Shao-shan Carol Huang Curriculum Vitae

September 2021

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Education and qualifications

2018 Workshop on Leadership in Biosciences

Cold Spring Harbor Laboratory, Cold Spring Harbor, NY, USA.

2011 Ph.D., Computational and Systems Biology

Massachusetts Institute of Technology, Cambridge, MA, USA

Advisor: Dr. Ernest Fraenkel

Thesis: A constraint optimization framework for discovery of cellular signaling and regulatory

networks

2005 B.Sc., Combined Honors Computer Science and Biology, Co-op Option

University of British Columbia, Vancouver, BC, Canada

Advisor: Dr. Wyeth Wasserman

Honors thesis: Computational identification of over-represented combinations of transcription factor

binding sites in sets of co-expressed genes

Current position

2020-present Mentoring Faculty, PhD progra	am in Developmental	Genetics, NY	'U School of Medicine
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2019-present Affiliate Faculty, Center for Data Science, New York University

2018-present Assistant Professor, Center for Genomics and Systems Biology, Department of Biology, New

York University

Past research positions

2011-2017	Postdoctoral associate, Genomic Analysis Laboratory & Plant Biology Laboratory
	The Salk Institute for Biological Studies
2011	Postdoctoral associate, Department of Biological Engineering
	Massachusetts Institute of Technology
2005-2011	Graduate student, Program in Computational and Systems Biology
	Massachusetts Institute of Technology
2004-2005	Undergraduate research assistant, Center for Molecular Medicine and Therapeutics
	University of British Columbia
2002 2005	Undergraduate research assistant Department of Mathematics

2003-2005 Undergraduate research assistant, Department of Mathematics

University of British Columbia

Funding

Ongoing Grant Support

NIH - 1R35GM138143 Huang (PI) 2020/07/01 - 2025/04/30

National Institutes of Health, National Institute of General Medical Sciences

Title: Dissecting natural variation in transcription factor - DNA interactions

Description: This grant aims to study the mechanisms of genome-wide transcription factor binding site changes and the resulting regulatory networks using diverse genomes and epigenomes of natural strains of Arabidopsis as model. We will build computational models of DNA sequence and non-sequence features for predicting binding variations, and experimentally validate the biological functions of selected binding site variants.

NSF - IOS-1916804 Huang (Co-PI) 2019/08/01 - 2023/07/31

National Science Foundation, Integrative Organismal Systems

Title: TRTech-PGR: Mapping and functional characterization of cis-regulatory module variation in plants **Investigators:** Andrea Gallavotti (Rutgers University; PI), Naden Krogan (American University; Co-PI) **Description:** This grant develops methods to map transcription factor-DNA interactions that enable comparative analysis of regulatory sequences in multiple genetic backgrounds. The methods will be applied in two species with different genomic properties: maize, a major monocot crop with a large genome, and Arabidopsis, a model eudicot with a compact genome.

Publications (*Equal contribution; \$NYU affiliation)

Preprints

Orduña, L., M. Li, D. Navarro-Payá, C. Zhang, A. Santiago, P. Romero, Ž. Ramšak, G. Magon, J. Höll, P. Merz, K. Gruden, A. Vannozzi, D. Cantu, J. Bogs, D. C. J. Wong, S.-s. C. Huang^{\$\\$}, and J. T. Matus (2021). Major orchestration of shikimate, early phenylpropanoid and stilbenoid pathways by Subgroup 2 R2R3-MYBs in grapevine. bioRxiv. eprint: https://www.biorxiv.org/content/early/2021/08/23/2020.12.31.424746.full.pdf.

