Hyundoo Jeong

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2007

| CURRENT POSITION | Chosun University, Gwangju, South Korea | |
|-----------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------|
| | Assistant professor, Electronic Engineering Sept. 20 | 018 - Present |
| EDUCATION & TRAINING | Baylor College of Medicine, Houston, Texas | |
| | Postdoctoral research associate Aug. 2017 | - Aug. 2018 |
| | • Advisors: Dr. Zhandong Liu and Dr. Joshua M. Shulman | |
| | Texas A&M University, College Station, Texas | |
| | Ph.D., Electrical and Computer Engineering | Aug. 2017 |
| | Dissertation: Probabilistic random walk models for comparative network analysis Advisors: Dr. Byung-Jun Yoon and Dr. Xiaoning Qian | |
| | Seoul National University, Seoul, South Korea | |
| | M.S., Electrical Engineering and Computer Science | Feb. 2009 |
| | Thesis: Resource allocation for multiuser OFDM relay networks a constraints Advisor: Dr. Jae Hong Lee | with fairness |
| | Inha University, Incheon, South Korea | |
| | B.S., Electronic Engineering Magna Cum Laude (Ranked in 2 out of 199 students) | Feb. 2007 |
| RESEARCH INTERESTS | Signal processing, machine learning, probabilistic graphical models, bioinformatics, single cell genomics, computational network biology, statistical network analysis, image processing, and large-scale data analysis. | |
| Honors and Awards | Young Scholar Award, Korean Institute of Intelligent Systems | Fall 2018 |
| | Travel Grants | M 0015 |
| | 14th annual MCBIOS conference, Little Rock, AR NSF supported grant for 3rd CNB-MAC workshop, Seattle, WA | Mar. 2017 Oct. 2016 |
| | • 13th annual MCBIOS conference, Memphis, TN | Mar. 2016 |
| | • NSF supported grant for 2nd CNB-MAC workshop, Atlanta, GA | Sept. 2015 |
| | NSF supported grant for XSEDE15 conference, St. Louis, MO 12th annual MCBIOS conference, Little Rock, AR | July 2015 Mar. 2015 |
| | Scholarship | Wiai. 2010 |
| | Graduate research assistantship, Texas A&M University Sept. 2013 to Aug. 2017 \$1,000 scholarship with an in-state tuition waive, Texas A&M University 2012 Samsung education scholarship 2008 BK21 (Brain Korea) research scholarships, National Research Foundation of Korea | |
| | 2007, 2008 Honor scholarships, Inha University English scholarship, Inha University | 005 and 2006 2005 |

Magna Cum Laude, Inha University

RESEARCH EXPERIENCE

Jan and Dan Duncan Neurological Research Institute

Houston, TX

Postdoctoral Research Associate

Aug. 2017 to Aug. 2018

- Developing single cell RNA sequencing deconvolution methods
- Developing an imputation method for single cell RNA-seq.
- Developing single cell embedding and 3D geometry reconstruction algorithms

Genomic Signal Processing Lab., Texas A&M University College Station, TX Research Assistant Sept. 2013 to Aug. 2017

- Developed novel random walk models to estimate the node-to-node correspondence across different networks (graphs).
- Developed global network alignment algorithms to identify homologous proteins in multiple protein-protein interaction (PPI) networks.
- Developed network querying algorithms to identify the conserved network modules in large-scale PPI networks.
- Analyzed statistical features of biological networks to construct benchmark datasets to assess the performance of network alignment and querying algorithms.
- Analyzed NGS (next-generation sequencing) data to identify differentially expressed genes with a statistical significance, (target species: *Mus musculus*).

AgriLife Research, Texas A&M University

College Station, TX

Research Assistant

Sept. 2013 to Aug. 2014

- Plant bioinformatics graduate training program
- Performed a downstream analysis for NGS data.
- Developed a quality assessment pipeline for de novo transcriptome assembly for improved gene annotations.

Institute of New Media & Communications

Seoul, South Korea

Research Assistant

Mar. 2007 to Jan. 2009

- Performed research on wireless communications (PHY layer).
- Developed a resource allocation algorithm for multiuser OFDM relay networks.
- Involved projects:
 - Adaptive Resource Allocation for Cooperative Diversity LG Electronics, 2008
 - Iterative Multiuser Detection for IDMA systems LG Electronics, 2007
 - Space Time Coded Cooperative Diversity

Work Experience

Patent Information Promotion Cente

Seoul, South Korea

KOSEF, 2007 to 2008

Patent Search Professional

Feb. 2012 to June 2012

- Searched and reviewed technical documents, and made review reports to assess the novelty of the patents.
- Specific areas: IPC H04 (electric communication technique) and G06 (computing; calculating; counting).

Samsung Electronics

Suwon, South Korea

Software Engineer at Mobile Communication Division

Feb. 2009 to Mar. 2011

- Developed smartphone system software for application processor based on Qualcomm Snapdragon chipset.
- Developed Windows phone device drivers: display (LCD and AMOLED), notification, proximity and ambient light sensor.
- Developed test applications to find hardware defectives in the manufacturing process.

