Advanced statistical genetics methods

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Learning objective

- Population structures in genetics data
 - Admixture model
 - Linear mixed effect model
- Linkage disequilibrium
 - Rare variant burden tests
 - Fine-mapping causal variants
- ► GWAS summary statistics
 - ► Transcriptome-wide association studies
 - ► LD-score regression

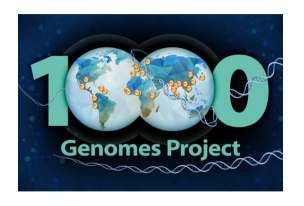
Today's lecture

Population structures in human genetics data

Linkage Disequilibrium: blessing and curse

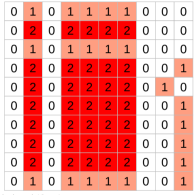
Systems genetics and summary statistics-based inference

The 1000 Genomes Project to investigate Human Genetic Variation



1KG contains whole genome sequencing data of 2,490 individuals sampled from 26 groups based on the origins and geographical locations (as of 2013 phase3).

Single Nucleotide Polymorphism (SNP) genotype information



first 10 individuals and 10 variants

Previously on the lecture 18:

- We will focus on biallelic variant (two allele, two different forms)
- Major and minor characters (depending on the frequency in reference data)
- We keep track of the number of the minor allele (0 to 2, due to diploid genome)

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▶ What is the mean of this variant?

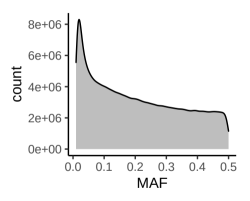
$$\widehat{\mathbb{E}}\big[X_j\big] = 2f_j$$

What is the variant of this variant?

$$\hat{\mathbb{V}}\big[X_j\big] = 2f_j(1 - f_j)$$

We can easily calculate MAF using bigsnpr

What is your interpretation?



Much of human genetics problems centre on two covariance matrices

For a standardized $n \times p$ genotype matrix X,

1. Genetic relatedness matrix (GRM)

a $n \times n$ matrix

$$K \approx XX^{\top}/n$$

The matrix K captures population structure/correlation across different individuals

- Kinship matrix; population admixture
- Human migration history

2. Linkage disequilibrium (LD)

a $p \times p$ matrix

$$R \approx X^{\top}X/n$$

The matrix R captures localized correlation patterns along the genomic axis within a chromosome.

- ▶ LD matrix
- The results of many, many recombination events

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What is this?

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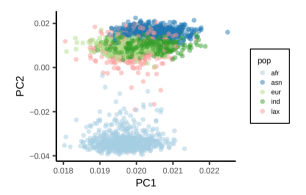
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$$\frac{1}{n}XX^\top = \frac{1}{n}UDV^\top VDU^\top = \frac{1}{n}UD^2U^\top$$
 sample × sample

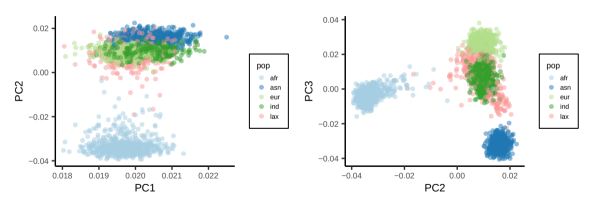
Let's take top 1000 most frequent variants

 ${\sf PCA} \ {\sf already} \ {\sf teaches} \ {\sf us} \ {\sf something} \ {\sf interesting}...$

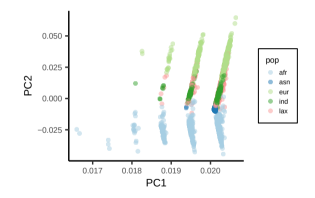


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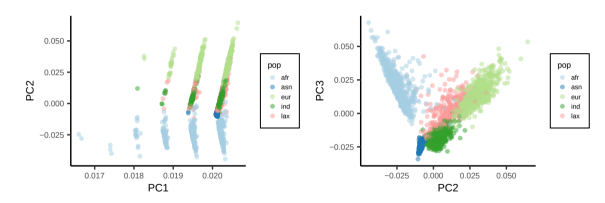
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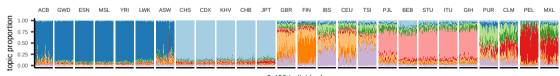
Estimating population admixture using top 10k variants

A potential generative model:

- lacksquare Sample each individual's topic proportion H_i
- Sample a topic membership for each variant j, say $Z_{ij} = k$ (could be implicitly handled)
- $lackbox{ Genotype } X_{ij}|Z_{ij}=k\sim ext{topic-specific } eta_{kj}$

$$\mathcal{L} = \prod_{i=1}^n \prod_{j=1}^{10\mathsf{k}} \left(\sum_{k=1}^9 H_{ik} \beta_{kj} \right)^{X_{ij}}$$

topic proportion H



2,490 individuals

Related work: Pritchard, Stephens, Donnelly, Genetics (2000)

