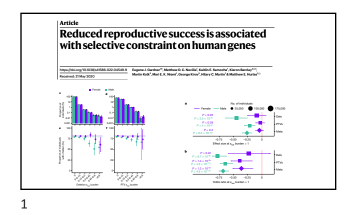


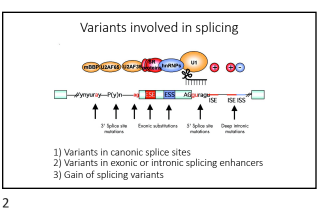




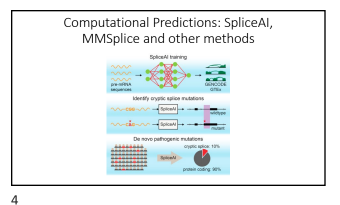


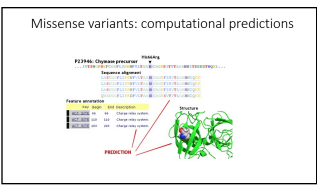


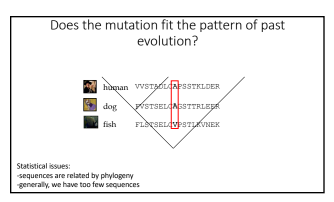




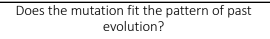
<figure><figure>



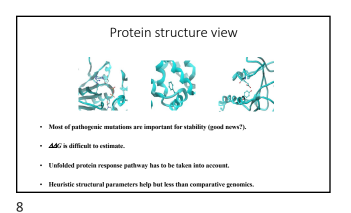


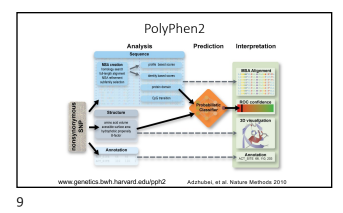




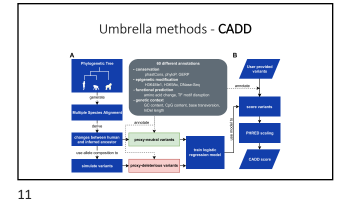


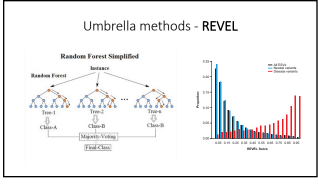
- We assume a constant fitness landscape: what is good for fish is good for human!
- We can estimate whether the mutation fits the pattern of amino acid changes.
- We can also estimate rate of evolution at the amino acid site

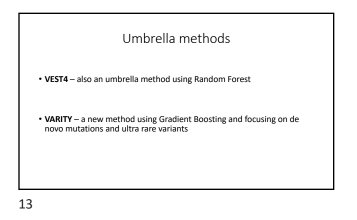




	<b>SIFT</b> is based on multiple sequence alignment
10	



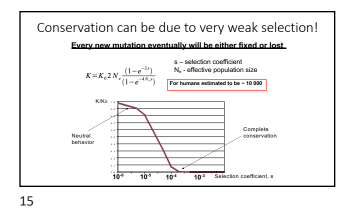


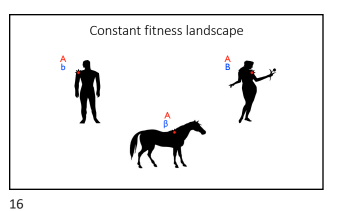


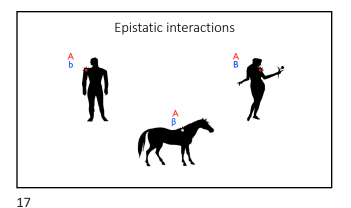
## Weakly deleterious mutations

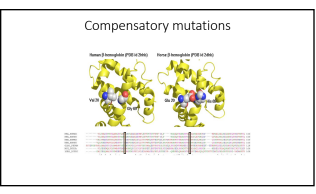
- Multiple independent lines of evidence suggest abundance of weakly deleterious alleles in humans
- · Weakly deleterious variants may occur in highly conserved positions
- Weakly deleterious alleles probably contribute to complex phenotypes but not to simple Mendelian phenotypes

