# Lecture 3: GWAS in Samples with Structure & Introduction to the REGENIE Software

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#### Introduction

- Genetic association studies are widely used for the identification of genes that influence complex traits.
- To date, hundreds of thousands of individuals have been included in genome-wide association studies (GWAS) for the mapping of both dichotomous and quantitative traits.
- Large-scale genomic studies often have high-dimensional data consisting of
  - ► Tens of thousands of individuals
  - Genotypes data on a million (or more!) SNPs for all individuals in the study
  - Many phenotypes of interest such as Height, BMI, HDL cholesterol, blood pressure, diabetes, etc.

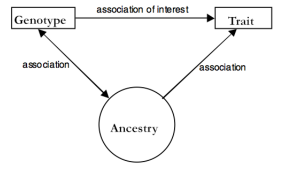
#### Introduction

- The vast majority of these studies have been conducted in populations of European ancestry
- Non-European populations have largely been underrepresented in genetic studies, despite often bearing a disproportionately high burden for some diseases.
- Recent genetic studies have investigated more diverse populations.

## **Confounding due to Ancestry**

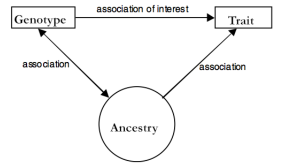
- ► The observations in association studies can be confounded by population structure
  - Population structure: the presence of subgroups in the population with ancestry differences
- Neglecting or not accounting for ancestry differences among sample individuals can lead to false positive or spurious associations!
- This is a serious concern for all genetic association studies.

## Confounding due to Ancestry



In statistics, a **confounding variable** is an extraneous variable in a statistical model that correlates with both the dependent variable and the independent variable.

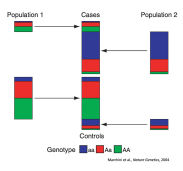
#### **Confounding due to Ancestry**



► Ethnic groups (and subgroups) often share distinct dietary habits and other lifestyle characteristics that leads to many traits of interest being correlated with ancestry and/or ethnicity.

## Spurious Association

- Association test aims to compare of allele frequency between cases and controls.
- Consider a sample from 2 populations:
  - No differences in allele frequencies between cases/controls within each population
  - Large differences in allele frequencies between populations
  - ▶ Population 2 is overrepresented among cases in the sample.
     ⇒ spurious association between disease and genetic marker



#### **Genomic Control**

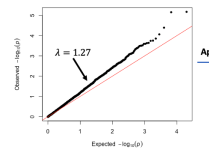
- Devlin and Roeder (1999) proposed correcting for substructure via a method called "genomic control."
- If there is no population structure, then at unlinked variants the test statistic  $T\sim\chi_1^2$  .
- If there is population structure, the statistic will deviate from a  $\chi_1^2$  distribution by an approximate constant factor  $T\sim \lambda\chi_1^2$  which is estimated as

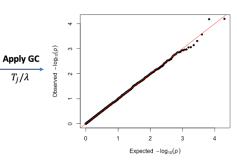
$$\lambda = \frac{\textit{median}(T)}{\textit{median}(\chi_1^2)} = \frac{\textit{median}(T)}{.456}$$

It is then applied to the test statistic values at all markers:

$$\tilde{T}_j = \frac{T_j}{\lambda}$$

#### **Genomic Control**





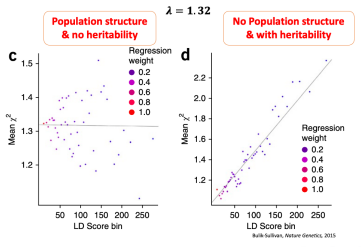
## LD Score Regression

- In practice,  $\lambda$  is computed using all variants
- ▶ Polygenicity can cause  $\lambda > 1$ 
  - $\blacktriangleright$  Hard to separate confounding from polygenicity when  $\lambda>1$
- ▶ LD score regression separates these by regressing "LD scores" L<sub>j</sub> on the test statistics