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We can include some proxy random variables for population structures in a linear GWAS model

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- We can include some proxy random variables for population structures in a linear GWAS model
- We might still need to consider uncertainty of the random effects

(digression) Useful facts on multivariate Gaussian distribution - 1

If we have ${f y}$

$$\mathbf{y} \sim \mathcal{N}(\mu, \Sigma)$$

then

$$\mathbb{E}[U^{\top}\mathbf{y}] = U^{\top}\mu, \quad \mathbb{V}[U^{\top}\mathbf{y}] = U^{\top}\Sigma U$$

and (affine transformation)

$$U^{\top}\mathbf{y} \sim \mathcal{N}(U^{\top}\mu, U^{\top}\Sigma U)$$

(digression) Useful facts on multivariate Gaussian distribution - 2

If we have two Gaussian random vectors, $\mathbf{y} \sim \mathcal{N}\big(\mu + \mathbf{u}, \Sigma_y\big)$ and $\mathbf{u} \sim \mathcal{N}(\mathbf{u}|\mathbf{0}, \Sigma_u)$

Bayesian integration:

$$\int \mathcal{N}\big(\mathbf{y}|\mu + \mathbf{u}, \Sigma_y\big) \, \mathcal{N}(\mathbf{u}|\mathbf{0}, \Sigma_u) \, d\mathbf{u} = \mathcal{N}\big(\mathbf{y}|\mu, \Sigma_y + \Sigma_u\big)$$

A key idea in the proof:

$$\left[\Sigma_y^{-1} - \Sigma_y^{-1} \left(\Sigma_y^{-1} + \Sigma_u^{-1}\right)^{-1} \Sigma_y^{-1}\right]^{-1} = \Sigma_y + \Sigma_u$$

by Woodbury identity.

A linear model with population-driven random effects

A linear regression model:

$$\mathbf{y} = \mathbf{x}_j \qquad \beta_j \qquad + \epsilon$$
 a fixed genetic effect

What are we missing? Can we assume homo-scedasticity, i.e.,

$$\epsilon \stackrel{?}{\sim} \mathcal{N}(\mathbf{0}, \sigma^2 I)$$

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A linear model with a random effect:

$$\mathbf{y} = \mathbf{x}_j \, eta_j + \mathbf{u}_{\mathsf{random effect}} + \epsilon$$

Note: There is no specific parameterization for this $n\times 1$ random vector ${\bf u}$. Now, we assume:

$$\epsilon \sim \mathcal{N}(\mathbf{0}, \sigma^2 I)$$

A linear model with population-driven random effects - 2

We want to capture unwanted population, cohort-specific random effects by $n \times 1$ vector \mathbf{u} and **remove** since our **goal** is to estimate the fixed genetic effect of a particular variant j.

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- 1. Note that \mathbf{u} shouldn't be tied to a particular variant (by definition)
- 2. Also, the covariation of ${\bf u}$ is primarily driven by relatedness among individuals, not the variants.

$$\mathbf{u} \sim \mathcal{N}(\mathbf{0}, \tau^2 K), \quad K pprox rac{1}{n} X X^{ op}$$

A linear mixed effect model (LMM) to test associations while adjusting population structure

We can define a hierarchical model:

$$\mathbf{y}|X,\beta,\mathbf{u},\sigma \sim \mathcal{N}(X\beta+\mathbf{u},\sigma^2I)$$

$$\mathbf{u}|\tau,K \sim \mathcal{N}(\mathbf{0},\tau^2K)$$
(1)

If we integrate out \mathbf{u} ,

$$\mathbf{y}|X, eta \sim \mathcal{N}\left(\mathbf{y} \left| Xeta, \underbrace{ au^2 K}_{ ext{genetic-relatedness matrix}} + \underbrace{\sigma^2 I}_{ ext{irreducible}}
ight)$$

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- Cumbersome computation required for matrix factorization or other latent variable modelling on a large genotype matrix
- ▶ We many not have a large matrix to learn about non-genetic confounders...
- One LMM estimation can substitute multiple matrix factorization steps
- ▶ We may have a good idea about relationships induced by random effects!

We can resolve maximum likelihood estimate of the parameters, β, τ, σ ,

$$\max \log \mathcal{N}(\mathbf{y} | X\beta, \sigma_2 (\delta K + I))$$

where $au^2=\delta\sigma^2$.

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$$\max \log \mathcal{N}(\mathbf{y} \,|\, X\beta, \sigma_2 \,(\delta K + I))$$

where $\tau^2 = \delta \sigma^2$.

