## Zeyun Lu

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#### Summary

PhD Student in Biostatistics interested in developing statistical methods to better understand genetic architecture for complex diseases using summary-based data.

#### Education

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Doctor of Philosophy in Biostatistics	08/2019 - Present
University of Southern California	Los Angeles, CA
Master of Science in Statistics	09/2015 - 12/2016
University of California, San Diego	La Jolla, CA
Bachelor of Science in Mathematics/Economics	09/2012 - 09/2014
University of California, Los Angeles	Los Angeles, CA
Associate in Arts Shoreline Community College	03/2010 - 06/2012 Shoreline, WA
Graduate Research Assistant Department of Population and Public Health Sciences	08/2020 - Present Los Angeles, CA

### **Professional** Experience

Keck School of Medicine, USC Mentor: Dr. Nicholas Mancuso

- Developing a Bayesian gene fine-mapping method for complex diseases using summary-based data and molecular data of multi-ethnic groups.
- Developing a method to test post-transcriptional/translational regulatory mechanisms for complex diseases.
- Constructed *cis*-protein Quantitative Loci reference panel using FUSION pipeline.

Statistician 04/2017 - 06/2019 Alzheimer's Therapeutic Research Institute (ATRI) San Diego, CA Keck School of Medicine, USC

Mentor: Dr. Rema Raman and Dr. Michael Donohue

- Participated as Biostatistician in study teams of several Alzheimer's Disease (AD) clinical trials and observational studies such as AD Neuroimaging Initiative 3 (ADNI3), Anti-Amyloid Treatment in Asymptomatic AD (A4), and Longitudinal Early-Onset AD Study (LEADS).
- Designed and generated different study reports for ADNI3, A4, and LEADS. Reports include but not limited to performing data analysis with mixed models, investigating, data anomalies, and implementing patient randomization.
- Designed, built, and administered R Shiny projects that created dynamic online clinical data reporting system and risk-based monitoring system for ADNI3.
- Mentored student intern in R Shiny, clinical trials, and statistical analysis.

Mathematics Graduate Research Internship 03/2016 - 06/2016 Department of Mathematics 09/2016 - 12/2016 University of California, San Diego La Jolla, CA Mentor: Dr. Anthony C. Gamst

 Analyzed several clinical trials data with visualization, tabulation, and statistical methods such as Linear Regression, Logistic Regression, and ANOVA etc. • Investigated data errors and proposed fixing schemes.

#### **Mathematics Tutor**

01/2011 - 06/2012 Shoreline, WA

Math Learning Center

Shoreline Community College

Mentor: Rosalie Tepper

• Tutored students mathematics from basic algebra to calculus and statistics on a weekly basis.

# Awards & Fellowship

3. **Reviewers' Choice**, American Society of Human Genetics 2021 10/2021 Abstract scored by reviewers in the top 10% of all poster abstracts.

- 2. Outstanding Overall Performance, Biostatistics Screening Exam 08/2020 In recognition of top two students in the annual screening exam for PhD students.
- 1. **Keck School of Medicine/Graduate School Fellowship** 2019/2020 In recognition of new PhD students' outstanding academic promise.

#### Articles

1. Integrative genomic analyses identify susceptibility genes underlying COVID-19 hospitalization. Pathak GA\*, Singh K\*, Miller-Fleming TW, Wendt FR, Ehsan N, Hou K, Johnson R, <u>Lu Z</u>, Gopalan S, Yengo L, Mohammadi P, Pasaniuc B, Polimanti R, Davis LK, Mancuso N. **Nature Communication**. July, 2021.