$$E[T_j] = \frac{Nh_g^2}{M} \cdot L_j + Na + 1$$

Slope → captures polygenicity Intecept → captures confounding

## **LD Score Regression**



## Correcting for Population Structure with PCA

- Principal Components Analysis (PCA) is the most widely used approach for identifying and adjusting for ancestry differences among sample individuals
- Consider the genetic relationship matrix  $\hat{\Psi}$  discussed in the previous lecture with components  $\hat{\psi}_{ij}$  for each pair of individuals as:

$$\hat{\psi}_{ij} = \frac{1}{M} \sum_{l=1}^{M} \frac{(G_{il} - 2\hat{\rho}_l)(X_{jl} - 2\hat{\rho}_l)}{\hat{\rho}_l(1 - \hat{\rho}_l)}$$

where  $G_{il} = \{0, 1, 2\}$  is the genotype value and  $\hat{p}_l$  is a corresponding allele frequency estimate at marker l

## Correcting for Population Structure with PCA

- Price et al. (2006) proposed correcting for structure in genetic association studies by applying PCA to  $\hat{\Psi}$ .
- ▶ They developed a method called EIGENSTRAT for association testing in structured populations where the top principal components (highest eigenvalues) are used as covariates in a linear regression model to correct for sample structure.

$$Y = \beta_0 + \beta_1 G + \beta_2 P C_1 + \beta_3 P C_2 + \beta_4 P C_3 + \dots + \epsilon$$
  
 $H_0: \beta_1 = 0 \text{ vs } H_a: \beta_1 \neq 0$ 

## Samples with Population Structure and Relatedness

- Relatedness (family structure or cryptic relatedness) in the sample can lead to spurious association in genetic association studies
- ► The EIGENSTRAT method was developed for unrelated samples with population structure
  - In the presence of relatedness, PCs may not fully capture this finer-scale structure
- Many genetic studies include relatedness & modeling it directly can lead to improvements in statistical power

# Association Testing in Samples with Population Structure and Relatedness

Linear mixed models (LMMs) have been demonstrated to be a flexible approach for association testing in structured samples. Consider the following model:

$$\mathbf{Y} = \mathbf{X}\boldsymbol{\beta} + \mathbf{G}_{\mathbf{s}}\gamma + \mathbf{g} + \boldsymbol{\epsilon}$$

- Fixed effects:
  - **X** is a  $n \times (k+1)$  matrix of covariates that includes an intercept
  - $\triangleright$   $\beta$  is the (k+1)-length vector of covariate effects
  - $ightharpoonup \gamma$  is the (scalar) association parameter of interest, measuring the effect of genotype on phenotype

#### **Linear Mixed Models for Genetic Association**

$$\mathbf{Y} = \mathbf{X}\boldsymbol{\beta} + \mathbf{G_s}\gamma + \mathbf{g} + \boldsymbol{\epsilon}$$

- Random effects:
  - **g** is a *n*-length vector of polygenic effects with  $\mathbf{g} \sim \mathcal{N}(\mathbf{0}, \sigma_g^2 \mathbf{\Psi})$ 
    - $\sigma_g^2$  represents additive genetic variance and  $\Psi$  is a  $n \times n$  matrix of pairwise measures of genetic relatedness (e.g. kinship matrix, GRM)
    - g should capture correlation between individuals due to genetic relatedness
  - $m{\epsilon}$  is a *n*-length vector with  $m{\epsilon} \sim N(\mathbf{0}, \sigma_e^2 \mathbf{I})$ 
    - $\sigma_e^2$  represents variance due to non-genetic effects assumed to be acting independently on individuals

