# Junghyun Jung

https://junghyunjj.github.io/

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Email: junghyun.jung@cshs.org

# **EDUCATION**

• Dongguk University

Seoul, Korea

Ph.D. in Life Science

Sept. 2014 - Feb. 2020

Mentor: Dr. Wonhee Jang

**Dissertation project:** Bioinformatics-based approaches for identifying novel genetic markers

• Dongguk University

Seoul, Korea

M.S. in Life Science

Mar. 2012 - Aug. 2014

Mentor: Dr. Wonhee Jang

**Dissertation project:** Studies on the *in silico* analyzing methods related to toxic effects and mechanisms of chemicals

• Dongguk University

Seoul, Korea

B.S. in Life Science

Mar. 2007 - Feb. 2012

# Professional and Academic Experience

#### • Cedars-Sinai Medical Center

Los Angeles, CA, USA

Research Data Scientist at Department of Computational Biomedicine

Jul. 2023 - present

Mentor: Dr. Kyoung Jae Won

- Develop novel computational tools
  - Develop novel computational tools for spatial omic datasets.
  - Develop novel statistical methods for single-cell genetics datasets (sc-eQTL).
- Large-scale integrative analysis of complex traits
  - Perform multi-omic data analysis for complex diseases.
  - Perform spatiotemporal transcriptomic data analysis for complex diseases

#### • University of Southern California

Los Angeles, CA, USA

Postdoctoral Researcher at Center for Genetic Epidemiology, Keck School of Medicine Aug. 2021 - Jun. 2023 Mentor: Dr. Nicholas Mancuso

- Large-scale integrative analysis of complex disease
  - Perform integrative analysis using GWAS with molecular QTL (molQTL) data for TWAS analysis.
  - Perform Single-cell RNA/ATAC/TCR-seq data analysis.
- o Inference of negative selection in the evolutionary architecture
  - Develop a novel statistical method using Variational Bayesian methods.
  - Understand the landscape of evolutionary architectures across many molecular phenotypes.

## • University of Southern California

Los Angeles, CA, USA

Postdoctoral Researcher at Center for Genetic Epidemiology, Keck School of Medicine Aug. 2020 - Jul. 2021 Mentor: Dr. Nicholas Mancuso

- o Genome-wide association analysis (GWAS) of multiple phenotypes in multi-ancestry
  - Perform multivariate GWAS analysis by analyzing correlated cytokines phenotypes.
  - Perform multi-ancestry GWAS analysis by analyzing correlated cytokines phenotypes.
- o Inference of negative selection in the evolutionary architecture
  - Duties similar to above.

Postdoctoral Researcher at Titus Family Department of Clinical Pharmacy, School of Pharmacy Mentor: Dr. Serghei Mangul

- Genome/Exome-wide association study to identify common/rare variants influencing COVID-19 outcomes using Middle Eastern cohort
  - Analyze Whole-Exome seq (WES) data to identify rare variants using the Saudi Human Genome Program cohort.
  - Perform GWAS analysis to identify common variants using the Saudi Human Genome Program cohort.
- o Rigorous benchmarking of HLA callers for RNA sequencing data
  - Evaluate performances of multiple HLA callers using gold standard data.
  - Consensus HLA typing based on multiple HLA callers using Genotype-Tissue Expression (GTEx) data.
  - HLA typing using multiple ancestry data.

# • Dongguk University

Seoul, Korea

Postdoctoral Researcher at Department of Computer Science and Engineering

Mar. 2020 - Jul. 2020

Mentor: Dr. Jong Wha J. Joo

- Set-based analysis to increase the power of association testing
  - Develop a statistical method to improve statistical power using multiple causal variants.
  - Multivariate phenotype analysis by analyzing many phenotypes simultaneously.

