Nothing to disclose.
Modeling tissue and gene co-regulation reveals causal tissues for disease

Tiffany Amariuta, PhD
Assistant Professor
University of California San Diego
Halıcıoğlu Data Science Institute / Department of Medicine
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Preprint of presented work: Amariuta 2022 bioRxiv
Outline

1. Background

2. Method and Simulations

3. Results on Real Traits
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Inferring causal tissues is an important goal

Knowing the causal disease tissue is crucial to
1) Describing the biological mechanism of disease
2) Selecting the cellular context in which to perform experiments
3) Determining the physiological target of pharmaceuticals

For most diseases, disease-associated cell types have been identified.

Using chromatin accessibility, bone chondrocytes are the most strongly associated cell type for height.

Finucane 2015 Nat Genet
Also see, Trynka 2013 Nat Genet, Hao 2018 Plos Genet
For most diseases, disease-associated cell types have been identified.

Using gene expression, the frontal cortex (BA9) of the brain is the most strongly associated tissue for schizophrenia.
For most diseases, disease-associated cell types have been identified.

Using enhancer predictions, CD4+ Th1 is the most strongly associated cell type for asthma.
High co-regulation across tissues means that many disease-associated tissues may not be causal

Gene expression is highly correlated across GTEx tissues (co-expression)

cis regulatory effects are highly correlated across GTEx tissues (co-regulation)

GTEx Consortium 2020 Science
Also see Ongen 2017 Nat Genet, Wainberg 2019 Nat Genet, Arvanitis 2022 AJHG
Colocalization of eQTLs with GWAS variants can implicate disease-critical genes and tissues

Gene X

Disease-associated variant from GWAS

Brain

<table>
<thead>
<tr>
<th>Allele affecting gene expression levels (eQTL)</th>
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<tbody>
<tr>
<td>Allele with no effect on gene expression</td>
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is an expression quantitative trait locus (eQTL)

Gene X expression

AA   AT   TT

Giambartolomei 2014 Plos Genet, Hormozdiari 2016 AJHG
Colocalization analysis is complicated by co-regulation

Disease-associated variant from GWAS

Brain → Gene X → disease

- Allele affecting gene expression levels (eQTL)
- Allele with no effect on gene expression

Wainberg 2019 Nat Genet
Siewert-Rocks 2022 AJHG
Colocalization analysis is complicated by co-regulation

Having the same causal variant

Disease-associated variant from GWAS

Brain

Heart

Gene X

Allele affecting gene expression levels (eQTL)

Allele with no effect on gene expression

Wainberg 2019 Nat Genet
Siewert-Rocks 2022 AJHG
Colocalization analysis is complicated by co-regulation

Disease-associated variant from GWAS

Brain

Heart

Muscle

Allele affecting gene expression levels (eQTL)

Allele with no effect on gene expression

Having causal variants in high linkage disequilibrium

Wainberg 2019 Nat Genet
Siewert-Rocks 2022 AJHG
Transcriptome-wide association studies (TWAS) perform polygenic colocalization of genes with disease

1. Learn SNP-gene weights from eQTL model.
2. Predict gene expression using new genotypes.

Gene A expression

SNP-gene estimated effects

Genetically predicted gene expression (W)

Gamazon 2015 Nat Genet, Gusev 2016 Nat Genet
TWAS association statistics are proportional to the amount of tagged causal effects due to co-regulation.

TWAS statistics include direct causal effects and tagging effects of co-regulated genes and tissues.
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Co-regulation across tissues and genes can be estimated using gene expression prediction models and a reference panel

1. Learn SNP-gene weights from eQTL model.

2. Predict gene expression using new genotypes.

3. Compute co-regulation score for a gene-tissue pair with a tissue $t'$.

$$r^2 = \text{squared correlation of } W \text{ between two genes.}$$

$$r^2_{g;g-1} + r^2_{g;g} + r^2_{g;g+1}$$
Tissue co-regulation score regression (TCSC) estimates tissue-specific contributions to disease

\[ E[\chi^2_{g,t}] = N \sum_{t'} \left[ l(g, t, t') \frac{h^2_{ge(t')}}{G_{t'}} \right] + 1 \]

Gene-tissue association with disease (TWAS)

GWAS sample size

Tissue co-regulation scores

# heritable genes in tissue t'

Estimand: Disease heritability explained by predicted gene expression in tissue t’
Tissue co-regulation score regression (TCSC) estimates tissue-specific contributions to disease

\[ E[\chi^2_{g,t}] = N \sum_{t'} [l(g, t, t') \frac{h^2_{ge(t')}}{G_{t'}}] + 1 \]

Our method determines that tissue \( t' \) causally contributes to disease if genes with \textbf{high co-regulation} to tissue \( t' \) have \textbf{higher TWAS }\( \chi^2 \) \textbf{statistics} and genes with \textbf{low co-regulation} to tissue \( t' \) have \textbf{lower TWAS }\( \chi^2 \) \textbf{statistics}.

