EDUCATION

Boston UNIVERSITY, Program of Bioinformatics, Boston, MA *Doctor of Philosophy in Bioinformatics*, May 2011

NATIONAL CHIAO TUNG UNIVERSITY, Institute of Bioinformatics, HsinChu, Taiwan *Master of Bioinformatics*, 2007

NATIONAL CENTRAL UNIVERSITY, Department of Computer Science, Jhongli City, Taiwan *Bachelor of Engineering in Computer Science*, 2005

TAINAN FIRST SENIOR HIGH SCHOOL, Tainan, Taiwan *Diploma*, 2001

Areas of Interest

- Next generation sequencing (NGS) analysis
- Small silencing RNA (miRNA, endo-siRNA, and piRNA) biogenesis
- Epigenetics (histone modification and DNA methylation) in stem cells and cancers
- Bioinformatics algorithms design and cloud/GPU computing
- Pathway enrichment analysis

RESEARCH EXPERIENCE

National Chiao Tung University, HsinChu, Taiwan

2012~present

Assistant professor, Institute of Bioinformatics and Systems Biology & Department of biological Science and Technology

- Conducted research on human Argonaute-small RNA sorting and tailing mechanisms, resulting in publication in **Molecular Cell**
- Conducted bioinformatics analysis to study mouse and human epigenetics. Results published in **Developmental Cell, Cell Report** and **PLoS Genetics**
- Developed algorithms and cloud applications for NGS analysis, published in Nuclear Acids Research

University of Massachusetts Medical School, Worcester, MA

2011~2012

Post doctoral researcher, Zhiping Weng's Lab, Program in Bioinformatics and Integrative Biology, UMass Med

- Designed algorithms to reconstruct hepatitis C virus quasi-species from NGS data
- Conducted projects pertaining small RNA tailing mechanisms, resulting in publication in <u>Cell</u> and Nature Methods
- Involved in research of mouse stem cell epigenetics, resulting in publication in Cell
- Published a review of enrichment analysis approaches on Briefings in Bioinformatics

University of Massachusetts Medical School, Worcester, MA

2009~2011

Research Assistant, Zhiping Weng's Lab, Program in Bioinformatics and Integrative Biology, UMass Med

- Devised novel suffix tree based algorithms to search for trimming and tailing in target RNA directed small RNA silencing effect.
- Designed analysis pipeline for NGS data.
- Discovered the putative model of trimming and tailing, resulting in publication in <u>Science</u>, <u>Genome</u>
 <u>Research</u> and <u>RNA</u>

Research Assistant, Charles DeLisi's Lab, Bioinformatics program, BU

• Developed network based pathway enrichment analysis algorithms and JAVA based tools. Results published on **Genome Biology** and **Nuclear Acids Research**

PUBLICATIONS

SCI journal:

- Chou M.-T., Han B. W., Hsiao C.-P., Zamore P. D., Weng Z. and **Hung J.-H.** *Tailor: A Computational Framework for Detecting Non-Templated Tailing of Small Silencing RNAs*. **Nucleic Acids Research**. Volume: 10. 1093. 2015.
- Reimão-Pinto M. M., Ignatova V., Burkard T. R., **Hung J.-H.**, Manzenreither R. A., Sowemimo I., Herzog V. A., Reichholf B., Fariña-Lopez S., Ameres S. L.. *Uridylation of RNA Hairpins by Tailor Confines the Emergence of MicroRNAs in Drosophila*. **Molecular Cell**. Volume 59, Issue 2, Pages 203–216, 2015.
- Tung Y.-T., Lu Y.-L., Peng K.-C., Yen Y.-P., Chang M., Li J., H. Jung, Thams S., Y.-P. Huang, **Hung J.-H.**, and Chen J.-A. *Mir-17~92 Governs Motor Neuron Subtype Survival by Mediating Nuclear PTEN*. **Cell Reports**. Volume: 11 Pages:1–14. 2015
- Li, Y.-L., Weng J.-C., Hsiao C.-C., Chou M.-T., Tseng C.-W. and **Hung J.-H**. *PEAT: an intelligent and efficient paired-end sequencing adapter trimming algorithm*. **BMC Bioinformatics**, 16:S2, 2015
- Carone RB., **Hung J-H**, Sarah J. Hainer, Min-Te Chou, Dawn M. Carone, Zhiping Weng, Thomas G. Fazzio, Oliver J. Rando. *High-Resolution Mapping of Chromatin Packaging in Mouse Embryonic Stem Cells and Sperm*. **Developmental Cell** Vol: 30(1), 2014
- Yildirim O., **Hung J-H**, Ryan J. Cedeno, Zhiping Weng, Christopher J. Lengner, Oliver J. Rando. *A System for Genome-Wide Histone Variant Dynamics In ES Cells Reveals Dynamic MacroH2A2 Replacement at Promoters*. **PLoS Genetics**, 10.1371, Aug. 2014
- Chen PB, **Hung J-H**, Hickman TL, Coles AH, Carey JF, Weng Z, Chu F, Fazzio TG. *Hdac6 regulates Tip60-p400 function in stem cells*. **eLIFE**, December 3, 2013
- Chih-Hung Chou, Feng-Mao Lin, Min-Te Chou, Sheng-Da Hsu, Tzu-Hao Chang, Shun-Long Weng, Sirjana Shrestha, Chiung-Chih Hsiao, **Hung J-H** and Hsien-Da Huang. *A computational approach for identifying microRNA-target interactions using high-throughput CLIP and PAR-CLIP sequencing*. **BMC Genomics**, 14:S2, 2013
- Jie Wang, Jiali Zhuang, Sowmya Iyer, Xin-Ying Lin, Melissa C. Greven, Bong-Hyun Kim, Jill Moor, Brian G. Pierce, Xianjun Dong, Daniel Virgil, Ewan Birney, **Hung J-H** and Zhiping Weng. Factorbook.org: a Wiki-based database for transcription factor-binding data generated by the ENCODE consortium. **Nucleic Acids Research**. 41(D1):D171-D176, 2013
- Ryuya Fukunaga, Bo W. Han, **Hung J-H**, Jia Xu, Zhiping Weng, Phillip D. Zamore. *Dicer Partner Proteins Tune the Length of Mature miRNAs in Flies and Mammals*. **Cell**. 2012 Oct 26;151(3):533-46
- Xie J, Ameres SL, Friedline R, **Hung J-H**, Zhang Y, Xie Q, Zhong L, Su Q, He R, Li M, Li H, Mu X, Zhang H, Broderick JA, Kim JK, Weng Z, Flotte TR, Zamore PD, Gao G. *Long-term, efficient inhibition of microRNA function in mice using rAAV vectors*. **Nature Methods**, 2012 Mar 4;9(4):403-9