#### LMM methods for Quantitative Traits

models for genome-wide

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association studies

#### TECHNICAL REPORTS TECHNICAL REPORTS nature genetics genetics Variance component model to account for sample Rapid variance components-based method for structure in genome-wide association studies whole-genome association analysis Hyun Min Kane<sup>1,2,8</sup>, Jae Hoon Sul<sup>3,8</sup>, Susan K Service<sup>4</sup>, Noah A Zaitlen<sup>5</sup>, Sit-vee Kone<sup>4</sup>, Nelson B Freimer<sup>4</sup>, Gulnara R Svishcheva<sup>1</sup>, Tatiana I Axenovich<sup>1</sup>, Nadezhda M Belonogova<sup>1</sup>, Cornelia M van Duiin<sup>2</sup> & Chiara Sabatti<sup>6</sup> & Eleazar Eskin<sup>1,5</sup> Yurii S Aulchenko<sup>1</sup> TECHNICAL REPORTS PLOS GENETICS OPEN & ACCESS Freely available online genetics Polygenic Modeling with Bayesian Sparse Linear Mixed Models Xiang Zhou1\*, Peter Carbonetto1, Matthew Stephens1,2\* 1 Department of Human Genetics University of Chicago, Chicago, Illinois, United States of America, 2 Department of Statistics, University of Chicago, Chicago, Illinois Genome-wide efficient mixed-model analysis for association studies Xiang Zhou1 & Matthew Stephens 1,2 TECHNICAL REPORTS BRIFF COMMUNICATIONS genetics FaST linear mixed Mixed linear model approach adapted for genome-wide

association studies

Zhiwu Zhang<sup>1</sup>, Ilhan Brooz<sup>1</sup>, Chao-Qiang Lai<sup>2</sup>, Rory J Todhunter<sup>3</sup>, Hemant K Tiwari<sup>4</sup>, Michael A Gore<sup>3</sup>, Peter J Bradbury<sup>6</sup>, Jiazming Yu<sup>2</sup>, Donna K Arnett<sup>4</sup>, Jose M Ordovas<sup>1,5</sup> & Edward S Backles<sup>1,6</sup>

## LMMs: Two Step Procedure

- Many LMM methods use a two-step procedure for GWAS
- Step 1 considers a null model without the tested SNP of interest (i.e.  $\gamma=0$ )

$$\mathbf{Y} = \mathbf{X}\boldsymbol{\beta} + \mathbf{g} + \boldsymbol{\epsilon}$$

- Obtain parameter estimates to get predictions for the polygenic effects g
- Same for all variants tested so only performed once which reduces the computational burden

#### LMMs: Two Step Procedure

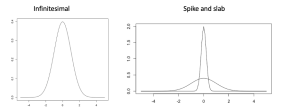
In Step 2, association testing of SNP and phenotype  $(H_0: \gamma = 0)$  is performed based on the model including the tested SNP

$$\mathbf{Y} = \mathbf{X}\boldsymbol{\beta} + \mathbf{G_s}\gamma + \mathbf{g} + \boldsymbol{\epsilon}$$

- ▶ A score test is performed using the null parameter estimates obtained from Step 1.
- Use Leave-One-Chromosome-Out (LOCO) scheme in Step 1 so polygenic term doesn't capture effects on tested chromosome (i.e. proximal contamination)

#### LMMs: Two Step Procedure

- Many methods differ mainly in Step 1 approach
  - Model used for the additive polygenic random effect term



- ► Algorithm used to obtain parameter estimates
  - Parameter estimates are obtained using various approaches (e.g. maximum likelihood, restricted maximum likelihood [REML],...)

#### LMMs on biobank scale data

- Largest biobanks have gathered data on 100,000s of individuals (e.g. UK Biobank at N = 500,000 individuals)
- Many LMM methods involved computationally expensive operations due to the  $N \times N$  GRM

Table 1 Computational cost of EMMAX, FaST-LMM, GEMMA, GRAMMAR-Gamma and GCTA

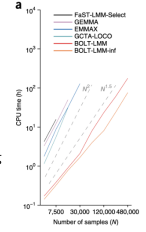
Method	<b>Building GRM</b>	Variance components	Association statistics
EMMAX	O(MN <sup>2</sup> )	O(N <sup>3</sup> )	O(MN <sup>2</sup> )
FaST-LMM <sup>a</sup>	$O(MN^2)$	O(N <sup>3</sup> )	$O(MN^2)$
GEMMA	$O(MN^2)$	O(N <sup>3</sup> )	$O(MN^2)$
GRAMMAR- Gamma	$O(MN^2)$	O(N <sup>3</sup> )	O(MN)
GCTA	$O(MN^2)$	$O(N^3)$	$O(MN^2)$

For each method, we list the computational cost of each step. alf M < N, the computational cost of FaST-LMM can be reduced to  $O(M^2 N)$ .

