CURRICULUM VITAE

**Il-Youp Kwak**

**Work**

Department of Applied Statistics

College of Business & Economics

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**EDUCATION**

2014 PhD, Statistics, University of Wisconsin-Madison (Advisor: Karl W. Broman)

2008 MS, Statistics, Korea University, Seoul, Korea (Advisor: Myung-Hoe Huh)

2006 BS, Mathematics, Korea University, Seoul, Korea

**PROFESSIONAL POSITIONS**

2019- Assistant Professor, Department of Applied Statistics, College of Business & Economics, Chung-Ang University, Seoul, Korea

2017-2019 Staff Engineer, Samsung Research, Seoul R&D Campus, Seoul, Korea

2016-2017 Research Associate, Lillehei Heart Institute, University of Minnesota, US

2014-2016 Postdoctoral Associate, Division of Biostatistics School of Public Health, University of Minnesota, US

**RESEARCH INTERESTS**

* Real-world big data analysis: Develop new service that combines enterprise big data
* Biostatistics: statistical methodology and software development for bio-data analysis
* Development of Statistical Software tools
* Deep learning applications (focus on voice classifications)

**DATA COMPETITIONS**

2020.3 [Winner of a Biomedical Data Competition] Our team (Il-Youp Kwak and Wuming Gong) won both Subchallenges 2 and 3 of the Allen Institute Cell Lineage Reconstruction DREAM Challenge

2020.7 Our CAU-ET team (Yerin Lee, Soyoung Lim and Il-Youp Kwak) ranked 7th and 9th place at the IEEE AASP challenge on Detection and Classification of Acoustic Scenes and Event (DCASE) Task 1 subtask B and A

**PUBLICATIONS**

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# Working Manuscripts

Gong W, Raz O, Salvador-Martinez I, **Kwak IY**, Jones MG, Khodaverdian A, Zhang R, Rao S, Wang R, Yosef N, Hu J, Liu Z, Zhang H, Guan Y, Retkute R, Garry DJ, Peng J, Han L, Shang X, Chung V, Mason M, DREAM consortium, Shendure J, Elowitz M, Telfold M, Shapiro E, Meyer P (2020+) Benchmarked machine learning approaches for reconstructions of large cell lineage trees in C.elegans and mouse (Submitted)

Published Papers

*2021* **Kwak IY**, Kwag S, Lee J, Huh J, Jeon Y, Hwang J, Yoon J, Lee C. (2021) RESMAX: Detecting Voice Spoofing Attacks with Residual Network and Max Filter Map, 25th International Conference on Pattern Recognition (ICPR2020), Milan, Italy

*2020* Ahmed E, **Kwak IY**, Huh J, Kim I, Oh T, Kim H (2020) Void: A fast and light voice liveness detection system, USENIX Security Symposium, Boston, USA, 2020

Yoon HY, **Kwak IY** (2020) Association Modeling on Keyword and Abstract Data in Korean Port Research, Journal of Korea Trade, 25 (5), 71-86

*2019* **Kwak IY**, Huh J, Kim I, Han S, Yoon J. (2019) Voice Presentation Attack Detection through Text-Converted Voice Command Analysis, Proceedings of the 2019 CHI Conference on Human Factors in Computing Systems

Magli A, Baik J, Mills LJ, **Kwak IY**, Dillon BS, Gonzalez RM, Stafford DA, Swanson SA, Stewart R, Thomson JA, Garry DJ, Dynlacht BD, Perlingeiro R (2019) Time-dependent Pax3-mediated chromatin remodeling and cooperation with Six4 and Tead2 specify the skeletal myogenic lineage in developing mesoderm, *PLoS biology*, 17 (2), e3000153

Magli A, Baik J, Pota P, Cordero CO, **Kwak IY**, Garry DJ, Love PE, Dynlacht BD, Perlingeiro R (2019) Pax3 cooperates with Ldb1 to direct local chromosome architecture during myogenic lineage specification, *Nature Communications,* 10, 2316

*2018* Gong W, **Kwak IY**(co-1st author), Pota P, Koyano-Nakagawa N, Garry DJ. (2018) DrImpute: imputing dropout events in single cell RNA sequencing data, *BMC Bioinformatics*, 19 (1), 220

Gong W, **Kwak IY,** Koyano-Nakagawa N, Pan W, Garry DJ. (2018) TCM visualizes trajectories and cell populations from single cell data, *Nature Communications*, 9 (1), 2749

*2017* **Kwak IY,** Pan W. (2017) Gene- and pathway-based association tests for multiple traits with GWAS summary statistics. *Bioinformatics*, 33 (1), 64-71.

Chen Z, Zhu W, Bender I, Gong W, **Kwak IY**, Yellamilli A, Hodges TJ, Nemoto N, Zhang J, Garry DJ, Berlo JH (2017) Pathologic Stimulus Determines Lineage Commitment of Cardiac C-kit+ Cells. *Circulation*, circulationaha. 117.030137

*2016* Wei P, Cao Y, Zhang Y, **Kwak IY**, Xu Z, Crosby J, Boerwinkle E, Pan W. (2016) On Robust Association Testing for Quantitative Traits and Rare Variants. *G3*, 116.035485.

**Kwak IY,** Pan W. (2016) Adaptive Gene- and Pathway-Trait Association Testing with GWAS Summary Statistics. *Bioinformatics*, 32 (8), 1178-1184.

*2015* Pan W, **Kwak IY**, Wei P. (2015) A Powerful Pathway-Based Adaptive Test for Genetic Association with Common or Rare Variants. *The American Journal of Human Genetics*, 97:86-98

**Kwak IY** Moore CR, Spalding EP, Broman KW. (2015) Mapping quantitative trait loci underlying function-valued traits using functional principal component analysis and multi-trait mapping. *G3*, 6 (1), 79-86.