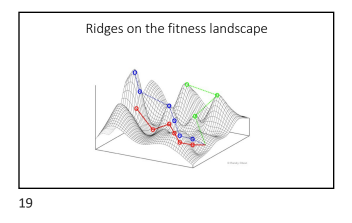
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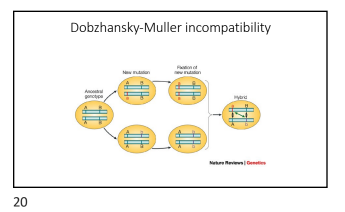


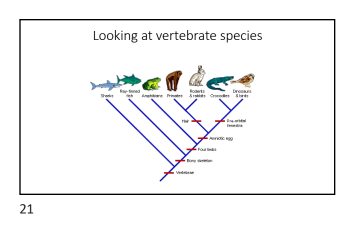


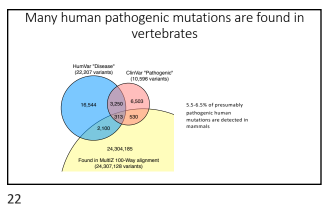


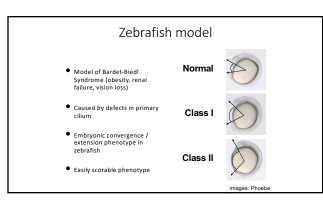


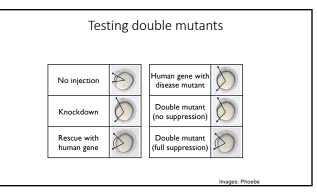


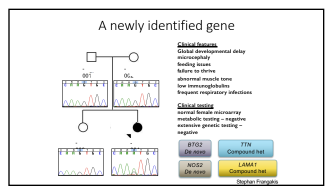








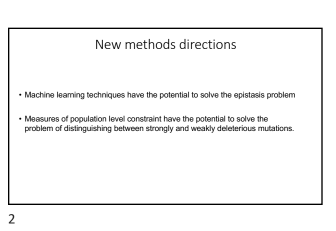




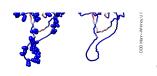


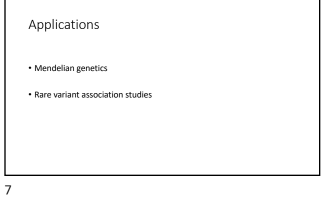


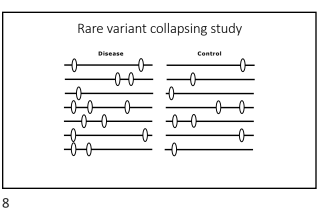
The mutation is a		evers estra			e m	amm	alian
BTG2	R80	L128	Q140	V141	L142		
H. sapiens P. troglodytes G. gorila M. musculus R. norvegicus H. glaber S. domesticus B. primigenius E. ferus caballus F. catus C. lupus familiaris D. novecnicctus G. gallus	<b>К • • × × × × × × × ×</b> ×	L • V V V V V V V V V V V V V V V V V V	Q • • • • • • • • •	V • MMMMMMMMMMMMMMMMMMMMMMMMMMMMMMMMMMM	L • M M M M M M M M M M		