Yang et al., Nature Genetics 2014

#### LMMs on biobank scale data

- ▶ Loh et al. (2015) proposed BOLT-LMM which used very efficient algorithms (Variational Bayes) to reduce scaling to ~ O(MN<sup>1.5</sup>) for Step 1 and could be applied to biobank-scale data
- ▶ Jiang et al. (2019) proposed fastGWA which made use of a sparse GRM leading to further improvements for Step 1  $\sim O(MN)$



Loh et al., Nature Genetics 2015

## LMMs & Whole Genome Regression

▶ LMMs are closely related to whole genome regression

$$Y = W\beta + g + \epsilon$$
  $\Leftrightarrow$   $Y = W\beta + \sum_{l=1}^{M} G_l \theta_l + \epsilon$ 
 $N(0, \sigma_g^2 \Psi)$   $N(0, \sigma_g^2/M)$ 
 $\Psi = GG^T/M$ 
GRM using M variants

## LMMs & Whole Genome Regression

► LMMs are closely related to whole genome regression

#### 1 parameter

$$Y = W\beta + \frac{g}{g} + \epsilon$$

$$\uparrow$$

$$N(0, \sigma_g^2 \Psi)$$

$$\Psi = GG^T/M$$

GRM using M variants

#### **M** parameters

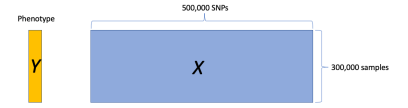
$$Y = W\beta + \sum_{l=1}^{M} G_l \frac{\theta_l}{\uparrow} + \epsilon$$

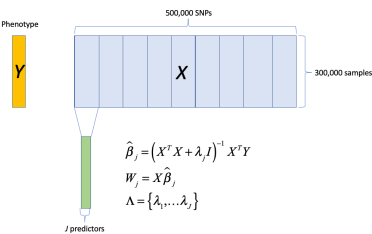
$$N(0, \sigma_g^2/M)$$

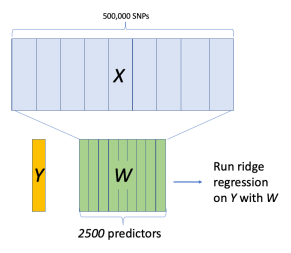
▶ Step 1: computationally efficient whole genome regression

$$\mathbf{Y} = \mathbf{X}\boldsymbol{\beta} + \sum_{l=1}^{M} G_l \theta_l + \epsilon$$

- ightharpoonup M is usually  $\sim$  500,000 SNPs across the genome
- REGENIE splits genetic data into blocks and run local regressions in each block to reduce computational burden







Step 1: computationally efficient whole genome regression

$$\mathbf{Y} = \mathbf{X}\boldsymbol{\beta} + \sum_{l=1}^{M} G_l \theta_l + \epsilon$$

- Divide into two levels of regressions
  - Reads genetic data in blocks and within each block fits ridge regression (penalized linear regression)
  - Fit another round of ridge regression on all the block predictors
- ▶ Polygenic predictions  $(\sum_{l=1}^{M} G_l \hat{\theta}_l)$  capture population structre, relatedness as well as polygenicity

Step 2: test the association parameter  $\gamma$  under the null hypothesis of  $H_0$ :  $\gamma = 0$ .

$$\mathbf{Y} = \mathbf{X}\boldsymbol{eta} + G_{s}\gamma + \sum_{l=1}^{M} G_{l}\hat{ heta}_{l} + \epsilon$$

- ► Test on millions of genetic variants (imputed or whole exome)
- Also works on binary traits where logistic regression is used instead of linear regression

https://rgcgithub.github.io/regenie/

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