# • Dongguk University

Seoul, Korea

Instructor at Convergence Software Institute Dongguk (Biological Information Software Convergence) Mar. 2018 - Aug. 2020

# • Dongguk University

Teaching assistant at Department of Life Science

Seoul, Korea

Sept. 2012 - Aug. 2013

# RESEARCH TECHNIQUE

#### • Computational Skills

- NGS data analysis: WES, RNA-seq, ATAC-seq, Chip-seq, and scRNA/scATAC/scTCR-seq data analysis
- Spatial omics analysis: Spatial transcriptome data analysis (Visium, Slide-seq, and Stereo-seq)
- Micoarray data analysis: SNP and expression array data analysis
- High performance computing: Slurm and Sun Grid Engine (SGE)
- Programming languages:: R, Python, and Bash
- Other software applications: Git, LaTeX, SPSS, and ImageJ

#### • Experimental Skills

- Mammalian cell culture
- Animal handling (mouse)

## Publications

- [1] W. Oh, **J. Jung**, and J. W. J. Joo, "MR-GGI: accurate inference of gene–gene interactions using Mendelian randomization," *BMC Bioinformatics*, vol. 25, no. 1, pp. 1–16, 2024.
- [2] J. Song, D. Kim, **J. Jung**, E. Choi, et al., "Elucidating immunological characteristics of the adenoma-carcinoma sequence in colorectal cancer patients in South Korea using a bioinformatics approach," Scientific reports, vol. 14, no. 10105, 2024.
- [3] D. Kim, J. Song, S. Mangul, N. Mancuso, <u>J. Jung</u>\*, and W. Jang\*, "Large-scale integrative analysis of juvenile idiopathic arthritis for new insight into its pathogenesis," *Arthritis Research & Therapy*, vol. 26, no. 1, p. 47, 2024.

- [4] Z. Zhang, **J. Jung**, A. Kim, N. Suboc, S. Gazal, and N. Mancuso, "A scalable approach to characterize pleiotropy across thousands of human diseases and complex traits using GWAS summary statistics," *The American Journal of Human Genetics*, vol. 110, no. 11, pp. 1863–1874, 2023.
- [5] <u>J. Jung</u>\*, Z. Lu, A. de Smith, and N. Mancuso\*, "Novel insight into the etiology of ischemic stroke gained by integrative multiome-wide association study," *Human Molecular Genetics*, vol. 33, no. 2, pp. 170–181, 2024.
- [6] G. Butler-Laporte, G. Povysil, J. A. Kosmicki, E. T. Cirulli, T. Drivas, S. Furini, C. Saad, A. Schmidt, P. Olszewski, U. Korotko, J. Jung, et al., "Exome-wide association study to identify rare variants influencing COVID-19 outcomes: Results from the Host Genetics Initiative," PLoS Genetics, vol. 18, no. 11, p. e1010367, 2022.
- [7] W. group lead Andrews Shea J. 6 Kanai Masahiro 3 Cordioli Mattia 7, M. analyses team lead, M. analyses team members: PHEWAS Polimanti Renato 1, M. analyses team members: Mendelian randomization Harerimana Nadia 8, M. analyses team members: methods development Pirinen Matti 7, g. p. Manuscript analyses team members: PC projection, P. management lead Liao Rachel G. 3, P. management support Chwialkowska Karolina 9 Trankiem Amy 3 Balaconis Mary K. 3, H. Nguyen, et al., "A first update on mapping the human genetic architecture of COVID-19," Nature, vol. 608, no. 7921, pp. E1–E10, 2022.