We need to deal with this unfriendly form of likelihood:

$$-\frac{1}{2}\left(n\log(2\pi\sigma^2) + \log|I+\delta K| + \frac{1}{\sigma^2}\left[\mathbf{y} - X\boldsymbol{\beta}\right]^\top (I+\delta K)^{-1}\left[\mathbf{y} - X\boldsymbol{\beta}\right]\right)$$

Instead, we can transform the underlying distribution using spectral decomposition of the genetic-relatedness matrix (GRM),

 $K = USU^{\top}$ where $U^{\top}U = I$, and S is a diagonal matrix.

$$U^{\top}\mathbf{y} \\ \text{projected output} \quad \sim \quad \mathcal{N}\bigg(U^{\top}X \\ \text{projected genotype} \beta, \sigma^2 U^{\top}(I+\delta K) U \bigg)$$

Lippert, Listgarten, ..., Heckerman, Nature Methods (2011)

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$$\begin{array}{ccc} U^{\top}\mathbf{y} & \sim & \mathcal{N}\bigg(\begin{matrix} U^{\top}X \\ \text{projected genotype} \end{matrix} \beta, \sigma^2 U^{\top}(I+\delta K) U \bigg) \\ \text{(by the affine transformation)} & \sim & \mathcal{N}\bigg(\begin{matrix} U^{\top}X \\ \text{projected genotype} \end{matrix} \beta, \sigma^2 (I+\delta S) \\ \text{diagonal matrix} \bigg) \end{array}$$

- \blacktriangleright We can find β by weighted least square
- \blacktriangleright We can find σ^2 and δ by fixing β

Lippert, Listgarten, ..., Heckerman, Nature Methods (2011)

A key research question in LMM: What covariance matrix?

If there were many types of random effects,

$$\mathbf{y} = X\beta + \mathbf{u} + \mathbf{w} + \dots + \epsilon$$
fixed random effects unknown

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If there were many types of random effects,

$$\mathbf{y} = X\beta + \mathbf{u} + \mathbf{w} + \ldots + \underset{\text{unknown}}{\epsilon}$$

We would need to many covariance matrices:

$$\mathbf{y}|\cdot \sim \mathcal{N}\left(X\beta,\, \sigma^2(I + \underbrace{\delta_u K_u + \delta_w K_w + \ldots}_{\text{random effects}})\right)$$

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If we only care about variance decomposition $\beta_i \sim \mathcal{N}(0, \tau)$:

$$\mathbf{y} \sim \mathcal{N} \Bigg(\mathbf{0}, \ \sigma^2 \left(\frac{\sigma_{\mathrm{genetic}}^2 X X^\top + I + \underbrace{\delta_u K_u + \delta_w K_w + \dots}_{\mathrm{random \ effects}} \right) \Bigg)$$

Should we worry about "over-fitting" in LMM?

An equivalent question for PCA-based confounder adjustment:

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An equivalent question for PCA-based confounder adjustment:

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For each chromosome $c\in\{1,\dots,22,{\rm X},{\rm Y}\}$, build a leave-one-chromosome-out (LOCO) kinship matrix, say K_{-c} :

$$\mathcal{N}\big(\mathbf{y}|\mathbf{x}_{j}\boldsymbol{\beta}_{j},\sigma^{2}(\delta K_{-c}+I)\big)$$

Yang, et al., Nature Genetics (2014)

Tucker, Price, Berger, Genetics (2014)

BSLMM: What will be a good prior for the effect variable in a LMM?

Bayesian Sparse LMM (BSLMM): Causal variants should have a higher, additional level of effect size (σ_a^2) than the background ones (σ_b^2) .

$$\beta_{j} \sim \pi \mathcal{N} \left(0, \frac{\sigma_{a}^{2} + \sigma_{b}^{2}}{p\tau} \right) + (1 - \pi) \mathcal{N} \left(0, \frac{\sigma_{b}^{2}}{p\tau} \right)$$

BVSR: spike-and-slab; BSLMM: mixture of two Gaussians; LMM: Gaussian

Zhou, Carbonetto, Stephens, PLoS Genetics (2014)

Today's lecture

Population structures in human genetics data

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Much of human genetics problems centre on two covariance matrices

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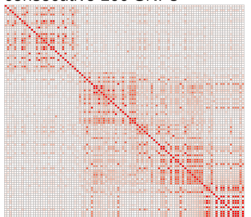
The matrix R captures localized correlation patterns along the genomic axis within a chromosome.

- ▶ LD matrix
- The results of many, many recombination events

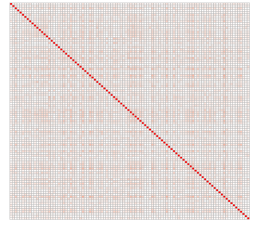
Let's discuss LD structures

Pairwise correlations between SNPs

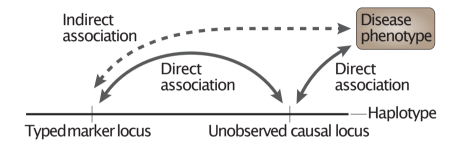
consecutive 100 SNPs



random 100 SNPs



GWAS fail to pinpoint exact locations associated with a disease



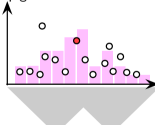
Balding, Nature Genetics Review (2006)

Common strategies to deal with LD structures

Strategy 1.