DOI: https://doi.org/10.1038/s41467-021-24824-z

## Posters & Abstracts

- Residual Proteome-wide Association Study Identifies Genes for Blood-Related Traits.
   <u>Lu Z</u>, Jung J, Gazal S, Mancuso N. American Society of Human Genetics.
   October, 2021 (Abstract accepted).
- 5. Integrative analyses identify genes and their functional consequences underlying COVID-19 hospitalization. Singh K, Pathak G, Miller-Fleming T, Wendt F, Ehsan N, Hou K, Johnson R, Lu Z, Gopalan S, Yang L, Mohammadi P, Pasaniuc B, Polimanti R, Davis L, Mancuso N. Molecular Genetics and Metabolism. April, 2021.
- 4. Multi-ethnic fine-mapping improves precision to identify causal genes in transcriptomewide association studies. Gopalan S\*, <u>Lu Z\*</u>, Jiang L, Conti DV, Gusev A, Pasaniuc P, Mancuso N. **Probabilistic Modeling In Genomics**. April, 2021.
- 3. Multi-ethnic fine-mapping improves precision to identify causal genes in transcriptome-wide association studies. Gopalan S, <u>Lu Z</u>, Conti DV, Gusev A, Pasaniuc B, Mancuso, N. American Society of Human Genetics. October, 2020.
- Brain Amyloid Burden, Sleep, and Circadian Rest/Activity Rhythms: Screening Findings from A4 and LEARN. Rosenberg PB, Zipunnikov V, <u>Lu Z</u>, Raman R, Carlsson CM, Mintzer JE, Marshall GA, Porsteinsson AP, Yaari R, Wanigatunga SK, Romano C, Wu MN, Aisen PS, Sperling RA, Spira AP. Alzheimer's & Dementia. November, 2018.
- 1. An Open-source Implementation of Data Standards for Alzheimer's Disease Clinical Trials. Sun CK, Donohue M, Ernstrom K, Jiang Y, <u>Lu Z</u>, Aisen P, Raman R. Clinical Trials on Alzheimer's Disease. Boston, MA. November, 2017.

<sup>\*</sup> Equal Contribution

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#### **Talks**

- Residual Proteome-wide Association Study Identifies Genes for Blood-Related Traits.
   <u>Lu Z.</u> Seminar in Biostatistics and Epidemiology. Department of Population and Public Health Sciences, Keck School of Medicine of USC. Los Angeles, CA. October, 2021 (Exp.).
- Multi-ethnic fine-mapping in transcriptome-wide association studies. <u>Lu Z. Seminar</u> in Biostatistics and Epidemiology. Department of Population and Public Health Sciences, Keck School of Medicine of USC. Los Angeles, CA. September, 2021 (Exp.).
- Probabilistic fine-mapping of transcriptome-wide association studies. <u>Lu Z. Seminar</u> in Biostatistics and Epidemiology. Department of Population and Public Health Sciences, Keck School of Medicine of USC. Online. September, 2020.
- 3. Proteome-Wide Association Study Identifies Independent Protein QTLs for Blood-Related Traits. <u>Lu Z.</u> Seminar in Biostatistics and Epidemiology. Department of Population and Public Health Sciences, Keck School of Medicine of USC. Online. September, 2020.
- 2. A4 Actigraphy Study Pre-randomization Analysis. <u>Lu Z.</u> ATRI Biostatistics Workshop. Alzheimer's Therapeutic Research Institute, San Diego, CA. November, 2018.
- Dynamic Reporting Web Framework for Clinical Trials Using R Shiny. Jain S, <u>Lu Z</u>.
   ATRI Biostatistics Workshop. Alzheimer's Therapeutic Research Institute, San Diego, CA. August, 2018.

## TAship Experience

2. PM-522A: Introduction to the Theory of Statistics
Department of Population and Public Health Sciences
Keck School of Medicine, University of Southern California

08/2021 - 12/2021

1. Analysis of Omics Data for Research Studies
Multi-omics Boot Camp of SHARPP Training
Mailman School of Public Health, Columbia University

06/2021

## Software

#### 3. FOCUS

- Python software to perform gene-trait fine-mapping from transcriptome-wide association studies in either single ethnic or multi-ethnic settings.
- https://github.com/mancusolab/focus

#### 2. ADNI3 and LEADS Dynamic Web Report

- Two Shiny apps that transform static reports to interactive web pages for clinical trails investigators' monitoring on enrollment, primary outcomes, and safety.
- Interval apps that are not public accessible.

### 1. ADNI3, A4, and LEADS

- Three R packages that crawl EDC API, transform data into CDISC format, and automatically build several reports (enrollment, safety etc) for clinical research.
- Interval packages that are not public accessible.

## Other Contributions

#### Reviewer Services

Reviewer for PLOS Computational Biology

Skills

R (Tidyverse, Shiny), Python, SAS, Stata, plink, gcta, bedtools, LDSC, Git, Latex, Linux, CDISC.