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

Postdoctoral fellow, *Advisor*: Xihong Lin

Boston, MA Aug 2019 - Present

o Developing robust Mendelian Randomization methods accounting for weak instrumental bias

o Developing models to quantify COVID-19 transmission patterns using U.S. states and county data

National Cancer Institute

Rockville, MD

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

Aug 2019 - Present

- o Developing trans-ethnic polygenic risk scores (PRSs) using Emprical Bayes and super learning algorithm
- o Developing PRSs for the risk of breast cancer using African American data

National Cancer Institute

Rockville, MD

Pre-doctoral fellow, Advisor: Montserrat García-Closas

July 2017 - May 2019

Led several multidisciplinary international collaborations using the largest breast cancer GWAS dataset

Johns Hopkins University, Department of Biostatistics

Research assistant, *Advisor*: Nilanjan Chatterjee

Baltimore, MD Oct 2015 - May 2019

- o Developed methods for testing of genetic associations with cancer risk incorporating tumor characteristics
- Built PRS models for breast cancer subtypes incorporating tumor characteristics
- o Collaborated on GWAS analysis for gallbladder cancer in India

Yale University, Department of Biostatistics

New Haven, CT

Research assistant, *Advisors*: Hongyu Zhao and Can Yang

Jul 2013 - Sep 2013

o Developed prediction models for environmental chemical exposure using GWAS data

The University of Queensland, Queensland Brain Institute

Research assistant, Advisors: Naomi Wray and Guobo Chen

Brisbane, Australia Jul 2013 - Sep 2013

o Conducted GWAS for lithium response in bipolar disorder patients

GRANTS AND FELLOWSHIPS

- 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: PI, Jan 2021-Dec 2025
- 2. **Pre-doctoral fellowship**, National Cancer Institute, Rockville, MD, USA, July 2017- May 2019

Honors and Awards

Johns Hopkins University:

- ASA Section on Statistics in Genomics and Genetics (SGG) 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, Hangzhou, China, 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. 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.

- 2. 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.
- 3. **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*, 1-10.
- 4. **Zhang, H.**, Zhao, N., Ahearn, T.U., Garcia-Closas, M., Chatterjee, N. (2020). A mixed-model approach for powerful testing of genetic associations with cancer risk incorporating tumor characteristics. *Biostatistics*.
- 5. 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.
- Zhao, N., Zhang, H., Clark, J., Maity, A., Wu, M. (2018). Composite Kernel Machine Regression based on Likelihood Ratio Test with Application for Combined Genetic and Gene-environment Interaction Effect. *Biometrics*, 75(2), 625-637.
- 7. 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.
- 8. 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.
- 9. 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. 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 to Breast Cancer Research*)
- 2. Ahearn, T.U.*, **Zhang, H.***, ..., Garcia-Closas, M. Chatterjee, N., 2020, Common breast cancer risk loci predispose to distinct tumor subtypes. (*Major revision at Breast Cancer Research*)

Presentations

Platform Talk:

Genome-wide Association Study (GWAS) Identifies 19 Novel Breast Cancer Loci From Analyses Accounting For Subtype Heterogeneity, American Society of Human Genetics Annual Conference (ASHG)
Platform Talk, Orlando, FL, USA, October 2017

Invited Talks:

- 1. Quantifying the Transmission dynamics of COVID-19 using U.S. States and County Level Data, *Joint Statistical Meetings (JSM)*, virtual meeting, Aug 2021
- 2. Developing Trans-ethnic Polygenic Risk Scores Through Empirical Bayes and Super Learning Algorithm, *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, *International Chinese Statistical Association (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, *Shanghai Jiao Tong University*, Shanghai, China, December 2019
- 6. Subtypes Specific GWAS and Risk Prediction Modeling on Breast Cancer Subtypes, *BCAC 23rd annual meeting*, Springdale, UT, USA, April 2019
- 7. Standard Approach and Methods Accounting for Subtype Heterogeneity Identified 33 Novel Breast Cancer Loci, *BCAC 22nd annual meeting*, Edinburgh, UK, June 2018
- 8. 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, September 2017

Topic Contributed Talks:

1. Testing for Genetic Associations in Case-control Studies Incorporating Multivariate Disease Characteristics, JSM, Baltimore, MD, USA, August 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, March 2018
- 5. The Mixture Approach to Estimating a Population-averaged Value, *ENAR*, Washington, D.C., USA, March 2017

Posters:

- 1. A mixed-model approach for powerful testing of genetic associations with cancer risk incorporating tumor characteristics, *ASHG*, San Diego, CA, USA, October 2018
- 2. Genome-wide association study (GWAS) identifies 19 novel breast cancer loci from analyses accounting for subtype heterogeneity, *Johns Hopkins Biostatistics Retreat*, Philadelphia, PT, USA, April 2018
- 3. Genome-wide association study (GWAS) identifies 19 novel breast cancer loci from analyses accounting for subtype heterogeneity, 14TH Annual NIH Graduate Student Research Symposium, Bethesda, MD, USA, February 2018
- 4. Genome-wide association study (GWAS) identifies 19 novel breast cancer loci from analyses accounting for subtype heterogeneity, *Delta Omega Poster Competition*, Baltimore, MD, USA, Feb 2018
- 5. Genome-wide association study (GWAS) identifies 19 novel breast cancer loci from analyses accounting for subtype heterogeneity, *The Maryland Genetics*, *Epidemiology*, *and Medicine Training Program (MD-GEM)*, Baltimore, MD, USA, February 2018
- 6. Methods for Analyzing Genome-wide Association Studies, *Johns Hopkins University, Department of Biostatistics*, 100 years celebration, Baltimore, MD, USA, July 2017
- 7. Testing for genetic association in case-control studies incorporating multivariate disease characteristics, *SAGES*, Philadelphia, PT, USA, June 2017
- 8. Testing for genetic association in case-control studies incorporating multivariate disease characteristics, *MD-GEM*, Baltimore, MD, USA, February 2017

Advising and Teaching

Advising:

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

Lab Instructor:

Responsibilities: teaching 3 to 4.5 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
- Survey Methodology, Undergraduate Level Class, taught by Peng Zhang, Hangzhou, China, Feb 2014 -Jun 2014

Software

R shiny app:

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

R package:

- 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

EDITORIAL ACTIVITIES

Ad hoc reviewers

Nature Communication, Plos Computational Biology, Plos One

Services

- 1. Statistical genetics group website maintainer (http://jhustatgen.org), Johns Hopkins University, Department of Biostatistics, Baltimore, MD, USA, August 2018 now
- 2. Session chair, Statistical Advances in the Design and Analysis of Sequence-Based Genetic Association Studies, Joint Statistical Meeting, 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**, Joint Statistical Meeting, Vancouver, BC, Canada, August 2018
- 6. **Johns Hopkins Biostatistics Mixer Organizer**, Eastern North American Regional meeting, 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 Researched micro-finance projects' impact on villagers' lives in Baroda teaming up with interns from 12 countries
 - o Surveyed 16 villages to collect data on sanitation, education and income levels

Programming Skills

R (50k-60k lines), C (10k lines), Shell Script, RcppEigen, Stan, Photoshop, MATLAB, PLINK, QCTOOL, GTOOL, STATA, SPSS, E-View, Origin