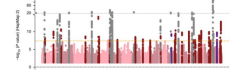
Large-scale genome-wide enrichment analyses of 31 human complex traits

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ICSA 2017, June 28

Examining associations between variables is a useful tool in genetics.

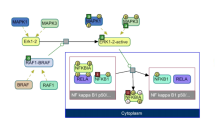
GWAS: SNP~Phenotype



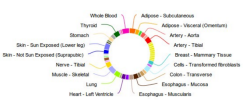
1000 Genomes: SNP~SNP



Biological Pathways: Gene~Gene



GTEx: Gene~Tissue



Enrichment analysis combines multiple sources of association.

- **SNP-trait:** GWAS summary statistics
- **SNP-SNP:** linkage disequilibrium (LD)
- Gene-Gene: biological pathways
- Gene-Tissue: RNA-seq across tissues

Let's keep this talk "jargon-free".

- 1. GWAS summary statistics
- 2. gene set enrichment analysis

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What are GWAS summary statistics?

- Data: phenotype Y and genotype X
- **Size:** n (>10K) individuals and p (>1M) variants (SNPs)

$$Y := \begin{bmatrix} y_1 \\ y_2 \\ \vdots \\ y_n \end{bmatrix} \quad X := \begin{bmatrix} x_{11} & \dots & x_{1j} & \dots & x_{1p} \\ x_{21} & \dots & x_{2j} & \dots & x_{2p} \\ \vdots & \vdots & \vdots & \dots & \vdots \\ x_{n1} & \dots & x_{nj} & \dots & x_{np} \end{bmatrix}$$

Model: single-SNP association analysis

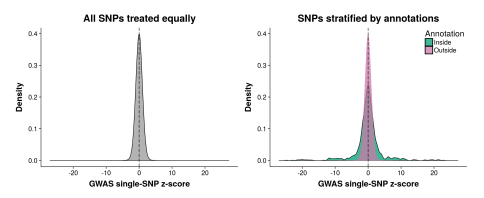
$$\begin{bmatrix} y_1 \\ y_2 \\ \vdots \\ y_n \end{bmatrix} \sim \begin{bmatrix} x_{1j} \\ x_{2j} \\ \vdots \\ x_{nj} \end{bmatrix} \longrightarrow \begin{cases} \hat{\beta}_j : \text{ marginal effect estimate} \\ \hat{\sigma}_j : \text{ standard error of } \hat{\beta}_j \end{cases}$$

• Availability: much more accessible than $\{Y, X\}$ (Nat. Genet., 2012)

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What is enrichment analysis?

- **Phenotype:** low-density lipoprotein (Teslovich *et al.*, 2010)
- Pathway: chylomicron-mediated lipid transport (17 genes)
- Annotation: Is the SNP "near" a pathway gene? (yes or no)



Recent reviews: de Leeuw et al. (2016); Pers (2016); Mooney et al. (2014); Wang et al. (2010).

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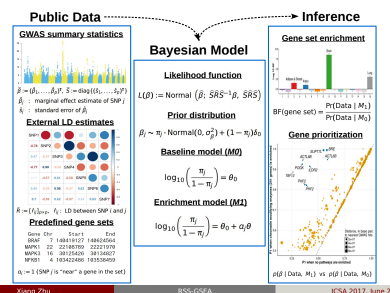
Introduction

The "enrichment" idea is simple, but there are (at least) two technical issues.

- 1. If the gene set is truly enriched, we should relax significance threshold for "green" SNPs, but how much to relax?
 - **Data-driven** threshold ← Function (Pathway, Phenotype)
 - Maintained type 1 error + Improved power
- 2. The "inflated" green curve may be driven by correlation between SNPs (LD), rather than enrichment of signal.
 - SNP 1 has a large genetic effect on a trait.
 - SNPs 2-100 have zero effect, but all are in high LD with SNP 1.
 - Thus, SNPs 1-100 all show very large **single-SNP** z-scores.

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We develop a method that systematically utilizes enrichment information.