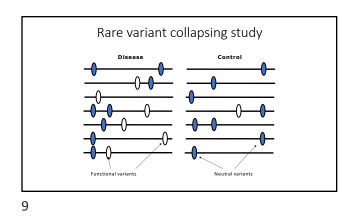


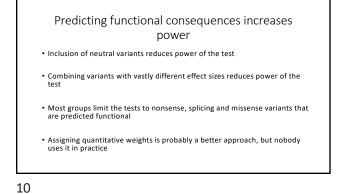
Large Language Models (VariPred) **EVE** – Variational Autoencoder \*\*\* ESM-1v ESM-2 ESM-1b 0.75 ŧ 0 € N 0.25 Evolutionary index •  $E_v \sim - \log \frac{P(x_v|\theta)}{P(x_{\rm WT}|\theta)}$ R AR ~ roximating the ve log-likelihood tant versus with er et al., Lin et el., bierxív 2023 Fra **№** 2021 3 4 1 BCINE EXE ESM ve PrimateAI-3D iteAl-3D 3D conv., (1,1,1) n. & ReLU M-G PHONE 3D conv., (3,3,3) & ReLU eat 3x m. & ReLU Dropout & sigmoid Loss fu Pathogenicity predictions for 20 amino acids 5 6

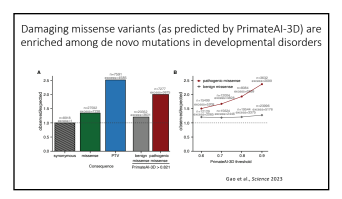


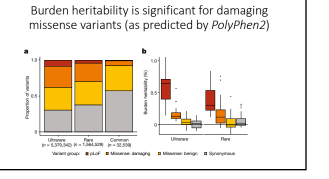




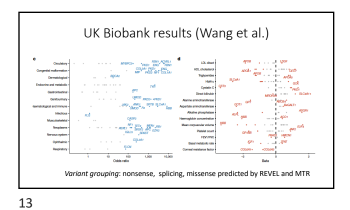


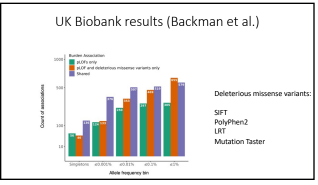




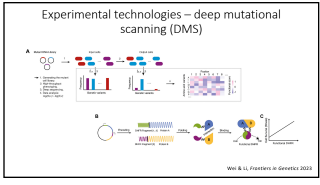




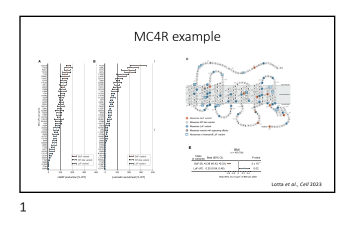


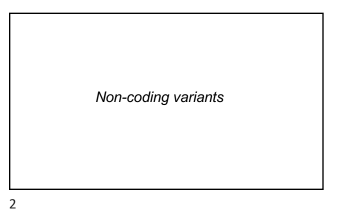


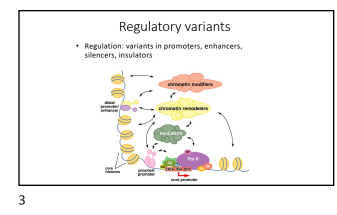


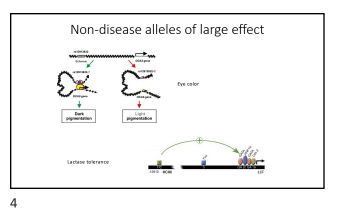


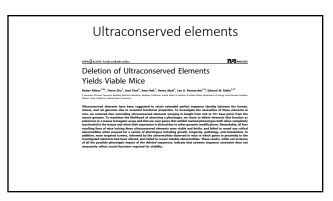


