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EDUCATION

- Ph.D.** Electronic Engineering (focus on bioinformatics and computational biology), 2013
City University of Hong Kong, Hong Kong
Advisor: Prof. Hong Yan
Thesis: Alpha shape based methods for analysis and prediction of biomolecular interactions.
- B.E.** Information Engineering (Talented student program), 2008
South China University of Technology, China

PROFESSIONAL EXPERIENCE

- Postdoctoral Fellow** Department of Biostatistics
(11/2013-present) Johns Hopkins Bloomberg School of Public Health
Advisor: Prof. Hongkai Ji

RESEARCH INTERESTS

Genomics; Data science; Single-cell genomics; Big data; Bioinformatics;
Computational biology.

My primary research interest is developing statistical and computational methods for integrative analysis of big data with focus on data science and genomics.

PUBLICATIONS

Preprints

1. **Zhou, W.**, Ji, Z. & Ji, H. Global Prediction of Chromatin Accessibility Using RNA-seq from Small Number of Cells. bioRxiv, 035816.
2. Stephens, K., **Zhou, W.**, Ji, Z., He, S., Ji, H., Guan, Y. & Taverna, S. Sex differences in gene regulation in the dorsal root ganglion after nerve injury. bioRxiv, 152652. [contribution: analyze RNA-seq data]

Refereed Journal Articles

1. Qiu, X. et al. (including **Zhou, W.**) Sequential Enhancer Sequestration Dysregulates Recombination Center Formation at the *IgH* Locus. *Molecular Cell* **70**, 21-33 (2018). [contribution: perform statistical and computational analysis of genomic data]
2. **Zhou, W.** et al. Genome-wide Prediction of DNase I Hypersensitivity Using Gene Expression. *Nature Communications* **8**, 1038 (2017).
3. Ji, Z. *, **Zhou, W.** * & Ji, H. Single-cell regulome data analysis by SCRAT. *Bioinformatics* **33**, 2930-2932 (2017). (*joint first authors)
4. **Zhou, W.**, Sherwood, B. & Ji, H. Computational Prediction of the Global Functional Genomic Landscape: Applications, Methods and Challenges. *Human Heredity* **81**, 88-105 (2016).
5. Jin, K. et al. (including **Zhou, W.**) HOXB7 Is an ER α Cofactor in the Activation of HER2 and Multiple ER Target Genes Leading to Endocrine Resistance. *Cancer Discov.* **5**, 944-959 (2015). [contribution: perform statistical and computational analysis of genomic data]
6. Wang, D., **Zhou, W.** & Yan, H. Mining of protein-protein interfacial residues from massive protein sequential and spatial data. *Fuzzy Sets Syst.* **258**, 101-116 (2015).
7. Fan, T. et al. (including **Zhou, W.**) Antagonistic effects of MYC and hypoxia in channeling glucose and glutamine into de novo nucleotide biosynthesis. *Cancer & Metabolism* **2** (Suppl 1), O10 (2014). [contribution: perform statistical and computational analysis of genomic data]
8. Wang, D. D., **Zhou, W.**, Yan, H., Wong, M. & Lee, V. Personalized prediction of EGFR mutation-induced drug resistance in lung cancer. *Scientific Reports* **3**, 2855 (2013).
9. Yang, X., **Zhou, W.**, Wang, D. D., Wu, Q. & Yan, H. A Survey on Structural Analysis of Nucleosome Core Particles. *Current Bioinformatics* **8**, 112-132 (2013).
10. **Zhou, W.**, Yan, H., Fan, X. & Hao, Q. Prediction of Protein-Protein Interactions Based on Molecular Interface Features and the Support Vector Machine. *Current Bioinformatics* **8**, 3-8 (2013).

