

CONTACT INFORMATION	13 Hang Kong Road, Wuhan, Hubei, China	Email: chaolong@hust.edu.cn Website: http://chaolongwang.github.io
RESEARCH INTEREST	Population genetics, statistical genetics/genomics, disease gene mapping, integrative genomics, next-generation sequencing data analysis, high dimensional data analysis	
PROFESSIONAL APPOINTMENTS	08/2018-Now: Professor , Epidemiology & Biostatistics School of Public Health, Tongji Medical College Huazhong University of Science and Technology, Wuhan, Hubei, China 01/2015-08/2018: Principal Investigator , Computational & Systems Biology Genome Institute of Singapore, A*STAR, Singapore 05/2016-08/2018: Adjunct Assistant Professor , Centre for Computational Biology Duke-NUS Medical School, Singapore 09/2012-12/2014: Research Fellow , Department of Biostatistics Harvard T.H. Chan School of Public Health, Harvard University, Boston, Massachusetts, USA • Postdoctoral research on statistical genetics with <i>Xihong Lin</i> and <i>Liming Liang</i>	
EDUCATION	08/2012: Ph.D. in Bioinformatics University of Michigan, Ann Arbor, Michigan, USA • Ph.D. advisor: <i>Noah Rosenberg</i> (Stanford University since 07/2011) • Thesis: Statistical methods for analyzing human genetic variation in diverse populations • 01/2012-08/2012: “Post”-doctoral research on statistical genetics with <i>Gonçalo Abecasis</i> 04/2011: M.A. in Statistics and M.S. in Bioinformatics University of Michigan, Ann Arbor, Michigan, USA 07/2008: B.S. in Physics Peking University, Beijing, China	
SELECTED AWARDS	- PQG Postdoc Travel Award, Program in Quantitative Genomics, Harvard University, 2014 - Charles J. Epstein Trainee Award semifinalist, American Society of Human Genetics, 2013 - Stellar Abstract Award in the 6th Annual PQG Conference, Harvard University, 2012 - HHMI International Student Research Fellowship, Howard Hughes Medical Institute, 2011-2012 - Rackham Predoctoral Fellowship, University of Michigan, 2011-2012 (declined) - DeLill Nasser Award, Genetics Society of America, 2011 - Rackham Conference Travel Grant, University of Michigan, 2009, 2010, 2011 - Fellowship (Program in Biomedical Sciences), University of Michigan, 2008-2009 - May Fourth Scholarship, Peking University, 2006	
SOFTWARE PACKAGES	This section lists bioinformatics software packages we developed. SEEKIN : a C++ program for estimating pairwise genetic relatedness in structured and admixed populations using sparse sequencing data. (Dou <i>et al.</i> 2017, <i>PLOS Genet</i>) LASER : a package written in C++ for estimating individual ancestry using either sequencing reads or genotyping data. (Wang <i>et al.</i> 2014, <i>Nat Genet</i> ; Wang <i>et al.</i> 2015, <i>Am J Hum Genet</i>) LASER Server : a web server based on the LASER method to estimate individual ancestry. (https://laser.sph.umich.edu ; Taliun <i>et al.</i> 2017, <i>Bioinformatics</i>) GMMAT : an R package to perform efficient genome-wide association tests based on generalized linear mixed models. (Chen*, Wang* <i>et al.</i> 2016, <i>Am J Hum Genet</i>) MicroDrop : a C++ program for estimating and correcting for allelic dropout in microsatellite data without replicated genotyping. (Wang <i>et al.</i> 2012, <i>Genetics</i>)	

MAJOR
PUBLICATIONS

This section includes first (*) and/or corresponding (#) author papers.

D Wu*, J Dou*, X Chai*, C Bellis, A Wilm, CC Shih, WWJ Soon, N Bertin, CB Lin, CC Khor, M DeGiorgio, S Cheng, L Bao, N Karnani, WYK Hwang, S Davila, P Tan, A Shabbir, A Moh, EK Tan, JN Foo, LL Goh, KP Leong, RSY Foo, CSP Lam, AM Richards, CY Cheng, T Aung, TY Wong, HH Ng, J Liu#, **C Wang#** on behalf of the SG10K Consortium (2019). Large-scale whole-genome sequencing of three diverse Asian populations in Singapore. *Cell*, accepted.

J Dou*, B Sun*, X Sim, JD Hughes, DF Reilly, ES Tai, J Liu, **C Wang#** (2017). Estimation of kinship coefficient in structured and admixed populations using sparse sequencing data. *PLOS Genetics*, **13**: e1007021.