Refereed research papers

- Plant Cell Atlas Consortium, S. Ghosh Jha*, A. T. Borowsky*, B. J. Cole*, N. Fahlgren*, A. Farmer*, S.-s. C. Huang*\$, P. Karia*, M. Libault*, N. J. Provart*, S. L. Rice, M. Saura-Sanchez*, P. Agarwal, A. H. Ahkami, C. R. Anderton, S. P. Briggs, J. A. Brophy, P. Denolf, L. F. Di Costanzo, M. Exposito-Alonso, S. Giacomello, F. Gomez-Cano, K. Kaufmann, D. K. Ko, S. Kumar, A. V. Malkovskiy, N. Nakayama, T. Obata, M. S. Otegui, G. Palfalvi, E. H. Quezada-Rodríguez, R. Singh, R. G. Uhrig, J. Waese, K. Van Wijk, C. R. Wright, D. W. Ehrhardt, K. D. Birnbaum, and S. Y. Rhee (2021). Science Forum: Vision, challenges and opportunities for a Plant Cell Atlas. *eLife* 10, e66877.
- 2. Scherma, M., J. S. Qvist, A. Asok, **S.-s. C. Huang**^{\$}, P. Masia, M. Deidda, Y. B. Wei, R. K. Soni, W. Fratta, P. Fadda, E. R. Kandel, D. B. Kandel, and P. A. Melas (2020). Cannabinoid exposure in rat adolescence reprograms the initial behavioral, molecular, and epigenetic response to cocaine. *Proceedings of the National Academy of Sciences* **117**(18), 9991–10002.
- 3. Procko, C., S. Morrison, C. Dunar, S. Mills, B. Maldonado, C. Cockrum, N. E. Peters, **S.-s. C. Huang**^{\$}, and J. Chory (2019). Big Data to the Bench: Transcriptome Analysis for Undergraduates. *CBE-Life Sciences Education* **18**(2), ar19.
- 4. Ursu, O., S. J. C. Gosline, N. Beeharry, L. Fink, V. Bhattacharjee, **S.-s. C. Huang**, Y. Zhou, T. Yen, and E. Fraenkel (Oct. 2017). Network modeling of kinase inhibitor polypharmacology reveals pathways targeted in chemical screens. *PLOS ONE* **12**(10), 1–22.
- 5. Trigg, S. A., R. M. Garza, A. MacWilliams, J. R. Nery, A. Bartlett, R. Castanon, A. Goubil, J. Feeney, R. O'Malley, S.-s. C. Huang, Z. Zhang, M. Galli, and J. R. Ecker (2017). CrY2H-seq: a massively multiplexed assay for deep-coverage interactome mapping. *Nature Methods* **14**(8), 819.
- 6. Bartlett, A., R. C. O'Malley, **S.-s. C. Huang**, M. Galli, J. R. Nery, A. Gallavotti, and J. R. Ecker (2017). Mapping genome-wide transcription-factor binding sites using DAP-seq. *Nature Protocols* **12**(8), 1659–1672.
- 7. Song, L., **S.-s. C. Huang**, A. Wise, R. Castanon, J. R. Nery, H. Chen, M. Watanabe, J. Thomas, Z. Bar-Joseph, and J. R. Ecker (2016). A transcription factor hierarchy defines an environmental stress response network. *Science* **354**(6312).
- 8. Kawakatsu, T. *, **S.-s. C. Huang***, F. Jupe*, E. Sasaki*, R. J. Schmitz, M. A. Urich, R. Castanon, J. R. Nery, C. Barragan, Y. He, H. Chen, M. Dubin, C.-R. Lee, C. Wang, F. Bemm, C. Becker, R. O'Neil, R. C. O'Malley, D. X. Quarless, The 1001 Genomes Consortium, N. J. Schork, D. Weigel, M. Nordborg, and J. R. Ecker (2016). Epigenomic Diversity in a Global Collection of *Arabidopsis thaliana* Accessions. *Cell* **166**(2), 492–505.
- 9. O'Malley, R. C. *, **S.-s. C. Huang***, L. Song, M. G. Lewsey, A. Bartlett, M. Galli, J. R. Nery, A. Gallavotti, and J. R. Ecker (2016). Cistrome and Epicistrome Features Shape the Regulatory DNA Landscape. *Cell* **165**(5), 1280–1292.