- Shigemizu D , Hu Z , **Hung J-H** , Huang C-L , Wang Y , et al. *Using Functional Signatures to Identify Repositioned Drugs for Breast, Myelogenous Leukemia and Prostate Cancer*. **PLoS Comput Biol** 8(2): e1002347. doi:10.1371/journal.pcbi.1002347
- Yildirim O, Li R, **Hung J-H**, Chen PB, Dong X, Ee L-Y, Weng Z, Rando OJ and Fazzio TG. *Mbd3/NURD complex regulates expression of 5-hydroxymethylcytosine marked genes in embryonic stem cells*. **Cell**. Volume 147, Issue 7, 1498-1510, 23 December 2011
- Han B, **Hung J-H**, Weng Z, Zamore PD, Ameres SL. *The 3'-to-5' Exoribonuclease Nibbler Shapes the 3' Ends of MicroRNAs Bound to Drosophila Argonaute1*. **Current Biology**, November 2011.
- **Hung J-H**, Yang TH, Hu Z, Weng Z, Delisi C. *Gene set enrichment analysis: performance evaluation and usage guidelines.* **Brief Bioinform**. 2011 Sep 7.
- Yang T-H, Kon M, **Hung J-H**, DeLisi C. Combinations of newly confirmed Glioma-Associated loci link regions on chromosomes 1 and 9 to increased disease risk. **BMC Medical Genomics** 2011, 4:63
- Berezikov E, Robine N, Samsonova A, Westholm JO, Naqvi A, **Hung J-H**, Okamura K, Dai Q, Bortolamiol-Becet D, Martin R, Zhao Y, Zamore PD, Hannon GJ, Marra MA, Weng Z, Perrimon N, Lai EC. *Deep annotation of Drosophila melanogaster micro RNAs yields insights into their processing, modification, and emergence.* **Genome Res.**, 2011; Feb; 21(2):203-15
- Ameres SL, **Hung J-H**, Xu J, Weng Z, Zamore PD. Target RNA-directed Tailing and Trimming purifies the sorting of endo-siRNA between the two Drosophila argonaute proteins. **RNA**. 2011; 17(1):54-63.
- Xie J, Xie Q, Zhang H, Ameres SL, **Hung J-H**, Su Q, He R, Mu X, Seher Ahmed S, Park S, Kato H, Li C, Mueller C, Mello CC, Weng Z, Flotte TR, Zamore PD, Gao G. *MicroRNA-regulated, Systemically Delivered rAAV9: A Step Closer to CNS-restricted Transgene Expression*. **Mol Ther** (PMID: 21179009)
- Ameres SL, Horwich MD, **Hung J-H,** Xu J, Ghildiyal M, Weng Z, Zamore PD. *Target RNA-directed trimming and tailing of small silencing RNAs.* **Science**. 320:1534-1539, 2010
- **Hung J-H,** Whitfield TW, Yang TH, Hu Z, Weng Z, DeLisi C. *Identification of functional modules that correlate with phenotypic difference: the influence of network topology*. **Genome Biology**.11:R23, 2010
- Hwang H, Vreven T, Pierce B, **Hung J-H**, Weng Z. *Performance of ZDOCK and ZRANK in CAPRI round 13-19*. **Protein**. 78(15):3104-10
- Hu Z., **Hung J-H,** Chang Y-C, Huang C-L, Huyck M. and DeLisi C. VisANT 3.5: multi-scale network visualization, analysis and inference based on the gene ontology. **Nucleic Acids Research**. 37: W115-W121, 2009
- Hung J-H, Hung J-H, and Chen S-G. A systematic Optimized Comparison Algorithm for Fast LDPC Decoding. ICACT 10th. Vol. 3, pp. 1513-1526. 2008.
- **Hung J-H,** Huang H-D, and Lee T-Y. *ProKware: integrated software for presenting protein structural properties in protein tertiary structures.* **Nucleic Acids Research**. 34: W89-W94, 2006.
- Lee T-Y, Huang H-D, **Hung J-H**, Yang Y-S, and Wang T-H. *dbPTM: An Information Repository of Protein Post-Translational Modification*. **Nucleic Acids Research**. 34: D622-D627. 2006.

Book Chapter:

- **Hung J-H**, and Weng Z. *Bioinformatics Essentials*. Molecular Cloning: A Laboratory Manual, 4th ed. Cold Spring Harbor Laboratory, 2012, ISBN-10: 1936113422.
- **Hung J-H**. *Enrichment Analysis*. Data Mining for Systems Biology. Methods in Molecular Biology series. Springer, 2013, ISBN 978-1-62703-107-3.

International Conference:

- **Hung J-H,** Hung J-H, and Chen S-G. A General Comparator Structure for LDPC Decoder by Novel Systematic Optimization Algorithms. ICASSP2007
- **Hung J-H,** Wang W-G, and Huang H-D. Systematic Identification of RNA Editing Site in Human Genome. ICS2006

International Patent:

- Hung J-H, **Hung J-H**, and Chen S-G. *An Efficient Set-Cyclic Comparison IP for Low Density Parity Check Code*. USA. (Patent No.: US 7945839B2)
- **Hung J-H,** Hung J-H, and Chen S-G. *An Efficient Cyclic Comparison IP for Low Density Parity Check Codes.* USA. (Patent No.: US 7966543B2)