TECHNICAL SKILLS

- Knowledge of signal processing, statistics, machine learning, bioinformatics, and communication theory
- Experienced in high performance computing (Linux CentOS 6, IBM x86 HPC Cluster, PBS/Torque and SLURM)
- Programming languages: Proficient in MATLAB, R, Rcpp, Python, and C/C++
- Bioinformatics tools: DEseq, EdgeR, Trinity, Oases, TopHat, Samtools, and IGV

JOURNAL IN PREPARATION

- 1. H.-M. Woo[†], **H. Jeong**[†], and B.-J. Yoon "Comprehensive updates in network synthesis models to benchmark comparative network analysis," in preparation. [†]Equally contributed
- 2. **H. Jeong** and Z. Liu, "PRIME: Probabilistic imputation and smoothing for single cell RNA sequencing," will be submitted. (Manuscript in preparation.)
- 3. C.-C Chen, **H. Jeong**, X. Qian, and B.-J Yoon, "TOPAS: network-based structural alignment of RNA sequences," Submitted to Bioinformatics (Under reviewing)
- 4. Y. Wang[†], **H. Jeong**[†], X. Qian, and B.-J Yoon, "ClusterM: a scalable algorithm for computational prediction of conserved protein complexes across multiple protein interaction networks," submitted to Science Advances (Under reviewing) [†]Equally contributed

JOURNAL PUBLICATIONS

- 1. N. Ghaffari[†], O. A. Arshad[†], **H. Jeong**[†], J. Thiltges, M.F. Criscitiello, B.-J. Yoon, A. Datta, and C. D. Johnson, "Examining de novo transcriptome assemblies via a quality assessment pipeline," IEEE/ACM Transaction on Computational Biology and Bioinformatics, vol.15, no.2, pp. 494 505, Mar. 2018. [†]Equally contributed
- 2. **H. Jeong**, X. Qian, and B.-J Yoon, "CUFID-query: random walk based network querying through network flow estimation," BMC Bioinformatics, 18(Suppl. 14):12, 2017.
- 3. **H. Jeong** and B.-J Yoon, "SEQUOIA: significance enhanced network querying through context-sensitive random walk and minimization of network conductance," BMC Systems Biology, 11(Suppl. 3):20, 2017.
- H. Jeong, X. Qian, and B.-J Yoon, "Effective comparative analysis of proteinprotein interaction networks by measuring the steady state network flow using a Markov model," BMC Bioinformatics, 17(Suppl. 13):395, 2016.
- H. Jeong and B.-J Yoon, "Effective estimation of node-to-node correspondence between different graphs," IEEE Signal Processing Letters, vol. 22, no. 6, pp. 661-665, June 2015.
- 6. **H. Jeong** and B.-J Yoon, "Accurate multiple network alignment through context-sensitive random walk," BMC Systems Biology, 9(Suppl. 1):S7, 2015.

Conference Publications

- 1. H.-M. Woo[†], **H. Jeong**[†], and B.-J Yoon, "Comprehensive updates in network synthesis models to create an improved benchmark for network alignment algorithms," 9th ACM-BCB, Washington, DC, USA, Aug., 2018. [†]Equally contributed
- 2. **H. Jeong** and B.-J Yoon, "SEQUOIA: Significance enhanced network querying through context-sensitive random walk and minimization of network conductance," 7th ACM-BCB, Seattle, WA, Oct., 2016.
- 3. **H. Jeong** and B.-J Yoon, "Identifying molecular complexes in biological networks through context sensitive network querying," 6th ACM-BCB, Atlanta, GA, Sept. 9, 2015.
- 4. **H. Jeong** and B.-J Yoon, "Accurate multiple network alignment through context-sensitive random walk," Asian Pacific Bioinformatics Conference (APBC) 2015, Hsinchu, Taiwan, Jan. 2015.
- 5. N. Ghaffari[†], O. A. Arshad[†], **H. Jeong**[†], et al., "De novo transcriptome assemblies and annotation for pacific whiteleg shrimp," IEEE Global Conference on Signal and Information Processing (GlobalSIP) 2014, Atlanta, GA, USA, Dec. 2014. [†]Equally contributed

- H. Jeong and B.-J Yoon, "A novel context-sensitive random walk model for estimating node correspondence between two biological networks," 5th ACM-BCB, Newport Beach, CA, USA, Sept., 2014.
- H. Jeong, J. H. Lee, and H. Seo, "Resource allocation for uplink multiuser OFDM relay networks with fairness constraints," IEEE Vehicular Technology Conference (VTC) 2009 Spring, Barcelona, Spain, Apr. 2009.