Fine-mapping to find a handful of causal ones

- Bayesian posterior estimation
- Overlap with epigenomics data



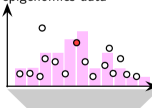
x-axis: genomic location; y-axis: -log10 p-value

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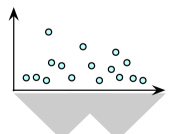
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Strategy 2.

Aggregation to combine all the information:

- Rare variant analysis
- Gene-level enrichment/association



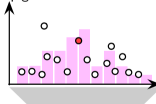
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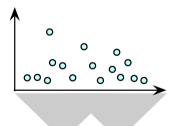
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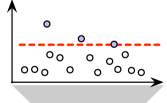
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Strategy 3. Pruning to remove somewhat redundant information (heuristics)

- p-value thresholding
- Useful in polygenic risk prediction

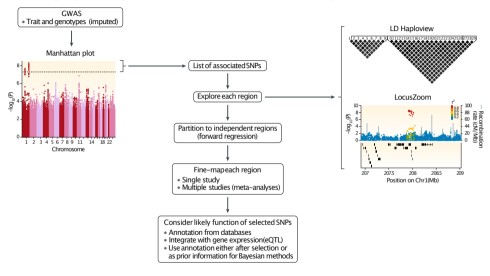


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Why fine-mapping?

A lead SNP^1 within a locus \neq a causal SNP

Fine-mapping typically follows GWAS meta-analysis



Fine-mapping could be done by a variable selection problem

If we had fully observed X and Y for the > 10k samples,

$$\mathbf{y} \sim \sum_{j=1}^{P} \mathbf{x}_{j} eta_{j} + \epsilon$$

1. A greedy forward selection method

- $\mathbf{y} \sim \mathbf{x}_k \beta_k$ (find the best)
- $\mathbf{y} \leftarrow \mathbf{y} \mathbf{x}_k \beta_k$ (take the residual)

(technically not a fine-mapping method)

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2. A brute-force combinatorial search

For all possible subsets of non-zero's $S \subset [p]$:

$$\max_{S} p(\mathbf{y}|\sum_{j\in S} \mathbf{x}_{j}\beta_{j})$$

(usually limit the search space $\left|S\right| < k$)

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- 3. **Bayesian prior**, L1 or spike-slab
- with a sparse prior $p(\beta)$

$$\min \|\mathbf{y} - X\beta\|^2 - \log p(\beta)$$

(normalization is required)

In practice, we don't have a full panel of genotypes!

But we have summary statistics of meta-analysis:

A generative model of SNP-level statistics

For each β_i 's, effect size, variance, z-score:

$$\hat{\beta}_j = \frac{\sum_i X_{ij} Y_i}{\sum_i X_{ij}^2}, \quad \hat{\mathbb{V}}[\beta_j] = \frac{\sigma_{\epsilon}^2}{\sum_{i=1} X_{ij}^2}, \quad Z_j = \frac{\hat{\beta}_j}{\sqrt{\hat{\mathbb{V}}[\beta_j]}}$$

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Although the underlying multivariate regression model:

$$\mathbf{y} = X\theta + \epsilon$$

where $\theta_i \neq \beta_i$

For simplicity, let's assume standardized genotype matrix X, i.e., $\bar{X}_j=0$ and $\hat{\sigma}^2_{X_j}=1$. The we have z-score

$$\hat{Z}_j = \sum_{i=1}^n X_{ij} Y_i / \sigma_\epsilon \sqrt{n}$$

for all $j \in [p]$.

For simplicity, let's assume standardized genotype matrix X, i.e., $\bar{X}_j=0$ and $\hat{\sigma}^2_{X_j}=1$. The we have z-score

$$\mathbf{z}_{p \times 1 \text{ univariate}} = \frac{1}{\sigma \sqrt{n}} X^{\top} \mathbf{y}$$

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$$\mathbf{z}_{p \times 1 \text{ univariate}} \ = \ \frac{1}{\sigma \sqrt{n}} X^{ op} \mathbf{y} = \frac{1}{\sigma \sqrt{n}} X^{ op} \underbrace{(X \theta + \epsilon)}_{\text{a multivariate mode}}$$

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$$= \frac{\sqrt{n}}{\sigma} \underbrace{\left(\frac{1}{n} X^{\top} X\right)}_{\text{LD}} \theta + \frac{1}{\sigma \sqrt{n}} X^{\top} \epsilon$$

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$$\begin{array}{ll} \mathbf{z} & = & \frac{1}{\sigma\sqrt{n}}X^{\top}\mathbf{y} = \frac{1}{\sigma\sqrt{n}}X^{\top}\underbrace{(X\theta + \epsilon)}_{\text{a multivariate model}} \\ & = & \mathbf{R}\frac{\sqrt{n}}{\sigma}\theta + \tilde{\epsilon}, \quad \tilde{\epsilon} \sim \mathcal{N}(\mathbf{0},\mathbf{R}) \end{array}$$

where $\mathbf{R} = n^{-1}X^{T}X$ is an empirical LD matrix.