D Taliun#, S Chothani, S Schönherr, L Forer, M Boehnke, GR Abecasis, **C Wang#** (2017). LASER server: ancestry tracing with genotypes or sequence reads. *Bioinformatics*, **33**: 2056-2058.

H Chen*, **C Wang***, MP Conomos, AM Stilp, Z Li, T Sofer, AA Szpiro, W Chen, JM Brehm, JC Cledon, S Redline, GJ Papanicolaou, TA Thornton, CC Laurie, K Rice, X Lin (2016). Control for population structure and relatedness for binary traits in genetic association studies via logistic mixed models. *American Journal of Human Genetics*, **98**: 653-666.

C Wang#, X Zhan, L Liang, GR Abecasis, X Lin (2015). Improved ancestry estimation for both genotyping and sequencing data using projection Procrustes analysis and genotype imputation. *American Journal of Human Genetics*, **96**: 926-937.

C Wang#*, X Zhan*, J Bragg-Gresham, HM Kang, D Stambolian, E Chew, K Branham, J Heck-enlively, The FUSION Study, RS Fulton, RK Wilson, ER Mardis, X Lin, A Swaroop, S Zöllner, GR Abecasis# (2014). Ancestry estimation and control of population stratification for sequence-based association studies. *Nature Genetics*, **46**: 409-415.

X Zhan*, DE Larson*, **C Wang***, DC Koboldt, Y Sergeev, 52 other coauthors, ER Mardis, A Swaroop, GR Abecasis (2013). Identification of a rare coding variant in Complement 3 associated with age-related macular degeneration. *Nature Genetics* **45**: 1375-1379.

C Wang#, KB Schroeder, NA Rosenberg (2012). A maximum-likelihood method to correct for allelic dropout in microsatellite data with no replicate genotypes. *Genetics* **192**: 651-669.

C Wang#, S Zöllner, NA Rosenberg (2012). A quantitative comparison of the similarity between genes and geography in worldwide human populations. *PLOS Genetics* **8**: e1002886. [Featured in *Science* 337: 1151, 2012]

C Wang, ZA Szpiech, J Degnan, M Jakobsson, TJ Pemberton, JA Hardy, AB Singleton, NA Rosenberg (2010). Comparing spatial maps of human population-genetic variation using Procrustes analysis. *Statistical Applications in Genetics and Molecular Biology* **9**: 13.

COAUTHORED
PUBLICATIONS

This section includes middle author papers.

A Teumer, Y Li, S Ghasemi, BP Prins, M Wuttke, T Hermle, ..., **C Wang**, ..., AS Butterworth, AM Hung, C Pattaro, A Köttgen (2019). Genome-wide association meta-analyses and fine-mapping elucidate pathways influencing albuminuria. *Nature Communications*, in press.

K Huang, T Yang, J Xu, L Yang, J Zhao, X Zhang, C Bai, J Kang, P Ran, H Shen, F Wen, W Yao, T Sun, G Shan, Y Lin, G Xu, S Wu, **C Wang**, ..., T Wu, KF Chung, J He, Chen Wang for the China Pulmonary Health (CPH) Study Group (2019). Prevalence, risk factors and management of asthma in China: a national cross-sectional study. *Lancet*, **394**: 407-418.

M Wuttke, Y Li, M Li, KB Sieber, MF Feitosa, M Gorski, A Tin, L Wang, ..., **C Wang**, ..., K Stefansson, AM Hung, IM Heid, M Scholz, A Teumer, A Köttgen, C Pattaro (2019). A catalog of

genetic loci associated with kidney function from analyses of a million individuals. *Nature Genetics*, **51(6)**: 957-972.

H Chen, JE Huffman, JA Brody, **C Wang**, S Lee, ..., CC Laurie, AC Morrison, KM Rice, X Lin (2019). Efficient variant set mixed model association tests for continuous and binary traits in large-scale whole genome sequencing studies. *American Journal of Human Genetics*, **104(2)**: 260-274.

F Takeuchi, M Akiyama, N Matoba, T Katsuya, M Nakatochi, ..., **C Wang**, ..., T Nabika, M Yokota, Y Kamatani, M Kubo, N Kato (2018). Interethnic analyses of blood pressure loci in populations of East Asian and European descent. *Nature Communications*, **9(1)**: 5052.

H Chen, BE Cade, KJ Gleason, AC Bjornes, AM Stilp, ..., **C Wang**, ..., X Zhu, SR Sunyaev, R Saxena, X Lin, S Redlin (2018). Multi-ethnic meta-analysis identifies RAI1 as a possible obstructive sleep apnea related quantitative trait locus in men. *American Journal of Respiratory Cell and Molecular Biology*, **58**: 391-401.

