HAOYU ZHANG, Ph.D.

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EDUCATION

Johns Hopkins University, Department of Biostatistics

Baltimore, MD

Ph.D. in Biostatistics, *Advisors*: Nilanjan Chatterjee and Ni Zhao

Sep 2014 - May 2019

Thesis title: Testing for Genetic Association and Building Risk Prediction Models for Cancers Incorporating Tumor Characteristics

Zhejiang University, Department of Mathematics

Bachelor of Science in Statistics

Hangzhou, China Sep 2010 - Jun 2014

Professional Experience

Harvard University, Department of Biostatistics

Boston, MA

Postdoctoral fellow, Advisor: Xihong Lin

Aug 2019 - Present

- o Develop robust Mendelian Randomization methods to identify causal risk factors for complex traits
- o Build models to quantify COVID-19 transmission patterns using U.S. states and county data
- o Co-advise a Ph.D. student with Dr. Lin on developing new polygenic risk score (PRS) methods

National Cancer Institute

Rockville, MD

Special volunteer, Advisors: Nilanjan Chatterjee and Montserrat García-Closas Aug 2019 - Present

- o Develop multi-ancestry PRS methods using empirical Bayes and super learning algorithm
- o Construct PRS for the risk of breast cancer in African Americans

National Cancer Institute

Rockville, MD

Pre-doctoral fellow, Advisor: Montserrat García-Closas

July 2017 - May 2019

- o Led analyses through multidisciplinary collaborations in the Breast Cancer Association Consortium
- o Discovered 32 novel breast cancer susceptibility loci through overall and subtype-specific GWAS
- o Revealed the complex etiological heterogeneity patterns of 173 known breast cancer risk loci

Johns Hopkins University, Department of Biostatistics

Baltimore, MD

Research assistant, Advisor: Nilanjan Chatterjee

Oct 2015 - May 2019

- o Developed methods for testing genetic associations with cancer risk incorporating tumor characteristics
- o Built PRS models for breast cancer subtypes incorporating tumor characteristics
- Collaborated on analyzing an Indian gallbladder cancer GWAS dataset

Johns Hopkins University, Department of Biostatistics

Baltimore, MD

Research assistant, *Advisor*: Thomas A. Louis

June 2015 - Sep 2015

o Developed nonparametric maximum likelihood model to estimate a population-averaged value

Yale University, Department of Biostatistics

New Haven, CT

Research assistant, Advisors: Hongyu Zhao and Can Yang

Jul 2013 - Sep 2013

o Built prediction models for human population responses to toxic compounds using GWAS datasets

The University of Queensland, Queensland Brain Institute

Research assistant, Advisors: Naomi Wray and Guobo Chen

o Conducted a GWAS for lithium treatment response in bipolar disorder patients

Brisbane, Australia Jul 2012 - Sep 2012

GRANTS AND FELLOWSHIPS

- 1. **K99 CA256513 NIH/NCI Pathway to Independence Award**, Methods for Mendelian randomization and mediation analysis using integrative genetic and genomic data for breast cancer, Role: Principal Investigator, Amount: \$192,702, Jan 2021- Dec 2022
- 2. Pre-doctoral fellowship, National Cancer Institute, Rockville, MD, USA, July 2017- May 2019

Honors and Awards

Harvard University

1. American Society of Human Genetics Annual Conference (ASHG) Reviewers' Choice poster award, virtual meeting, Oct 2021

Johns Hopkins University

- American Statistical Association Section on Statistics in Genomics and Genetics Distinguished Student Paper Award, Denver, CO, USA, Aug 2019
- 2. Delta Omega Scholarship, Baltimore, MD, USA, May 2019
- 3. Delta Omega Public Health Honor Society, Alpha Chapter, Baltimore, MD, USA, April 2019
- 4. Louis I. and Thomas D. Dublin Award, Baltimore, MD, USA, March 2019
- 5. DCEG Fellows Award for Research Excellence (DFARE), National Cancer Institute, Rockville, MD, USA, Nov 2018 (only graduate student winner in 2018)
- 6. Joseph Zeger Travel award to ENAR, Baltimore, MD, USA, March 2018
- 7. Symposium on Advances in Genomics, Epidemiology, and Statistics (SAGES) Poster Award, University of Pennsylvania, Philadelphia, PA, USA, June 2017

Zhejiang University

- 1. Meritorious Winner in the American Mathematical Contest in Modeling, virtual competition, 2013
- 2. First Class Scholarship for Outstanding Merit, Hangzhou, China, 2011
- 3. National Scholarship, Ministry of Education, Hangzhou, China, 2011 (Top 2 %)
- 4. Excellent Student Honor, Hangzhou, China, 2011