- [8] Y. Shim, M. Shin, **J. Jung**, B. Koo, and W. Jang, "An in-silico approach to studying a very rare neurodegenerative disease using a disease with higher prevalence with shared pathways and genes: Cerebral adrenoleukodystrophy and Alzheimer's disease," *Frontiers in molecular neuroscience*, vol. 15, 2022.
- [9] J. Song, D. Kim, S. Lee, **J. Jung**, J. W. J. Joo, and W. Jang, "Integrative transcriptome-wide analysis of atopic dermatitis for drug repositioning," *Communications biology*, vol. 5, no. 1, pp. 1–13, 2022.
- [10] J. L. Wiemels, R. Wang, M. Zhou, H. Hansen, R. Gallant, J. Jung, N. Mancuso, A. J. De Smith, C. Metayer, S. C. Kogan, et al., "Cytomegalovirus proteins, maternal pregnancy cytokines, and their impact on neonatal immune cytokine profiles and acute lymphoblastic leukemogenesis in children," *Haematologica*, 2022.
- [11] A. Kousathanas, E. Pairo-Castineira, K. Rawlik, A. Stuckey, C. A. Odhams, S. Walker, C. D. Russell, T. Malinauskas, Y. Wu, J. Millar, J. Jung, et al., "Whole genome sequencing reveals host factors underlying critical Covid-19," Nature, pp. 1–10, 2022.
- [12] C. Fallerini, N. Picchiotti, M. Baldassarri, K. Zguro, S. Daga, F. Fava, E. Benetti, S. Amitrano, M. Bruttini, M. Palmieri, J. Jung, et al., "Common, low-frequency, rare, and ultra-rare coding variants contribute to COVID-19 severity," Human genetics, pp. 1–27, 2021.
- [13] M. D'Antonio, J. P. Nguyen, T. D. Arthur, H. Matsui, B. M. Neale, M. Daly, A. Ganna, C. Stevens, G. A. Pathak, S. J. Andrews, J. Jung, et al., "SARS-CoV-2 susceptibility and COVID-19 disease severity are associated with genetic variants affecting gene expression in a variety of tissues," Cell reports, vol. 37, no. 7, p. 110020, 2021.
- [14] B. Lee, M. K. Shin, I.-W. Hwang, J. Jung, Y. J. Shim, G. W. Kim, S. T. Kim, W. Jang, and J.-S. Sung, "A Deep Learning Approach with Data Augmentation to Predict Novel Spider Neurotoxic Peptides," International Journal of Molecular Sciences, vol. 22, no. 22, p. 12291, 2021.
- [15] G. Povysil, G. Butler-Laporte, N. Shang, C. Wang, A. Khan, M. Alaamery, T. Nakanishi, S. Zhou, V. Forgetta, R. J. Eveleigh, J. Jung, et al., "Rare loss-of-function variants in type I IFN immunity genes are not associated with severe COVID-19," The Journal of Clinical Investigation, vol. 131, no. 14, 2021.
- [16] F. Hormozdiari<sup>†</sup>, <u>J. Jung</u><sup>†</sup>, E. Eskin, and J. W. J. Joo, "MARS: leveraging allelic heterogeneity to increase power of association testing," *Genome biology*, vol. 22, pp. 1–26, April 2021.
- [17] D. Kim, J. Song, S. Lee, **J. Jung**, and W. Jang, "An Integrative Transcriptomic Analysis of Systemic Juvenile Idiopathic Arthritis for Identifying Potential Genetic Markers and Drug Candidates," *International Journal of Molecular Sciences*, vol. 22, no. 2, p. 712, 2021.