Fine-mapping is to find a sparse multivariate heta

Input:

- Summary statistics $p \times 1$ z-score vector: **z**
- ightharpoonup Reference panel $p \times p$ LD matrix: R

Goal:

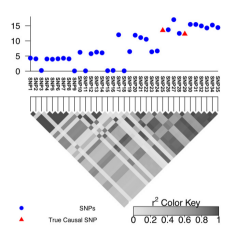
$$\max_{\theta} \mathcal{N}\left(\mathbf{z} \middle| \frac{\sqrt{n}}{\sigma} R \theta, R\right)$$

where

$$\theta_j \sim \pi \delta_0(\theta_j) + (1-\pi) \mathcal{N} \big(0,\tau^{-1}\big)$$

Zhu and Stephens, Annals of Applied Statistics (2017)

Not all GWA-significant variants are causal



A reasonable fine-mapping approach:

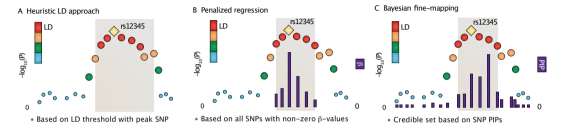
- ldentify GWAS loci (with p $< 5 \times 10^{-8}$)
- ► Find neighbouring SNPs in each GWAS locus
- Convert p-values to z-scores (caution: We should take into account major/minor allele directions)
- ightharpoonup Take an appropriate local LD matrix R
- **E**stimate θ in the following model:

$$\mathbf{z} \sim \mathcal{N}(R\theta, R)$$

▶ We may construct 95% credible set:

$$\{j:\,\hat{p}(\theta_j\neq 0|R,\mathbf{z})>.95\}$$

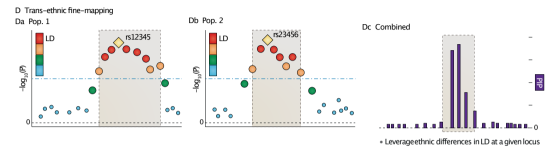
Fine-mapping approaches - 1: heuristics and Bayesian approach



- In almost all cases, Bayesian method outperforms
- Note: This is a "statistical" fine-mapping

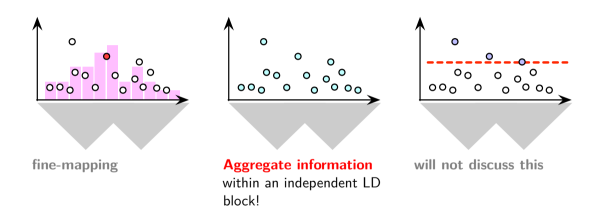
Schaid et al., Nature Review Genetics (2018)

Fine-mapping approaches - 2: trans-ethnic analysis



- Additional information across multiple GWAS summary statistics
- "Causal triangulation" to combine multiple lines of orthogonal evidence

How should we deal with LD structures?



Burden test: Can we aggregate all the SNPs to boost power?

Motivation:

- ightharpoonup Summary statistics $p \times 1$ z-score vector: \mathbf{z}
- ightharpoonup Reference panel $p \times p$ LD matrix: R
- Unfortunately, none of the SNPs make GWA significance

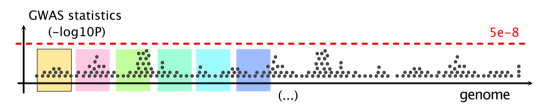
Question:

- ▶ Should we give up on this GWAS result (z-scores)?
- Alternatively, is there any way to reduce the number of hypothesis?

Gene-level aggregate test statistics

Assume underlying multivariate effect θ for the observed z-score vector:

$$\mathbf{z} \sim \mathcal{N}(R\theta, R)$$



- Can we aggregate information over many SNPs within each gene (box)?
- ▶ Although each genetic variant can occur rarely (hence, very weak association statistics), they may implicate the same target gene.