5. Scholarship for Distinguished Social Work, Hangzhou, China, 2011

Publications

- * represents joint first author
- 1. Shi, A., Gaynor, S. M., Dey, R., **Zhang, H.**, Quick, C., Lin, X. (2022). COVID-19 Spread Mapper: A multi-resolution, unified framework and open-source tool. *Bioinformatics*.
- 2. Ahearn, T.U.*, **Zhang, H.***, ..., Garcia-Closas, M. Chatterjee, N., (2022). Common variants in breast cancer risk loci predispose to distinct tumor subtypes. *Breast Cancer Research*, 24(1), 1-13.
- 3. **Zhang, H.**, Zhao, N., Ahearn, T.U., Garcia-Closas, M., Chatterjee, N. (2021). A mixed-model approach for powerful testing of genetic associations with cancer risk incorporating tumor characteristics. *Biostatistics*, 22(4), 772-788.
- 4. Mhatre, S., Richmond, R. C., Chatterjee, N., Rajaraman, P., Wang, Z., **Zhang, H.**, ..., Dikshit, R. P. (2021). The role of gallstones in gallbladder cancer in India: a Mendelian randomization study. *Cancer Epidemiology and Prevention Biomarkers*, 30(2), 396-403.
- 5. Zhang, Y. D., Hurson, A. N., **Zhang, H.**, ..., Chanock, S. J., Chatterjee, N., Garcia-Closas, M. (2020). Assessment of polygenic architecture and risk prediction based on common variants across fourteen cancers. *Nature communications*, 11(1), 1-13.
- 6. **Zhang, H.***, Ahearn, T. U.*, ..., Easton, D., Chatterjee, N., Garcia-Closas, M. (2020). Genome-wide association study identifies 32 novel breast cancer susceptibility loci from overall and subtype-specific analyses. *Nature Genetics*, 52(6), 572-581.
- 7. Koutros, S., Kogevinas, M., Friesen, M. C., ..., **Zhang, H.**, ..., Tardon, A. (2020). Diesel exhaust and bladder cancer risk by pathologic stage and grade subtypes. *Environment International*, 135, 105346.
- 8. Zhao, N., **Zhang, H.**, Clark, J., Maity, A., Wu, M. (2018). Composite kernel machine regression based on likelihood ratio test for joint testing of genetic and gene–environment interaction effect. *Biometrics*, 75(2), 625-637.
- 9. Hu, Q., Ding, Z., **Zhang, H.**, He, Y. (2018). The superior glenohumeral joint capsule alone does not prevent superior translation of the humeral head: an in vitro biomechanical study. *Arthroscopy: The Journal of Arthroscopic & Related Surgery*, 34(11), 2962-2970.
- 10. Derkach, A., **Zhang, H.**, Chatterjee, N. (2017). Power Analysis for Genetic Association Test (PAGEANT) provides insights to challenges for rare variant association studies. *Bioinformatics*, 34(9), 1506-1513.
- 11. Mhatre, S., Wang, Z., Nagrani, R., ..., **Zhang, H.**, ..., Dikshit, R., Chatterjee, N., Rajaraman, P. (2017). Common genetic variation and risk of gallbladder cancer in India: a case-control genome-wide association study. *The Lancet Oncology*, 18(4), 535-544.

PREPRINTS

- * represents joint first author
- 1. **Zhang, H.**, Zhan, J., Jin, J.,..., Chatterjee, N., (2022). Novel Methods for Multi-ancestry Polygenic Prediction and their Evaluations in 3.7 Million Individuals of Diverse Ancestry. *Biorxiv*.
- 2. Zhu, J., **Zhang, H.**, Niu, Z., Song, J., Jiang, X., (2020). Do mental health disorders influence the risk of female cancer? Results from a genetic study. (*Submitted*)
- 3. Jiang, Z., **Zhang H.**, Ahearn T.U., Garcia-Closas, M., Chatterjee N., Zhan, X., Zhao, N., (2021), The sequence kernel association test for multi-categorical outcomes with the application to a breast cancergenome-wide association study. (*Submitted*)