J Liu, X Wan, **C Wang**, C Yang, X Zhou, C Yang (2017). LLR: A latent low-rank approach to colocalizing genetic risk variants in multiple GWAS. *Bioinformatics*, **33**: 3878-3886.

H Liu, Z Wang, Y Li, G Yu, X Fu, ..., **C Wang**, ..., S Chen, J Liu, F Zhang (2017). Genome-wide analysis of protein-coding variants in leprosy. *Journal of Investigative Dermatology*, **137**: 2544-2551.

X Wang, Z Zhang, M Nathan, T Cai, S Lee, **C Wang**, TW Yu, CA Walsh, X Lin (2017). Rare variant association test in family based sequencing studies. *Briefings in Bioinformatics*, **18**: 954-961.

W Dai, M Yang, **C Wang**, T Cai (2017). Sequence robust association test (SRAT) for familial data. *Biometrics*, **73**: 876-884

Z Wang, BC Henn, **C Wang**, Y Wei, L Su, R Sun, H Chen, PJ Wagner, Q Lu, X Lin, R Wright, D Bellinger, M Kile, M Mazumdar, MM Tellez-Rojo, L Schnaas, DC Christiani (2017). Genome-wide gene by lead exposure interaction analysis identifies UNC5D as a candidate gene for neurodevelopment. *Environmental Health*, **16**: 81.

BE Cade, H Chen, AM Stilp, KJ Gleason, T Sofer, ..., **C Wang**, PC Zee, CL Hanis, SR Sunyaev, SR Patel, CC Laurie, X Zhu, R Saxena, X Lin, S Redline (2016). Genetic Associations with Obstructive Sleep Apnea Traits in Hispanic/Latino Americans. *American Journal of Respiratory and Critical Care Medicine*, **194**: 886-897.

X Lin, S Lee, M Wu, **C Wang**, H Chen, Z Li, X Lin (2016). Test for rare variants by environment interactions in sequencing association studies. *Biometrics*, **72**: 156-164.

TJ Pemberton, **C Wang**, JZ Li, NA Rosenberg (2010). Inference of unexpected genetic relatedness among individuals in HapMap Phase III. *American Journal of Human Genetics* **87**: 457-464. [Featured in *Am J Hum Genet* 87: 447-448, 2010 and *Genetics* 186(2): NP, 2010]

JT Mosher, TJ Pemberton, K Harter, **C Wang**, EO Buzbas, P Dvorak, C Simon, SJ Morrison, NA Rosenberg (2010). Lack of population diversity in commonly used human embryonic stem-cell lines. *New England Journal of Medicine* **362**: 183-185. [Featured in *Nature* 462: 945, 2009]

NM Kopelman, L Stone, **C Wang**, D Gefel, MW Feldman, J Hillel, NA Rosenberg (2009). Genomic microsatellites identify shared Jewish ancestry intermediate between Middle Eastern and European populations. *BMC Genetics* **10**: 80.

L Huang, **C Wang**, NA Rosenberg (2009). The relationship between imputation error and statistical power in genetic association studies in diverse populations. *American Journal of Human Genetics* **85**: 692-698. [Featured in *Am J Hum Genet* 85: 539-540, 2009 & *Nat Rev Genet* 10: 817, 2009]

PROFESSIONAL ACTIVITIES	2009-Now: Member of the American Society of Human Genetics (ASHG) 2016-Now: Member of the International Genetic Epidemiology Society (IGES) 2009-Now: Reviewer for <i>American Journal of Human Genetics</i> , <i>Annals of Human Genetics</i> , <i>Bioinformatics</i> , <i>Biometrics</i> , <i>BMC Bioinformatics</i> , <i>BMC Genetics</i> , <i>Clinical Chemistry</i> , <i>Gene</i> , <i>Genetics</i> , <i>Genome Biology</i> , <i>Genome Research</i> , <i>Human Biology</i> , <i>Human Heredity</i> , <i>Journal of Human Genetics</i> , <i>Molecular Biology and Evolution</i> , <i>Molecular Ecology</i> , <i>PLoS Genetics</i> , <i>PLoS ONE</i> , <i>Proceedings of the National Academy of Sciences (PNAS)</i> , <i>Scientific Reports</i> 2017: Reviewer for Student Paper Award Competition, American Statistical Association (ASA) Section on Statistics in Genomics and Genetics
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