Sequence Kernel Association Test (SKAT) to aggregate rare variant info

In a model $\mathbf{y} \sim X^{(g)}\theta_g$, where X is constructed within a window around a specific gene g, we want to test

$$H_0:\,\theta_q=0\quad\text{vs.}\quad H_1:\,\theta_q=0$$

In a sense, it is the same as doing model comparison (after integrate out θ):

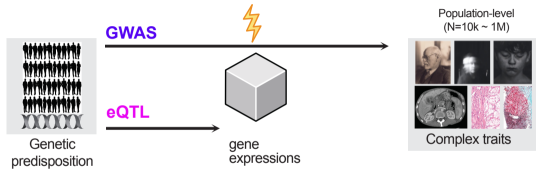
$$H_0: \, \mathcal{N} \Bigg(\mathbf{y} \Bigg| \, \mathbf{0}, \, \underbrace{\tau^2 \mathbf{K}}_{\text{e.g., local kinship}} + \sigma^2 I \Bigg) \quad \text{ vs.} \quad H_1: \, \mathcal{N} \big(\mathbf{y} | \mathbf{0}, \, \sigma^2 I \big)$$

where $K \propto n^{-1} X^{(g)} {X^{(g)}}^{\top}$ or we can substitute it with a different type of kernel matrix. \implies We want to test $H_0: \tau=0$ or not.

Wu, Lee, ..., Lin, AJHG (2011)

Can we aggregate genetic association statistics using prior knowledge?

Expression Quantitative Trait Loci (eQTL) results provide necessary context to interpret GWAS signals



$$\mathbf{m}_g \sim X \alpha_g + \epsilon_g \quad \implies \quad \mathbf{y}_{\text{GWAS}} = \sum_{g \in \text{causal genes}} \mathbf{m}_g \beta_g + \epsilon_y$$

The same type of a linear model for a phenotype vector \mathbf{y}

$$\mathbf{y} \sim X\theta + \epsilon_y,$$

where X is constructed within a cis-window around a specific gene g (say \pm 500kb).

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We also have a gene expression vector \mathbf{m} :

$$\mathbf{m}_{q} \sim X\alpha_{q} + \epsilon_{m}$$

The same type of a linear model for a phenotype vector ${f y}$

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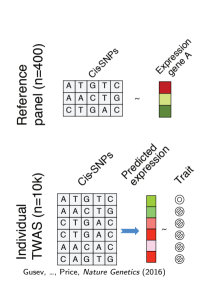
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We also have a gene expression vector \mathbf{m} :

$$\mathbf{m}_{a} \sim X\alpha_{a} + \epsilon_{m}$$

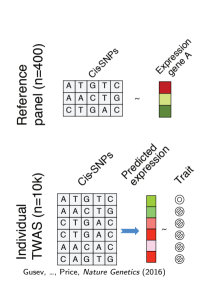
A key question: Are they correlated?

$$H_0: \mathbb{E}[\mathbf{y}^{\mathsf{T}}\mathbf{m}] = 0$$
 vs. $H_1: \mathbb{E}[\mathbf{y}^{\mathsf{T}}\mathbf{m}] \neq 0$



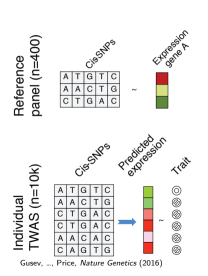
Goal: hypothesis testing of non-zeroness

$$\frac{1}{n}\mathbf{m}^{\top}\mathbf{y} = \frac{1}{n}(X\alpha + \epsilon_m)^{\top}(X\theta + \epsilon_y)$$



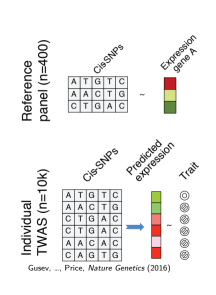
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$$\begin{split} \frac{1}{n}\mathbf{m}^{\top}\mathbf{y} &= \frac{1}{n}(X\alpha + \epsilon_m)^{\top}(X\theta + \epsilon_y) \\ &= \alpha^{\top}\Big(\frac{1}{n}X^{\top}X\Big)\theta + \dots \end{split}$$



Goal: hypothesis testing of non-zeroness

$$\begin{split} \frac{1}{n}\mathbf{m}^{\top}\mathbf{y} &= \frac{1}{n}(X\alpha + \epsilon_m)^{\top}(X\theta + \epsilon_y) \\ &= \alpha^{\top}\Big(\frac{1}{n}X^{\top}X\Big)\theta + \dots \\ &= \alpha^{\top}\underbrace{\left(\frac{1}{n}X^{\top}X\right)\theta}_{\text{GWAS z-score}} + \dots \end{split}$$



Goal: hypothesis testing of non-zeroness

$$\begin{split} \frac{1}{n}\mathbf{m}^{\top}\mathbf{y} &= \frac{1}{n}(X\alpha + \epsilon_m)^{\top}(X\theta + \epsilon_y) \\ &= \alpha^{\top}\Big(\frac{1}{n}X^{\top}X\Big)\theta + \dots \\ \text{(we saw this)} &= \alpha^{\top}\underbrace{\Big(\frac{1}{n}X^{\top}X\Big)\theta}_{\text{GWAS z-score}} + \dots \end{split}$$

TWAS statistic:

$$T_g = \frac{\alpha_g^{\top} \mathbf{z}}{\sqrt{\alpha_g^{\top} R \alpha_g}} \sim \mathcal{N}(0, 1)$$

where $\alpha_q=$ multivariate eQTL for a gene g.