We may identify tissue-specific contributions to the \textbf{covariance} of two diseases by regressing products of TWAS z-scores on co-regulation scores.
TCSC is powerful, well-calibrated, and unbiased in simulations.

TCSC has substantially higher power than the Ongen 2017 Nat Genet method.

Bars represent 95% CI.
Mancuso Lab TWAS simulator
Amariuta 2022 bioRxiv (see manuscript for many more simulations)
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Applying TCSC to real gene expression and trait data

1. We built gene expression prediction models across 48 GTEx tissues, retaining heritable protein coding genes.

2. We analyzed 78 European GWAS summary statistics (average N = 302K).

3. TCSC
   • Finds 27 causal tissue-disease pairs at 10% FDR.
   • Increases the specificity of known tissue-disease associations.
TCSC identifies causal tissue-disease pairs

1. Aorta artery → Glaucoma:
   - High blood pressure is a known risk factor for glaucoma.

2. Esophagus muscularis → FEV1/FVC:
   - Strength of esophageal muscles likely impacts air expulsion rate (FEV1).
     Analysis of composite traits identified no association with lung capacity (FVC).

3. Heart ventricle → Platelet count:
   - Platelets cause blood clots in response to damaged blood vessels; the left ventricle pumps blood out of the heart potentially modifying platelet counts in serum.

Where $\pi(t') = \text{proportion of disease heritability explained by predicted expression in tissue } t'$
TCSC increases specificity of known tissue-disease pairs

1. Adipose (subcutaneous) → HDL:
   • No causal link to adipose visceral omentum.
   Link between HDL and subcutaneous may involve adiponectin.

2. Adipose (subcutaneous) → WHRadjBMI:
   • No causal link with any other metabolic tissue.

3. Brain (cerebellum) → BMI:
   • Previous studies have identified generic associations with the central nervous system.

Where \( \pi(t') \) = proportion of disease heritability explained by predicted expression in tissue \( t' \)
Other methods are less specific in implicating tissue-disease pairs

Black lines separate tissues with high genetic correlation. Purple circle is TCSC tissue-disease pair.

The remaining complex traits and diseases have similar patterns.

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Higher resolution makes causal inference more difficult, but TCSC can be applied here.

Tissues
- Lung (510)
- Broad mammary tissue (398)
- Pancreas (370)
- Liver (368)
- Adrenal gland (333)
- Kidney cortex (37)
- Kidney medulla (10)
- Visceral omentum (498)
- Small intestine terminal ileum (379)
- Fallopian tube (10)
- Uterus (22)
- Notochord (57)
- Endoderm (350)
- Ectoderm (37)
- Vagina (34)
- Sun-exposed skin (lower leg) (609)
- Cultured fibroblasts (480)
- Subcutaneous adipose (588)
- Skeletal muscle (708)

Cell types
- Minor salivary gland (44)
- Thyroid (374)
- Aorta (387)
- Atrial appendage (372)
- Coronary artery (353)
- Left ventricle (386)
- Esophagus mucosa (497)
- Esophagus muscularis (486)
- Gastric fundus (404)
- Splenic (227)
- Stomach (324)
- Transverse colon (308)
- Small intestine (606)
- Bladder (77)
- Prostate (222)
- Testes (376)
- White blood (670)
- ES-derived lymphocytes (447)

Populations of single cells

GTEx Consortium 2020 Science
Ulirsch 2019 Nature Genetics

from Nathan 2022 Nature
Perez 2022 Science
Yazar 2022 Science
Cross-trait TCSC identifies tissue-disease covariance pairs

Where \( \pi(t') (\text{resp. } \xi(t')) \) = proportion of disease heritability (resp. covariance) explained by predicted expression in tissue \( t' \)

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Conclusions

1. Co-regulation scores and TWAS statistics can be used to infer the causal tissue(s) underlying disease heritability (covariance).

2. TCSC identifies new, biologically plausible tissue-disease pairs including the aorta artery and glaucoma.

3. TCSC may be more informative when applied to dynamic eQTL datasets, case/control eQTL, and single cell gene expression datasets.
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Please check out our manuscript:
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I’m looking for graduate students and postdocs interested in statistical and population genetics to join my group.
website: amariutalab.org contact: tamariuta@gmail.com