- [18] J. H. Choi, T. Kim, **J. Jung**, and J. W. J. Joo, "Fully automated web-based tool for identifying regulatory hotspots," *BMC genomics*, vol. 21, no. 10, pp. 1–7, 2020.
- [19] G. J. Lee, S. M. Park, J. Jung, and J. W. J. Joo, "A Fully Automated Parallel-Processing R Package for High-Dimensional Multiple-Phenotype Analysis Considering Population Structure," *International Journal of Fuzzy Logic and Intelligent Systems*, vol. 20, no. 3, pp. 219–226, 2020.
- [20] J. Song, D. Kim, J. Hong, G. W. Kim, J. Jung, S. Park, H. J. Park, J. W. J. Joo, and W. Jang, "Meta-analysis of polymyositis and dermatomyositis microarray data reveals novel genetic biomarkers," Genes, vol. 10, no. 11, p. 864, 2019.
- [21] <u>J. Jung</u>, G. W. Kim, B. Lee, J. W. J. Joo, and W. Jang, "Integrative genomic and transcriptomic analysis of genetic markers in Dupuytren's disease," *BMC medical genomics*, vol. 12, no. 5, pp. 1–10, 2019.
- [22] E. Lee, H. Jeon, M. Lee, J. Ryu, C. Kang, S. Kim, **J. Jung**, and Y. Kwon, "Molecular origin of AuNPs-induced cytotoxicity and mechanistic study," *Scientific reports*, vol. 9, no. 1, pp. 1–13, 2019.
- [23] <u>J. Jung</u>, G. W. Kim, W. Lee, C. Mok, S. H. Chung, and W. Jang, "Meta-and cross-species analyses of insulin resistance based on gene expression datasets in human white adipose tissues," *Scientific reports*, vol. 8, no. 1, pp. 1–13, 2018.
- [24] H. Kim, J. Yoo, J. Shin, Y. Chang, J. Jung, D.-G. Jo, J. Kim, W. Jang, C. J. Lengner, B.-S. Kim, et al., "Modelling APOE 3/4 allele-associated sporadic Alzheimer's disease in an induced neuron," Brain, vol. 140, no. 8, pp. 2193–2209, 2017.
- [25] J. Yoo, E. Lee, H. Y. Kim, D.-H. Youn, J. Jung, H. Kim, Y. Chang, W. Lee, J. Shin, S. Baek, et al., "Electromagnetized gold nanoparticles mediate direct lineage reprogramming into induced dopamine neurons in vivo for Parkinson's disease therapy," Nature nanotechnology, vol. 12, no. 10, pp. 1006–1014, 2017.
- [26] J. Jung, K. Hah, W. Lee, and W. Jang, "Meta-analysis of microarray datasets for the risk assessment of coplanar polychlorinated biphenyl 77 (PCB77) on human health," *Toxicology and Environmental Health Sciences*, vol. 9, no. 2, pp. 161–168, 2017.
- [27] J. Jung, C. Mok, W. Lee, and W. Jang, "Meta-analysis of microarray and RNA-Seq gene expression datasets for carcinogenic risk: An assessment of Bisphenol A," *Molecular & Cellular Toxicology*, vol. 13, no. 2, pp. 239–249, 2017.
- [28] E. Lee, J. Jung, D. Jung, C. S. Mok, H. Jeon, C.-S. Park, W. Jang, and Y. Kwon, "Inhibitory effects of novel SphK2 inhibitors on migration of cancer cells," Anti-Cancer Agents in Medicinal Chemistry (Formerly Current Medicinal Chemistry-Anti-Cancer Agents), vol. 17, no. 12, pp. 1689–1697, 2017.
- [29] H. G. Kang, D.-H. Kim, S.-J. Kim, Y. Cho, J. Jung, W. Jang, and K.-H. Chun, "Galectin-3 supports stemness in ovarian cancer stem cells by activation of the Notch1 intracellular domain," *Oncotarget*, vol. 7, no. 42, p. 68229, 2016.