11. Zhu, Y., **Zhou, W.**, Dai, D. & Yan, H. Identification of DNA-Binding and Protein-Binding Proteins Using Enhanced Graph Wavelet Features. *IEEE/ACM Transactions on Computational Biology and Bioinformatics (TCBB)* **10**, 1017-1031 (2013).
12. Guan, P. P., **Zhou, W.** & Yan, H. The relationship between geometric patterns of hydrogen bonds and periodic dinucleotides in nucleosome structures. *J. Theor. Biol.* **313**, 136-141 (2012).
13. **Zhou, W.** & Yan, H. Alpha shape and Delaunay triangulation in studies of protein-related interactions. *Briefings in Bioinformatics* **15**(1), 54-64 (2012).
14. **Zhou, W.**, Yan, H. & Hao, Q. Analysis of surface structures of hydrogen bonding in protein-ligand interactions using the alpha shape model. *Chemical Physics Letters* **545**, 125-131 (2012).
15. Wu, Q., **Zhou, W.**, Wang, J. & Yan, H. Correlation between the flexibility and periodic dinucleotide patterns in yeast nucleosomal DNA sequences. *J. Theor. Biol.* **284**, 92-98 (2011).
16. **Zhou, W.** & Yan, H. Prediction of DNA-binding protein based on statistical and geometric features and support vector machines. *Proteome science* **9**, 1-6 (2011).
17. **Zhou, W.** & Yan, H. Relationship between periodic dinucleotides and the nucleosome structure revealed by alpha shape modeling. *Chemical Physics Letters* **489**, 225-228 (2010).
18. **Zhou, W.** & Yan, H. A discriminatory function for prediction of protein-DNA interactions based on alpha shape modeling. *Bioinformatics* **26**, 2541-2548 (2010).

Refereed Conference Papers

1. **Zhou, W.**, Wang, D. & Yan, H. Prediction of anti-EGFR drug resistance base on binding free energy and hydrogen bond analysis. *2013 IEEE Symposium on Computational Intelligence in Bioinformatics and Computational Biology*, Singapore, April 2013.
2. **Zhou, W.** & Yan, H. Analysis of Ligand Binding Sites Using Alpha Shapes. *2012 IEEE International Conference on System, Man, and Cybernetics*, Seoul, Korea, October 2012.
3. **Zhou, W.**, Yan, H., Fan, X. & Hao, Q. Prediction of protein-protein interactions using alpha shape modeling. *2011 International Symposium on Computational Models for Life Sciences*, Toyama City, Japan, October 2011, pp 244-252.
4. **Zhou, W.** & Yan, H. Prediction of DNA-binding protein based on alpha shape modeling. *2010 IEEE International Conference on Bioinformatics and Biomedicine (BIBM)*, Hong Kong, December 2010, pp 23-28.
5. **Zhou, W.**, Xu, X. & Huang, W. Shape and Boundary Analysis for Classification of Breast Masses. *2008 International Symposium on Computational Intelligence and Design*, Wuhan,

China, October 2008, Vol. 2, pp 42-46.

6. Tang, X., Xu, X., **Zhou, W.** & Mao, Y. A Novel Semantic based CAD System for Mammography. *2008 International Conference on Computer Science and Information Technology*, Singapore, August 2008, pp 97-101.

Posters

1. **Zhou, W.**, Du, F., Bai, J., Ying M., Laterra J. & Ji, H. Computational prediction of cistrome with application to stem cells. *The 7th Annual Maryland Stem Cell Research Symposium*, Silver Spring, MD, USA, December 2014.
2. **Zhou, W.**, Ji, Z. & Ji, H., Analysis and prediction of single-cell functional genomic data. *CSHL Single Cell Analyses meeting*, Cold Spring Harbor, NY, USA, November 2017.