For a standardized $n \times p$ genotype matrix X,

1. Genetic relatedness matrix (GRM)

a $n \times n$ matrix

$$K \approx XX^{\top}/n$$

The matrix K captures population structure/correlation across different individuals

- Kinship matrix; population admixture
- Human migration history

2. Linkage disequilibrium (LD)

a $p \times p$ matrix

$$R \approx X^{\top}X/n$$

The matrix R captures localized correlation patterns along the genomic axis within a chromosome.

- LD matrix
 - The results of many, many recombination events

Today's lecture

Population structures in human genetics data

Linkage Disequilibrium: blessing and curse

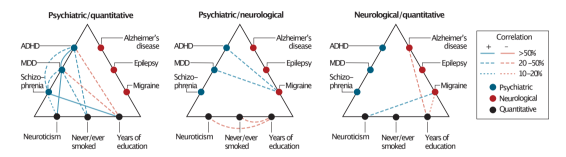
Systems genetics and summary statistics-based inference

GWAS is only the beginning of a post-GWAS analysis.

Post-GWAS analysis example: genetic correlations across many traits

Psychiatric disorders				Neurological disorders			
Disorder	Source	Cases	Controls	Disorder	Source	Cases	Controls
Attention deficit hyperactivity disorder	PGC-ADD2	12,645	84,435	Alzheimer's disease	IGAP	17,008	37,154
Anorexia nervosa	PGC-ED	3495	10,982	Epilepsy	ILAE	7779	20,439
Anxiety disorders	ANGST	5761	11,765	Focal epilepsy	"	4601*	17,985*
Autism spectrum disorder	PGC-AUT	6197	7377	Generalized epilepsy	"	2525*	16,244*
Bipolar disorder	PGC-BIP2	20,352	31,358	Intracerebral hemorrhage	ISGC	1545	1481
Major depressive disorder			153,234	Ischemic stroke	METASTROKE	10,307	19,326
Obsessive-compulsive disorder	PGC-OCDTS	2936	7279	Cardioembolic stroke	"	1859*	17,708*
Posttraumatic stress disorder	PGC-PTSD	2424	7113	Early onset stroke			
Schizophrenia	PGC-SCZ2	33,640	43,456	Large-vessel disease	"	1817*	17,708*
Tourette syndrome	PGC-OCDTS	4220	8994	Small-vessel disease	"	1349*	17,708*
				Migraine	IHGC	59,673	316,078
				Migraine with aura	"	6332*	142,817*
				Migraine without aura	"	8348*	136,758*
				Multiple sclerosis	IMSGC	5545	12,153
				Parkinson's disease	IPDGC	5333	12,019
Total psychiatric		158,028	365,993	Total neurologic		107,190	418,650

Post-GWAS analysis example: genetic correlations across many traits



How did they measure correlations?

The Brainstorm Consortium, Science (2018)

Several benefits of post-GWAS (or systems genetics) analysis

- Inherited genetic information is usually stable over a lifetime.
- It is hard to test/measure all the disease-related phenotypes for all individuals.
- ► Full, unlimited access to individual-level information is often unnecessary in a post-GWAS analysis.
- ▶ A post-GWAS analysis is often computationally more efficient than an individual-level analysis.

LD-score regression: a model-based summary-statistics analysis

What is a generative model for a $\chi_i^2 (\equiv Z_i^2)$ statistics vector?

We have seen that

$$Z_j = \frac{\sqrt{n}}{\sigma} \sum_k R_{jk} \theta_k + \epsilon_j$$

where $\epsilon \sim \mathcal{N}(0,1)$.

LD-score regression: a model-based summary-statistics analysis

What is a generative model for a $\chi_j^2 (\equiv Z_j^2)$ statistics vector?

$$\mathbb{E}\left[\chi_{j}^{2}\right] = \mathbb{E}\left[Z_{j}^{2}\right] = \mathbb{E}\left(\sqrt{n}\sum_{k}R_{jk}\theta_{k} + \epsilon_{j}\right)^{2}$$

Bulik-Sullivan et al., Nature Genetics (2014); Finucane et al., Nature Genetics (2015)

LD-score regression: a model-based summary-statistics analysis

What is a generative model for a $\chi_i^2 (\equiv Z_i^2)$ statistics vector?