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Address Issue 1: Learning enrichment from data

Model-based approach:

- Assume that SNP j is trait-associated with probability π_j
- Represent π_i as a function of SNP-level annotation a_i

$$\log_{10}\left(\frac{\pi_j}{1-\pi_j}\right) := \theta_0 + a_j\theta$$

• Estimate enrichment parameter θ from data

Data-adaptive threshold:

- "Enriched" data $\leadsto \theta > 0 \leadsto$ larger π_j for $a_j = 1 \leadsto$ increased power
- "Null" data $\rightarrow \theta \approx 0 \rightarrow \text{unchanged } \pi_j \rightarrow \text{maintained type 1 error}$

References: Veyrieras *et al.* (2008); Carbonetto & Stephens (2013); Pickrell (2014); Kichaev *et al.* (2014); Chen *et al.* (2016); Li & Kellis (2016); Wen *et al.* (2017).

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Address Issue 2: Modeling linkage disequilibrium

Genome-wide multiple-SNP likelihood function:

$$L_{\mathsf{rss}}(\beta) := \mathsf{Normal}(\widehat{\beta}; \widehat{S}\widehat{R}\widehat{S}^{-1}\beta, \widehat{S}\widehat{R}\widehat{S})$$

- lacksquare multiple-SNP parameter: $oldsymbol{eta}:=(oldsymbol{eta}_1,\ldots,oldsymbol{eta}_{oldsymbol{
 ho}})^{\intercal}$
- lacksquare single-SNP summary data: $\widehat{oldsymbol{eta}}:=(\widehat{eta}_1,\ldots,\widehat{eta}_p)^{\intercal}$
- $\qquad \widehat{S} := \operatorname{diag}(\widehat{s}), \ \widehat{s} := (\widehat{s}_1, \ldots, \widehat{s}_\rho)^{\mathsf{T}}, \ \widehat{s}_j^2 := \widehat{\sigma}_j^2 + n^{-1}\widehat{\beta}_j^2 \approx \widehat{\sigma}_j^2$
- R: the shrinkage estimate of LD (Wen & Stephens, 2010; Li & Stephens, 2003)
- "Big Data" Genetics: jointly analyze 1.1 million common SNPs

Reference: Zhu and Stephens (2017 To appear). http://dx.doi.org/10.1101/042457.

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Methods

We develop a method that systematically utilizes enrichment information.

➤ Inference **Public Data GWAS summary statistics** Gene set enrichment **Bayesian Model** Likelihood function $\hat{\beta} := (\hat{\beta}_1, \dots, \hat{\beta}_n)^T, \hat{S} := \text{diag}\{(\hat{s}_1, \dots, \hat{s}_n)^T\}$ $L(\beta) := Normal(\widehat{\beta}; \widehat{SRS}^{-1}\beta, \widehat{SRS})$ $\hat{\beta}_i$: marginal effect estimate of SNP i Pr(Data | M₁) ŝ_i: standard error of β̂_i BF(gene set) = Prior distribution Pr(Data | M₀) External LD estimates $\beta_j \sim \pi_j \cdot \text{Normal}(0, \sigma_{\beta}^2) + (1 - \pi_j)\delta_0$ Gene prioritization Baseline model (M0) $\log_{10}\left(\frac{\pi_j}{1-\pi_i}\right) = \theta_0$ 0.7 -0.56 0.62 -0.57 0.34 0.63 SNP7 Enrichment model (M1) $\hat{R} := [\hat{r}_{ij}]_{p \times p}, \hat{r}_{ij} : LD \text{ between SNP } i \text{ and } j$ Predefined gene sets $\log_{10}\left(\frac{\pi_j}{1-\pi_i}\right) = \theta_0 + a_j\theta$ P1 when no pathways are enriched $a_i := 1 \{SNP | is "near" a gene in the set \}$

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 $p(B \mid Data, M_1)$ vs $p(B \mid Data, M_0)$

We apply the method to analyze 31 complex traits and 4,026 gene sets.

