# YUHUA ZHANG

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#### **EDUCATION**

## University of Michigan, Ann Arbor, Michigan

2019.9-2023(expected)

Ph.D. in Biostatistics

Dissertation Topic: Statistical Modeling of Large-scale Social Network Data

Supervised by: Dr. Walter Dempsey & Dr. Sebastian Zoellner

# University of Michigan, Ann Arbor, Michigan

2017.9-2019.4

M.S. in Biostatistics

Selected Coursework: Linear Regression, Statistical Inference, Bayesian Modeling, Numerical Linear Algebra, Statistical Computing, Stochastic Process, Linear Programming, Machine Learning

## Shanghai Jiaotong University, Shanghai, China

2013.9-2017.6

B.S. in Biology

Thesis: Identifying Causal Variants of Complex Diseases Based on GWAS Data

#### RESEARCH INTERESTS

Large-scale Network Data Analysis, Cluster Anaysis, Bayesian Nonparametric Method, Statistical Inference for High Dimensional Data, Causal Inference

#### PROFESSIONAL SKILLS SUMMARY

#### Computing Skills

- · Computing language: Python, R, SQL
- · Able to work under Linux environment
- · Able to learn new tools in a decent amount of time

#### Statistical Modeling

- · Bayesian Nonparametric Method
- $\cdot$  Statistical Inference
- · (Generalized) Linear Regression Model

#### Data Experience

- $\cdot$  Large-scale Network Data Interaction data from the TalkLife Platform
- · High dimensional data/Genomic Data Exome sequence data from Prechter Project and MGI Project

#### **Communication Skills**

- · Demonstrated ability to present the research work and analysis results clearly and precisely
- · Experience with collaborating and communicating efficiently with team members

# Language:

· Chinese (native), English (working)

#### PROFESSIONAL EXPERIENCE

## University of Michigan, Ann Arbor

Graduate Student Research Assistant

2018.9 - present

- · Identified the underlying community structures of the network data from TalkLife to help the platform implement proper interference over certain users.
- · Pre-processed and analyzed the exome sequence data in Prechter Project for Bipolar Disorder.

· Analyzed the 15-yr longitudinal data collected from 1,159 bipolar patients to improve clinical diagnosis.

# Yale University

Research Assistant 2016.9 – 2017.5

- · Integrated functional annotation to prioritize GWAS summary stats.
- · Proposed an innovative fine-mapping model under Bayesian framework.
- · Compared polygenic risk scores in different simulation settings.

# **PUBLICATIONS**

Ryan, K. A., Han, P., **Zhang, Y.**, Marshall, D. F., Yocum, A. K., McInnis, M. G., Zöllner, S. (2021). Stability of personality traits in bipolar disorder: Findings from a longitudinal cohort. Journal of Affective Disorders, 283, 1-10.

**Zhang, Y.**, Quick, C., Yu, K., Barbeira, A., Luca, F., Pique-Regi, R., ... & GTEx Consortium. (2019). Investigating tissue-relevant causal molecular mechanisms of complex traits using probabilistic TWAS analysis. bioRxiv, 808295.

Hu, Y., Lu, Q., Liu, W., **Zhang, Y.**, Li, M., & Zhao, H. (2017). Joint modeling of genetically correlated diseases and functional annotations increases accuracy of polygenic risk prediction. PLoS genetics, 13(6), e1006836.

## Manuscripts in Preparation

**Zhang, Y.**, Dempsey, W. An edge exchangeable model based community detection method for large-scale network data

Zhang, Y., Zoellner, S. Intergrating gene co-expression network communities into the effect size estimation

#### LEADERSHIP AND SERVICE

## Department of Biostatistics, University of Michigan, Ann Arbor

- Co-founder and member of the DEI Student Committee, August 2020- April 2021
- Co-founder and member of the Student Research Conflict Committee, August 2020- April 2021
- Departmental DEI committee member, January 2019- June 2020