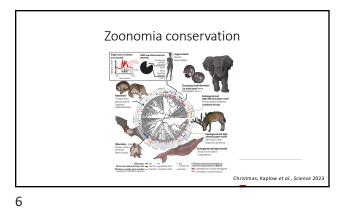


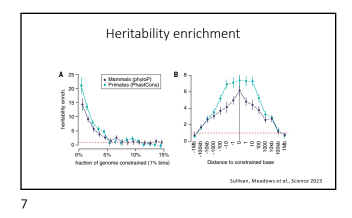


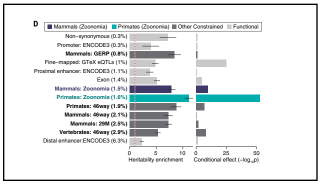


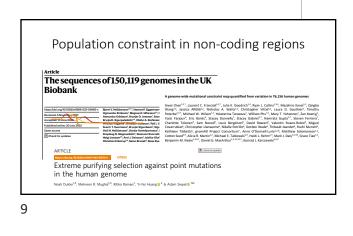


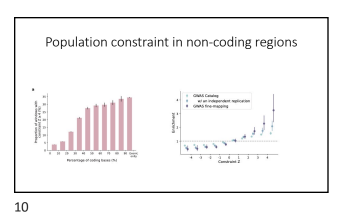


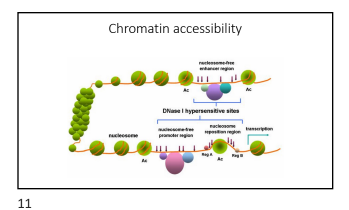


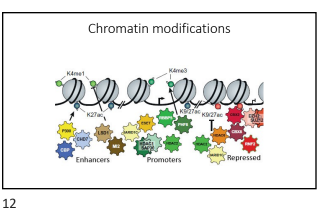


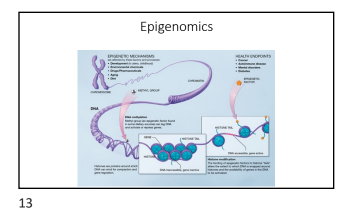


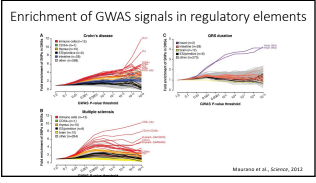




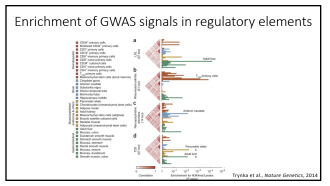




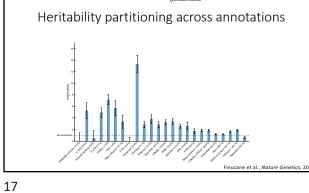


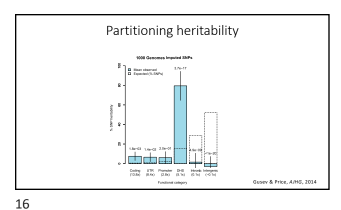




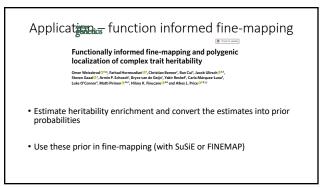


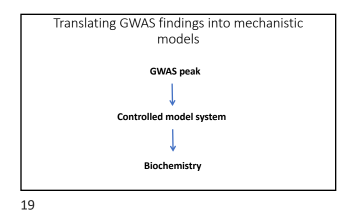


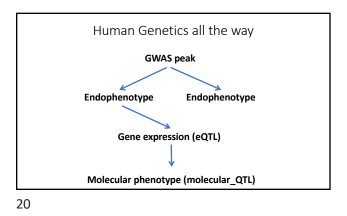


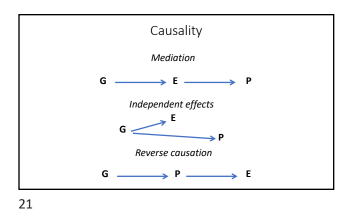


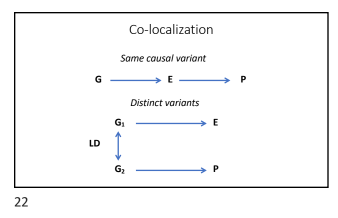


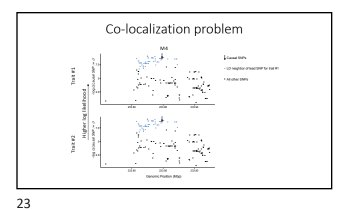


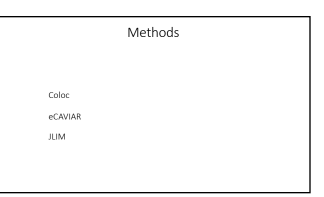








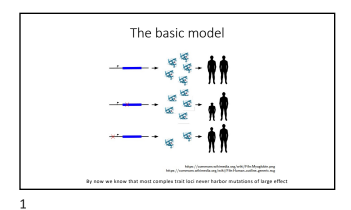




## Genetic variants differ between Mendelian and complex traits

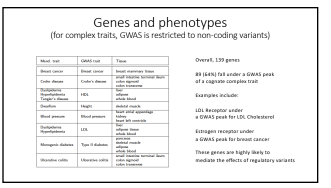