# Conference Presentations - Oral Presentation

- Z. Zhang, J. Jung, N. Suboc, S. Gazal, N. Mancuso. A variational Bayesian approach to characterize pleiotropic components across thousands of human diseases and complex traits using GWAS summary statistics. ASHG, Los Angeles, CA, USA, 2022
- J. Jung, G.W. Kim, B. Lee, J.W.J. Joo., W. Jang. Integrative genomic and transcriptomic analysis for genetic markers in Dupuytren's disease. TBC/BIOINFO 2018. Seoul, Korea, 2018
- J. Jung, G.W. Kim, J.H. Park, S.H. Chung, W. Jang . Meta-analysis of insulin resistance based on gene expression datasets in human adipose tissues. The 13th international congress on obesity (ICO). Vancouver, Canada, 2016.

• G.W. Kim, **J. Jung**, H.K. Jo, K.J. Jeong, W. Jang, S.H. Chung. Elucidation of suppressive mechanism of betulinic acid on adipogenesis using global gene expression profiling. The 13th international congress on obesity (ICO). Vancouver, Canada, 2016.

# Conference Presentations - Poster Presentation

- J. Jung, Z. Lu, N. Mancuso. Novel insight into the etiology of ischemic stroke gained by integrating human transcriptome-wide association study with rodent expression data. ASHG, Virtual, 2021
- Z. Zhang, J. Jung, N. Suboc, S. Gazal, N. Mancuso. A variational Bayesian approach to characterize latent genetic components using GWAS summary statistics. ASHG, Virtual, 2021
- Z. Lu, **J. Jung**, S. Gazal, N. Mancuso. Residual proteome-wide association study identifies genes for blood-related traits. ASHG, Virtual, 2021
- J. Jung, R. Brown, W. Jang, B. Pasaniuc, J.W.J. Joo and E. Eskin. GGmend: A Mendelian randomization method for finding gene-on-gene regulatory effects in the presence of unobserved confounders. 23th RECOMB, Washington, D.C., USA, 2019.
- F. Hormozdiari, J. Jung, E. Eskin and J.W.J. Joo. Leveraging allelic heterogeneity to increase the power of association testing. 23th RECOMB, Washington, D.C., USA, 2019.
- J. Song, W. Jang, J. Jung, D.E. Kim and J.Y. Hong W. Jang. Meta-analysis of polymyositis and dermatomyositis microarray data sets. 23th RECOMB, Washington, D.C., USA, 2019.
- J. Jung, G.W. Kim, J.H. Park, S.H. Chung, W. Jang. Cross-species analysis reveals molecular mechanisms of insulin resistance in white adipose tissues. 12th International Conference on Toxicogenomics (ICT), Seoul, Korea, 2016.
- J. Jung, G.W. Kim, J.H. Park, S.H. Chung, W. Jang. Integrative cross-species analysis of insulin resistance expression profile in white adipose tissues. 9th International Conference on Environmental Health Sciences (ICoEHS), Incheon, Korea, 2016.
- J. Jung, J.H. Park, W. Jang. Meta-analysis of microarray gene expression datasets for silver nanoparticles. 11th International Conference on Toxicogenomics (ICT), Seoul, Korea, 2015.
- J. Jung, J.H. Park, W. Jang. Meta-analysis of microarray and RNA-Seq gene expression datasets for carcinogenic risk assessment of Bisphenol A. 8th International Conference on Environmental Health Sciences (ICoEHS), Incheon, Korea, 2015.
- J. Jung, J.H. Park, W. Jang. Meta-analysis of microarray gene expression datasets for in silico risk assessment of Bisphenol A. 6th International Conference on Environmental Health Sciences (ICoEHS), Incheon, Korea, 2013.

## PATENTS

- Composition for diagnosing Insulin Resistance and its uses. 2018.03.21. (1020180032865)
- Diagnostic kit comprising biomarkers for bisphenol A exposure diagnosis. 2018.12.21 (1019336950000)
- Diagnostic kit comprising biomarkers for genistein exposure diagnosis. 2018.12.21 (1019336940000)
- Diagnostic kit comprising biomarkers for polychlorinated biphenyl 77 exposure diagnosis. 2018.12.21 (1019336920000)
- Genetic markers for diagnosing juvenile idiopathic arthritis and diagnosis method using these markers. 2021.09.02 (1020210116775)

# Honors and Awards

- 2014 Dongguk Global Researcher Scholarship
- 13th International Conference on Toxicogenomics (ICT), Best poster, The Korea Society of Toxicogenomics and Toxicoproteomics, 2016
- 9th International Conference on Environmental Health Sciences (ICoEHS), Best poster, Society of Environmental Risk Assessment and Health Science, 2016
- 6th International Conference on Environmental Health Sciences (ICoEHS), Best poster, Society of Environmental Risk Assessment and Health Science, 2013

# TEACHING EXPERIENCE

• Invited lecturer on Statistical Methods in Human Genetics (PM 570)

University of Southern California (spring 2023)

I introduce cutting-edge tools for genome-wide association and genetic perturbations analysis that have been recently published.

• Workshop on Genomic Data Analysis and Deep Learning

LINC+ Donguk University (Winter 2019)

A 1-month workshop introducing attendees to the basics of Omics data analysis. Topics include preprocessing of RNA-seq and expression microarray data, differentially expressed genes (DEGs) analysis and functional enrichment analysis.

• Instructor for Big Data Processing/Analysis using Public Databases (BSI4002-01 and BSI4001-01)

Convergence Software Institute Dongguk (2018 - 2020)

This course provides the basic principles of computer analysis programming and the basics of biological data analysis using Python and R.

## PROFESSIONAL SERVICE

• Review Editor: Frontiers in Molecular Bioscience (June 2023 -)

Review editor for Frontiers in Molecular Bioscience (RNA Networks and Biology section).

• Reviews

Genome medicine, Translational Stroke Research, BMC genomics, Scientific Reports, Molecular Neurobiology, European Journal of Medical Research