PRESENTATIONS

Invited Talks

- 1. Methods for risk prediction using integrative multi-ancestry genetic and genomic datasets
 - o PRIMED Consortium Method Working Group seminar, Apr 2022
 - o Program in Quantitative Genomics (PQG) seminar series, Harvard T.H. Chan School of Public Health, Mar 2022
 - o Department of Human Genetics, Emory University, Feb 2022
 - o Rutgers Cancer Institute of New Jersey, Rutger University, Jan 2022
 - o Department of Biostatistics, University of Michigan, Jan 2022
 - o Division of Cancer Epidemiology & Genetics, National Cancer Institute, Jan 2022
 - o Institute for Genomic Health, Icahn School of Medicine at Mount Sinai, Jan 2022
 - o Department of Biostatistics, Boston University, Jan 2022
 - o Biostatistics & Computational Biology Branch, National Institute of Environmental Health Sciences, Jan 2022
 - o Department of Biostatistics, New York University, Jan 2022
 - o Department of Mathematical Sciences, New Jersey Institute of Technology, Dec 2021
- 2. Developing transethnic polygenic risk scores through empirical Bayes and super learning algorithm
 - International Chinese Statistical Association (ICSA) Applied Statistics Symposium, virtual meeting, Sep 2021
 - Program in Genetic Epidemiology and Statistical Genetics (PGSG) seminar series, virtual meeting, May 2021
- 3. Quantifying the transmission dynamics of COVID-19 using U.S. states and county level data
 - o Joint Statistical Meetings (JSM), virtual meeting, Aug 2021
 - o ICSA Applied Statistics Symposium, virtual meeting, Dec 2020
- 4. Methods for developing ancestry specific polygenic risk scores, *Breast Cancer Association Consortium* (*BCAC*) 25th annual meeting , virtual meeting, Sep 2020

- 5. A mixed-model approach for powerful testing of genetic associations with cancer risk incorporating tumor characteristics
 - Department of Data Science, Southwestern University of Finance and Economics, Chengdu, China, Jan 2020
 - o School of life Sciences and Biotechnology, Shanghai Jiao Tong University, Shanghai, China, Dec 2019
- 6. Subtypes specific GWAS and risk prediction modeling on breast cancer subtypes, *BCAC 23rd annual meeting*, Springdale, UT, USA, Apr 2019
- 7. Standard approach and methods accounting for subtype heterogeneity identified 32 novel breast cancer loci, *BCAC 22nd annual meeting*, Edinburgh, UK, Jun 2018
- 8. Genome-wide association study identifies 19 novel breast cancer loci from analyses accounting for subtype heterogeneity, *ASHG Platform Talk*, Orlando, FL, USA, Oct 2017
- 9. "Testing for genetic associations in case-control studies incorporating multivariate disease characteristics", *National Cancer Institute, Division of Cancer Epidemiology & Genetics, Integrative Tumor Epidemiology Branch*, Rockville, MD, USA, Sep 2017

Contributed Talks

- 1. Robust Mendelian randomization (MR) analysis with weak instruments, *JSM*, virtual meeting, Aug 2020
- 2. A mixed-model approach for powerful testing of genetic associations with cancer risk incorporating tumor characteristics, *JSM*, Denver, CO, USA, Aug 2019
- 3. Developing polygenic risk prediction models for cancer subtypes incorporating multivariate disease characteristics, *JSM*, Vancouver, BC, Canada, July 2018
- 4. The mixture approach to estimating a weighted population-averaged Value, *Eastern North American Regional meeting of the International Biometric Society (ENAR)*, Atlanta, GA, USA, Mar 2018
- 5. The mixture approach to estimating a population-averaged Value, *ENAR*, Washington, D.C., USA, Mar 2017
- 6. Testing for genetic associations in case-control studies incorporating multivariate disease characteristics, JSM, Baltimore, MD, USA, Aug 2017

Posters

- 1. Developing transethnic polygenic risk scores through empirical Bayes and super learning algorithm, *ASHG*, virtual meeting, Oct 2021
- 2. A mixed-model approach for powerful testing of genetic associations with cancer risk incorporating tumor characteristics, *ASHG*, San Diego, CA, USA, Oct 2018
- 3. Genome-wide association study identifies 19 novel breast cancer loci from analyses accounting for subtype heterogeneity

- o Johns Hopkins Biostatistics Retreat, Philadelphia, PT, USA, April 2018
- o 14TH Annual NIH Graduate Student Research Symposium, Bethesda, MD, USA, Feb 2018
- o Delta Omega Poster Competition, Baltimore, MD, USA, Feb 2018
- o The Maryland Genetics, Epidemiology, and Medicine Training Program (MD-GEM), Baltimore, MD, USA, Feb 2018
- 4. Methods for analyzing genome-wide association Studies, *Johns Hopkins University, Department of Biostatistics*, 100 years celebration, Baltimore, MD, USA, Jul 2017
- 5. Testing for genetic association in case-control studies incorporating multivariate disease characteristics, *SAGES*, Philadelphia, PT, USA, Jun 2017
- 6. Testing for genetic association in case-control studies incorporating multivariate disease characteristics, *MD-GEM*, Baltimore, MD, USA, Feb 2017