This application is not small:

```
# of Parameters = 31 \times (3,913+113) \times 1.1 Million \approx 137 Billion
```

- 31 human phenotypes
- 3,913 biological pathways
- 113 tissue-based gene sets
- 1.1 million common SNPs

One student can get this done, aided by:

- Publicly available input data: GWAS + LD + gene sets
- Variational inference; Squared extrapolation method (SQUAREM)
- Parallel computing; Hierarchical data format (HDF5)

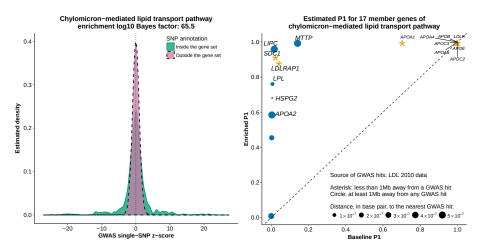
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Our full results are publicly available.

- Results
 https://xiangzhu.github.io/rss-gsea/results
- Software https://github.com/stephenslab/rss
- Demonstration
 https://stephenslab.github.io/rss/Example-5
- R package (in progress) https://github.com/stephenslab/rssr(N. Knoblauch)

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Example: Low-density lipoprotein & MTTP

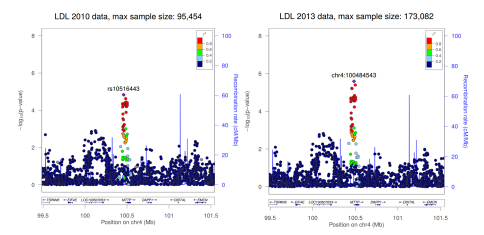


- Gene detection: $P_1 := 1 \Pr(\beta_j = 0, \forall j \in \text{locus} \mid \text{Data})$
- *MTTP*: baseline $P_1 = \mathbf{0.14}$; enriched $P_1 = \mathbf{0.99}$

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Example: Low-density lipoprotein & MTTP

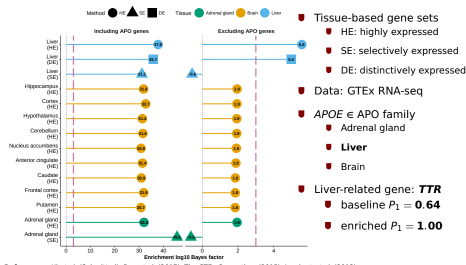
- Total sample size: **95,454** (2010) \longrightarrow **173,082** (2013)
- GWAS p-value: 1.5×10^{-5} (2010) $\longrightarrow 2.6 \times 10^{-6}$ (2013) "<" 5×10^{-8}



References: Global Lipids Genetics Consortium (2013); Teslovich et al. (2010).

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Example: Alzheimer's disease & Liver



References: Xi et al. (Submitted); Dey et al. (2017); The GTEx Consortium (2015); Lambert et al. (2013).

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Discussion

What's next?

We develop a new enrichment analysis method that:

- uses **publicly available** data as input
- **efficiently** assesses thousands of gene sets
- **automatically** identifies trait-associated genes

What do we plan to do next?

- one annotation at a time \rightarrow jointly analyze **many** annotations
- gene-based annotations → "finer-scale" annotations
- spike-and-slab prior → more **flexible** effect size distributions

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GWAS summary statistics

Teslovich *et al.* (2010); Manning *et al.* (2012); Morris *et al.* (2012); van der Harst *et al.* (2012); Köttgen *et al.* (2013); Lambert *et al.* (2013); Okada *et al.* (2014); Ripke *et al.* (2014); Wood *et al.* (2014); Day *et al.* (2015); Liu *et al.* (2015); Locke *et al.* (2015); Nikpay *et al.* (2015); Shungin *et al.* (2015); Okbay *et al.* (2016); van Rheenen *et al.* (2016)

GTEx Consortium

https://gtexportal.org





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RESEARCH COMPUTING CENTER

Thank you!

Preprint: https://doi.org/10.1101/160770

Software: https://github.com/stephenslab/rss

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