INVITED TALKS

ORAL & POSTER PRESENTATIONS

- 1. **H. Jeong**, "Network-based signal processing in bioinformatics," Korean Institute of Intelligent Systems Fall Conference, Wonju, S. Korea 2018.
- H. Jeong, X. Qian, and B.-J Yoon, "CUFID-query: Accurate network querying through estimation of network flow between query and target networks based on random walk," 14th Annual Conference of the MidSouth Computational Biology and Bioinformatics Society (MCBIOS), Little Rock, AR, USA, Mar. 2017.
- H. Jeong, X. Qian, and B.-J Yoon, "CUFID-multialign: Enhanced multiple alignment of biological networks through estimation of steady-state flow using a random walk model," 14th Annual Conference of the MidSouth Computational Biology and Bioinformatics Society (MCBIOS), Little Rock, AR, USA, Mar. 2017.
- 3. H. Jeong and B.-J Yoon, "SEQUOIA: Significance enhanced network querying through context-sensitive random walk and minimization of network conductance," 3rd Int'l Workshop on Computational Network Biology: Modeling, Analysis, and Control (CNB-MAC), Seattle, WA, USA, Oct. 2016.
- 4. B.-J Yoon, and **H. Jeong**, "Predicting potential functional module in biological networks through context-sensitive random walk based network querying," Qatar Foundation-Annual Research Conference (QF-ARC), Doha, Qatar, Mar. 2016.
- 5. **H. Jeong** and B.-J Yoon, "Effective comparative analysis of protein-protein interaction networks by measuring the steady state network flow using a Markov model," 13th Annual Conference of the MidSouth Computational Biology and Bioinformatics Society (MCBIOS), Memphis, TN, USA, Mar. 2016.
- C.-C Chen, H. Jeong, X. Qian, and B.-J Yoon, "Pairwise global structural alignment for RNA sequences through topological networks," 13th Annual Conference of the MidSouth Computational Biology and Bioinformatics Society (MCBIOS), Memphis, TN, USA, Mar. 2016.
- H. Jeong and B.-J. Yoon, "Identifying molecular complexes in biological networks through context sensitive network querying," 2nd Int'l Workshop on Computational Network Biology: Modeling, Analysis, and Control (CNB-MAC), Atlanta, GA, USA, Sep. 2015.
- H. Jeong, O. A. Arshad, N. Ghaffari, J. Thiltges, P. Blood, M. F. Criscitiello, B.-J. Yoon, A. Datta, and C. D. Johnson, "Comparative de novo transcriptome assemblies for Pacific whiteleg shrimp empowered by XSEDE resources," Extreme Science Engineering Discovery Environment (XSEDE), Saint Louis, MO, USA, July. 2015.
- 9. **H. Jeong** and B.-J Yoon, "Querying networks using context-sensitive random walk," 12th Annual Conference of the MidSouth Computational Biology and Bioinformatics Society (MCBIOS), Little Rock, AR, USA, Mar. 2015.
- H. Jeong and B.-J. Yoon, "A novel context-sensitive random walk model for estimating node correspondence between two biological networks," Int'l Workshop on Computational Network Biology: Modeling, Analysis, and Control (CNB-MAC), Newport Beach, CA, USA, Sep. 2014.
- 11. **H. Jeong** and B.-J. Yoon, "A novel random walk model for comparing protein interactions networks," IEEE Int'l. Workshop on Genomic Signal Processing and Statistics (GENSIPS) 2013, Houston, TX, USA, Nov. 2013.

Patents

1. J. H. Lee and H. Jeong et. al "Method of allocating resources for cooperative diversity," 10-1523098, South Korea.

SERVICE

Reviewer

IEEE/ACM Transactions on Computational Biology and Bioinformatics, PLOS ONE, BMC Bioinformatics, Pattern Recognition Letters, ACM-BCB (2017), EURASIP Journal on Bioinformatics and Systems Biology, Journal of Biological Systems, and Communications for Statistical Applications and Methods.

Other

Korean Student Association at Texas A&M University 2014 - 2017 Student volunteer, XSEDE 2015 2015

Teaching

Fall 2018

EXPERIENCE

- Electronic Circuit 2
- Introduction to Programming (Python)
- Advanced Programming (C)
- Digital Logic Lab.

GRADUATE LEVEL Texas A&M University

Coursework

Pattern Recognition, Machine Learning with Networks, Neural Networks, Stochastic Systems, Introduction to Optimization, Probabilistic Graphical Model, Graph Theory, Introduction to Classical Analysis, Linear Network Analysis, Statistics in Research I, Engineering Applications to Genomics, Algorithm for Structural Bioinformatics, Digital Image Processing, Advanced Discrete Time Systems, and Mobile Wireless Network.

Seoul National University

Stochastic and Random Processes, Information Theory, Detection and Estimation, Channel Coding Theory, Adaptive Signal Processing, Multi-Rate Signal Processing, Advanced Digital Communication, Wireless and Satellite Communications, and Advanced Wireless Communication.

Last updated: November 2, 2018