*2014* **Kwak IY,** Moore CR, Spalding EP and Broman KW. (2014) A simple regression-based method to map quantitative trait loci underlying function-valued phenotypes. *Genetics*, 197(4):1409-1416

*2013* Moore CR, Johnson LS, **Kwak IY**, Livny M, Broman KW, Spalding EP (2013) High-throughput computer vision introduces the time axis to a quantitative trait map of a plant growth response. *Genetics* 195:1077–1086

*2010* Billings T, Sargent EE, Szatkiewicz JP, Leahy N, **Kwak, IY**, Bektassova N, Walker M, Hassold T, Graber JH, Broman KW, Petkov PM (2010) Patterns of recombination activity on mouse chromosome 11 revealed by high resolution mapping. *PLoS One* 5:e15340

*2008* **Kwak IY**, Huh MH (2008) Andrews’ Plot for Extended Uses, *The Korean Communications in Statistics*, 15: 87–94

**Kwak IY**, Huh MH (2008) Parallel Coordinate Plots of Mixed-Type Data, *The Korean Communications in Statistics*, 15: 587–595

Park MR, **Kwak IY**, Huh MH (2008) Applications of Parallel Coordinate Plots for Visualizing Gene Expression Data, *The Korean Journal of Applied Statistics*, 21(6): 911–921

**SOFTWARE DEVELOPMENT**

R/DCLEAR An R package for R package for Distance based Cell LinEAge Reconstruction (DCLEAR). These codes are created during the participation of Cell Lineage Reconstruction DREAM challenge. Our team achieved the best performance on subchallenges 2 and 3 of the competition.

R/Drimpute An R package for imputing dropout events. Many statistical methods in cell type identification, visualization and lineage reconstruction do not account for dropout events ('PCAreduce', 'SC3', 'PCA', 't-SNE', 'Monocle', 'TSCAN', etc). 'DrImpute' can improve the performance of such software by imputing dropout events.

R/aSPU An R package for the (adaptive) Sum of Powered Score ('SPU' and 'aSPU') tests, inverse variance weighted Sum of Powered score ('SPUw' and 'aSPUw') tests and some gene-based and pathway based association tests (aSPUs, aSPUsPath, MTaSPUsSet, MTaSPUsSetPath, etc). (<https://cran.r-project.org/web/packages/aSPU/index.html>).

R/funqtl An R package with add-on functions for the R/qtl package to deal with QTL mapping for function-valued traits. ([github.com/ikwak2/funqtl](http://github.com/ikwak2/funqtl)).

R/xoi Tools for analyzing crossover interference. ([github.com/kbroman/xoi](http://github.com/kbroman/xoi)). Assisted Karl Broman to develop the package using C language.

R/APCP An R package that plots Andrews type parallel coordinate plot. ([github.com/ikwak2/APCP](http://github.com/ikwak2/APCP)).

**PRESENTATIONS**

*2020* RESMAX: Detecting Voice Spoofing Attacks with Residual Network and Max Filter Map, *Spring Conference,* *The Korean Statistical Society*, Cheonan, Korea

*2019* A survey on voice spoofing detection methods, *Autumn Conference,* *The Korean Statistical Society*, Seoul, Korea

*2019* Voice Presentation Attack Detection through Text-Converted Voice Command Analysis, *Spring Conference,* *The Korean Statistical Society*, Kangwon, Korea

*2019* Voice Presentation Attack Detection through Text-Converted Voice Command Analysis*, 2019 CHI Conference on Human Factors in Computing Systems*, Glasgow, UK

*2017* Gene- and pathway-based association tests for multiple traits with GWAS summary statistics, *2017 ENAR*

*spring meeting*, Washington, DC

*2016* Adaptive Gene- and Pathway-Trait Association Testing with GWAS Summary Statistics, *2016 ENAR*

*spring meeting*, Austin, TX

*2015* Regression-Based Methods to Map Quantitative Trait Loci Underlying Function-Valued Phenotypes, *2015*

*ENAR spring meeting*, Miami, FL

*2015* Adaptive Gene- and Pathway-Trait Association Testing with GWAS Summary Statistics, *2015 3rd Annual BICB Symposium*, Minneapolis, MN

*2013* QTL analysis with Function-valued Trait Data, *2013 ENAR Spring meeting*, Olando, FL

2008 Parallel Coordinate Plots of Mixed-type Data, *2008 Korea-Japan Student Conference*, Seoul, Korea

2007 Andrews’ Plot for Extended Uses, *Autumn Conference, The Korean Statistical Society*, Seoul, Korea

**POSTERS**

2021 RESMAX: Detecting Voice Spoofing Attacks with Residual Network and Max Filter Map, 25th International Conference on Pattern Recognition (ICPR2020), Milan, Italy

2016 Gene-based association tests for multiple traits with GWAS summary statistics, TC ASA Fall Research Meeting, Mounds View, MN

2015 Adaptive Gene- and Pathway-Trait Association Testing with GWAS Summary Statistics, TC ASA Fall Research Meeting, Rochester, MN

2013 QTL analysis with Function-valued Trait Data, Complex Trait Community annual meeting, Madison, WI

**JOURNAL MANUSCRIPT REVIEWS**

Molecular Ecology Resources, Scientific Reports, Biometrics, IEEE/ACM Transactions on Computational Biology and Bioinformatics, Bioinformatics, Journal of Korean Statistical Society, Communications for Statistical Applications and Methods, iScience, IEEE Transactions on Mobile Computing