Complex trait variants

- Mendelian & somatic cancer variants
- Small effect size
- Large effect sizes
  Small number of loci
- Extremely large number of loci
   Mostly non-coding (regulatory)
   Small number of Mostly coding
  - Are in "putatively causative" genes

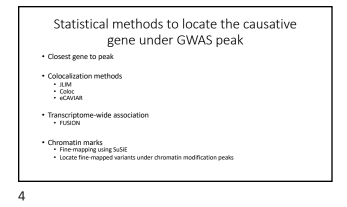


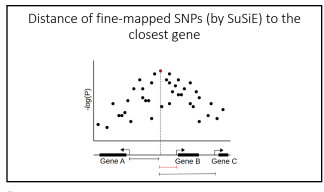


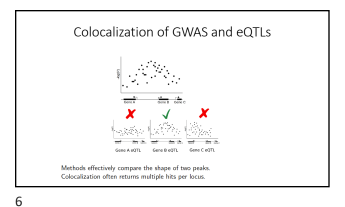
- Most genes involved in Mendelian components of complex traits are also causative for cognate common forms.
- Variants involved in common forms alter regulatory sequence of these genes.
- This in turn induces changes in gene expression; regulatory variants are *eQTLs*.

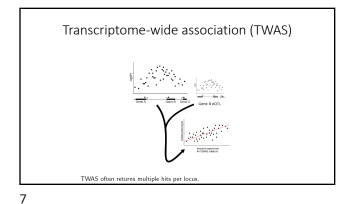


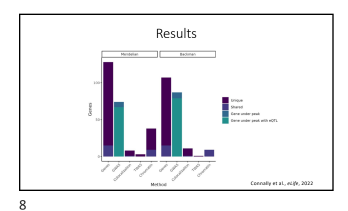


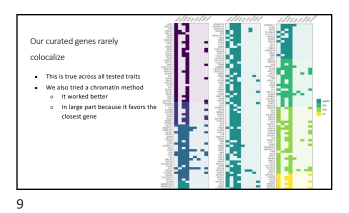


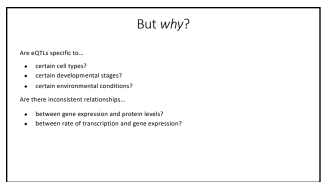


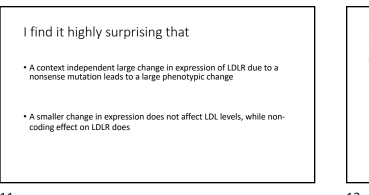


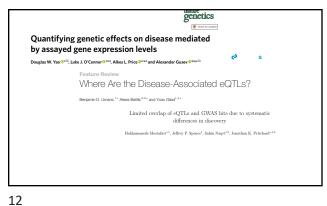












k  $$h_{\rm mel}$$   $$h_{\rm mel}$$   $$h_{\rm mel}$$   $$h_{\rm mel}$$ 