Conference presentations

1. **Zhou, W.** & Ji, H. Big Data Regression and Prediction in Functional Genomics. *The 2015 INFORMS Annual Meeting*, Philadelphia, PA, USA, November 2015.
2. **Zhou, W.** et al. Big data regression and prediction for high-throughput genomic data. *Joint Statistical Meetings*, Chicago, IL, July 2016.
3. **Zhou, W.**, Ji, Z. & Ji, H. Global Prediction of Chromatin Accessibility Using RNA-seq from Single Cell and Small Number of Cell. *2017 ICSA Applied Statistics Symposium*, Chicago, IL, June 2017.
4. **Zhou, W.**, Ji, Z. & Ji, H. Next Generation Analysis Tools for Single-Cell Functional Genomic Data. *Joint Statistical Meetings*, Baltimore, MD, August 2017.

Invited Talks

1. Prediction and Analysis of High Dimensional Functional Genomics Data, Institute for Basic Biomedical Sciences at Johns Hopkins School of Medicine, Baltimore, MD, December 2016.
2. Genome-wide prediction of DNase I Hypersensitivity Using Gene Expression, *11th Annual Symposium and Poster Session on Genomics and Bioinformatics*, Center for Computational Genomics at Johns Hopkins, Baltimore, MD, October 2017.
3. Prediction and analysis of functional genomic data: from bulk to single cell, The Lieber Institute for Brain Development, Baltimore, MD, April 2018.

Patent Applications

1. **Zhou, W.**, & Yan, H. METHODS FOR MODELING AND ANALYSIS OF INTERFACE BETWEEN POINT PATTERNS. US Patent Application 20130024175, publication date: January 24, 2013.

SOFTWARE AND DATABASE

1. BIRD: Big data Regression for predicting DNase I hypersensitivity.
<https://github.com/WeiqiangZhou/BIRD>
2. PDDB: Predicted DNase I Hypersensitivity Database.
<http://jilab.biostat.jhsph.edu/~bsherwo2/bird/index.php>
3. SCRAT: Single-cell regulome analysis toolbox.
<https://zhiji.shinyapps.io/scrat/>
4. SCDV: Single-cell differential variance analysis.
<https://github.com/WeiqiangZhou/SCDV>

PROFESSIONAL ACTIVITIES

Reviewer	Nature Methods, Nature Communications, Nucleic Acid Research, Biostatistics, BMC Bioinformatics, Proteome Science, IEEE Transactions on NanoBioscience, The Eleventh Asia Pacific Bioinformatics Conference 2013, The Tenth Asia Pacific Bioinformatics Conference 2012.
Session chair	Joint Statistical Meetings, 2016 Joint Statistical Meetings, 2017
Thesis examiner	Hamidreza Khataee Gavgani. 2016. Theoretical Investigation of Intracellular Transport by Molecular Motors. Ph.D. Thesis, Griffith University, Brisbane.

TEACHING ACTIVITIES

2012	Teaching assistant in EE5806: Topics in Image Processing City University of Hong Kong
2010- 2011	Teaching assistant in EE2000: Logic Circuit Design City University of Hong Kong
2010	Teaching assistant in EE3206: Java Programming & Applications City University of Hong Kong

MENTORING EXPERIENCE

- 2017 Runzhe Li, summer intern, Johns Hopkins University
Predicting chromatin accessibility using gene expression across different platforms.
current position: PhD student, Johns Hopkins University
- 2015 Steffen Cornwell, summer intern, Johns Hopkins University
Predicting tissue-specific locations of cis-regulatory elements using chromatin immunoprecipitation sequencing data.
current position: undergraduate student, University of Pennsylvania

ACADEMIC HONORS & AWARDS

- Research Tuition Scholarships** (2009-2010, 2010-2011, 2011-2012), Chow Yei Ching School of Graduate Studies, City University of Hong Kong, Hong Kong.
- Outstanding Academic Performance Award for Research Degree Students** (2009-2010, 2010-2011, 2011-2012), Chow Yei Ching School of Graduate Studies, City University of Hong Kong, Hong Kong.
- Postgraduate Studentship** (2009-2013), University Grants Committee, Hong Kong.