Advising and Teaching

Advising

- 1. Tony Chen, Ph.D. student, Department of Biostatistics, Harvard University, April 2021 Present (co-advise with Xihong Lin)
 - o Developing PRS methods to improve risk predictions for complex traits
 - o Performing mediation analysis using PRS as mediators

Lab Instructor

Responsibilities: teaching 3 to 4 hours labs per week, beta-test, proctor and grade exams

- 1. Statistical Methods in Public Health II-III (Biostat 622-623), Master Level Class, taught by Marie Diener-West and Karen Bandeen-Roche, Baltimore, MD, USA, Oct 2018 Mar 2019
- 2. Statistical Methods in Public Health II-III (Biostat 622-623), Master Level Class, taught by Marie Diener-West and Karen Bandeen-Roche, Baltimore, MD, USA, Oct 2017 Mar 2018

Teaching Assistant

Responsibilities: hold office hour, grade homework and exams, and hold some classes

- Statistical Methods in Public Health I-IV (Biostat 621-624), Master Level Class, taught by Marie Diener-West, Karen Bandeen-Roche, John McGready and James Tonascia, Baltimore, MD, USA, Sep 2016 May 2017
- 2. Statistical Theory I-IV (Biostat 731-734), PhD Level Class, taught by Elizabeth L. Ogburn, Constantine Frangakis and Michael Rosenblum, Baltimore, MD, USA, Sep 2015 May 2016
- 3. Survey Methodology, Undergraduate Level Class, taught by Peng Zhang, Hangzhou, China, Feb 2014 Jun 2014

SOFTWARE AND PROGRAMMING SKILLS

R shiny app

1. PAGEANT: Power Analysis for GEnetic AssociatioN Tests Software link: https://andrewhaoyu.shinyapps.io/PAGEANT/Github link: https://github.com/andrewhaoyu/PAGEANT

R package

- 1. **CKLRT**: Composite Kernel Machine Regression Based on Likelihood Ratio Test CRAN link: https://cran.r-project.org/web/packages/CKLRT/index.html Github link: https://github.com/andrewhaoyu/CKLRT
- 2. **TOP**: Two-stage polytomous logistic regression Github link: https://github.com/andrewhaoyu/TOP
- 3. **CT-SLEB**: A multi-ancestry polygenic risk score approach Github link: https://github.com/andrewhaoyu/CTSLEB

Programming skills

R (80k-90k lines), C (10k lines), Shell Script, RcppEigen, Stan, Photoshop, MATLAB, PLINK, QCTOOL, GTOOL, STATA, SPSS, E-View, Origin

EDITORIAL ACTIVITIES

Ad hoc reviewer

Nature Communication, Plos Computational Biology, Plos One, Frontiers in Genetics, Plos Genetics

Services

- Statistical genetics group website maintainer (http://jhustatgen.org), Johns Hopkins University, Department of Biostatistics, Baltimore, MD, USA, August 2018 May 2019
- 2. Session chair, Statistical Advances in the Design and Analysis of Sequence-Based Genetic Association Studies, JSM, Baltimore, MD, USA, August 2018
- 3. **Student representative**, Johns Hopkins University, Department of Biostatistics, Baltimore, MD, USA, August 2017 August 2018
 - o Hold the student lunches with topics including passing qualify exam and finding summer internship
 - o Organizes sessions to help students writing cover letter for summer internship
 - o Attend faculty monthly meetings
- 4. TA training day planning committee, Johns Hopkins University, Department of Biostatistics, Baltimore, MD, USA, August 2018
- 5. Johns Hopkins Biostatistics Mixer Organizer, JSM, Vancouver, BC, Canada, August 2018
- 6. **Johns Hopkins Biostatistics Mixer Organizer**, ENAR, Atlanta, GA, USA, August 2018

- 7. TA training day planning committee, Johns Hopkins University, Department of Biostatistics, Baltimore, MD, USA, August 2017
- 8. TA training day planning committee, Johns Hopkins University, Department of Biostatistics, Baltimore, MD, USA, August 2016
- 9. **Volunteer service**, Sharadadevi Gramudyog Utpadak Mandali Ltd (SGS), **Baroda, India**, June 2011 August 2011
 - o Collaborated with volunteers form 12 countries to create a detailed survey to examine multiple aspects of village life
 - o Interviewed locals in 16 villages to gather data about education, hygiene, income and saving, etc.
 - o Analyzed collected data and presented recommendations about serving the needs of nearby communities