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If "true" multivariate effect for each variant is independent of other variants' effects, i.e., $\mathbb{E}\left[\theta_k\theta_j\right]=0$ for all $k\neq j$,

$$\mathbb{E}\left[\chi_{j}^{2}\right] = n \underbrace{\sum_{k} R_{jk}^{2} \mathbb{E}[\theta_{k}^{2}] + 1}_{\text{LD-score}}$$

Baseline LD-score regression to measure polygenic heritability

If we assume $\theta_k=\tau/p$, assuming all the variants can exert weak effects, and defining an LD score for a variant j, $l_j=\sum_k R_{jk}^2$, we have

$$\mathbb{E}\big[\chi_j^2\big] = n \underbrace{\sum_k R_{jk}^2 \, \mathbb{E}\big[\theta_k^2\big] + 1}_{\text{LD-score}}$$

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$$\mathbb{E}\big[\chi_j^2\big] = n \underbrace{\sum_k R_{jk}^2}_{\text{LD-score}} \mathbb{E}\big[\theta_k^2\big] + 1 = \underset{\text{sample size}}{n} l_j \underbrace{\frac{\tau}{p}}_{\text{per SNP heritability}} + 1$$

where p is the total number of SNPs.

Baseline LD-score regression to measure polygenic heritability

If we assume $\theta_k=\tau/p$, assuming all the variants can exert weak effects, and defining an LD score for a variant j, $l_j=\sum_k R_{jk}^2$, we have

$$\mathbb{E}\big[\chi_j^2\big] = \underset{\text{sample size}}{n} l_j \qquad \frac{\tau}{p} + 1$$

We estimate the parameters by regressing the observed χ^2 statistics on the reference LD scores l_i :

$$\begin{pmatrix} \chi_1^2 \\ \vdots \\ \chi_j^2 \\ \vdots \end{pmatrix} \sim n \begin{pmatrix} l_1 \\ \vdots \\ l_j \\ \vdots \end{pmatrix} \frac{\tau}{p} + n\phi + 1 \\ \text{genomic inflation} + 1 \\ \text{per SNP heritability}$$

Stratified LD-score regression

When genome is partitioned by annotations (e.g., epigenetic tracks)

$$\mathbb{E}\left[\chi_{j}^{2}\right] = \frac{n}{p} \sum_{t} l_{jt} \frac{\tau_{t}}{\text{stratified heritability}} + \frac{n\phi}{\text{genomic inflation}} + \frac{1}{\text{null}}$$

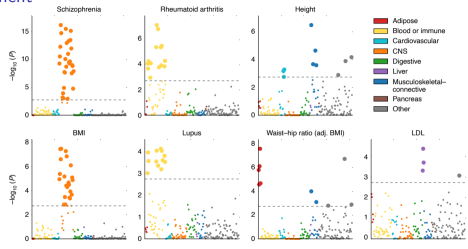
where we use partitioned LD-scores for each annotation type t

$$l_{jt} = \sum_{k} R_{jk}^{2} I\left\{k \in \mathcal{A}_{t}\right\}.$$

More explicitly,

$$\begin{pmatrix} \chi_1^2 \\ \vdots \\ \chi_j^2 \\ \vdots \end{pmatrix} \sim \frac{n}{p} \begin{pmatrix} l_{11} & l_{12} & l_{1t} & \dots \\ & \vdots & & \\ l_{j1} & l_{j2} & l_{jt} & \dots \\ \vdots & & \\ \vdots & & \\ \text{stratified heritability} \end{pmatrix} + \underbrace{n\phi}_{\text{genomic inflation}} + \underbrace{1}_{\text{null}}$$

Stratified LD-score regression can identify tissue-specific heritability enrichment



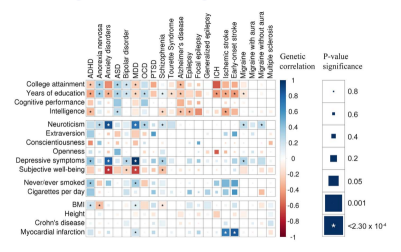
Bivariate LD-score regression

Instead of one χ^2 vector, we need to deal with the element-wise product of two vectors of z-scores (between a trait 1 and 2):

$$\begin{pmatrix} z_1^{(1)}z_1^{(2)}l_1 \\ \vdots \\ z_j^{(1)}z_j^{(2)}l_j \\ \vdots \end{pmatrix} \sim \frac{\sqrt{N_1N_2}}{p} \begin{pmatrix} l_1 \\ \vdots \\ l_j \\ \vdots \end{pmatrix}_{\substack{\text{genetic correlation} \\ \text{sample sharing}}} + \frac{\rho_0N_s}{\sqrt{N_1N_2}}$$

where N_1 and N_2 count sample size of the GWAS 1 and 2; N_s is the number of control individuals shared between the two traits.

Bivariate LD-score regression to test genetic correlations



Learning objective

- Population structures in genetics data
 - Admixture model
 - Linear mixed effect model
- Linkage disequilibrium
 - Rare variant burden tests
 - Fine-mapping causal variants
- ► GWAS summary statistics
 - Transcriptome-wide association studies
 - ► LD-score regression