- 10. Pedmale, U. V., **S.-s. C. Huang**, M. Zander, B. J. Cole, J. Hetzel, K. Ljung, P. A. Reis, P. Sridevi, K. Nito, J. R. Nery, J. R. Ecker, and J. Chory (2016). Cryptochromes Interact Directly with PIFs to Control Plant Growth in Limiting Blue Light. *Cell* **164**(1-2), 233–245.
- 11. Chang, K. N., S. Zhong, M. T. Weirauch, G. Hon, M. Pelizzola, H. Li, **S.-s. C. Huang**, R. J. Schmitz, M. A. Urich, D. Kuo, et al. (2013). Temporal transcriptional response to ethylene gas drives growth hormone cross-regulation in Arabidopsis. *eLife* **2**, e00675.
- 12. **Huang**, **S.-s. C.**, D. C. Clarke, S. J. Gosline, A. Labadorf, C. R. Chouinard, W. Gordon, D. A. Lauffenburger, and E. Fraenkel (2013). Linking proteomic and transcriptional data through the interactome and epigenome reveals a map of oncogene-induced signaling. *PLoS Computational Biology* **9**(2), e1002887.
- 13. Tuncbag, N., A. Braunstein, A. Pagnani, **S.-S. C. Huang**, J. Chayes, C. Borgs, R. Zecchina, and E. Fraenkel (2013). Simultaneous reconstruction of multiple signaling pathways via the Prize-Collecting Steiner Forest problem. *Journal of Computational Biology* **20**(2), 124–136.
- 14. Qiao, H., Z. Shen, **S.-s. C. Huang**, R. J. Schmitz, M. A. Urich, S. P. Briggs, and J. R. Ecker (2012). Processing and subcellular trafficking of ER-tethered EIN2 control response to ethylene gas. *Science* **338**(6105), 390–3.
- 15. Tuncbag, N., S. McCallum, **S.-s. C. Huang**, and E. Fraenkel (2012). SteinerNet: a web server for integrating 'omic' data to discover hidden components of response pathways. *Nucleic Acids Research* **40**(W1), W505–W509.
- 16. **Huang**, **S.-s. C.** and E. Fraenkel (2009). Integrating proteomic, transcriptional, and interactome data reveals hidden components of signaling and regulatory networks. *Science Signaling* **2**(81), ra40.

Reviews

- 1. Shojaee, A., M. Saavedra, and **S.-s. C. Huang**^{\$} (2021). Potentials of single-cell genomics in deciphering cellular phenotypes. *Current Opinion in Plant Biology* **63**, 102059.
- 2. Qian, Y. and **S.-s. C. Huang**^{\$} (2020). Improving plant gene regulatory network inference by integrative analysis of multi-omics and high resolution data sets. *Current Opinion in Systems Biology* **22**, 8–15.
- 3. **Huang**, **S.-s. C.** and J. R. Ecker (2018). Piecing together cis-regulatory networks: insights from epigenomics studies in plants. *WIREs Systems Biology and Medicine* **10**(3), e1411.

Book chapter

1. **Huang**, **S.** and E. Fraenkel (2012). Swimming upstream: identifying proteomic signals that drive transcriptional changes using the interactome and multiple '-omics' datasets. *Computational Methods in Cell Biology* **110**, 57.

Patent

1. Fraenkel, E., **S.-S. C. Huang**, D. R. Karger, L. Riva, and E. Yeger-Lotem (Dec. 2012). *Identifying biological response pathways*. US Patent No. 8612160. Issued December 17, 2013.

Invited talks (\$NYU affiliation)

- 1. \$ Annual Meeting of Mid-Atlantic section of the American Society of Plant Biologists (ASPB). College Park, MD. (May 2020). Cancelled due to COVID-19.
- 2. \$ University of Cambridge Sainsbury Laboratory. Cambridge, UK. (November 27, 2019).
- 3. \$ University of Maryland Department of Cell Biology and Molecular Genetics. College Park, MD. (September 20, 2019).
- 4. \$ Iowa State University Bioinformatics and Computational Biology Symposium. Ames, IA. (April 29, 2019).
- 5. SCNRS Jacques Monod Conference: First International Plant Systems Biology Meeting. Roscoff, France. (September 12, 2018).
- 6. \$ Workshop on Construction of Gene Regulatory Networks in Arabidopsis in the 29th International Conference on Arabidopsis Research. Turku, Finland. (June 26, 2018).
- 7. \$ Boyce Thompson Institute. Ithaca, NY. (May 8, 2018).
- 8. University of California, Riverside Data Science Center. Riverside, CA. (March 22, 2017).
- 9. Cornell University School of Integrative Plant Science. Ithaca, NY. (March 9, 2017).

- 10. New York University Department of Biology. New York, NY. (March 7, 2017).
- 11. Washington University at St. Louis Department of Biology. St. Louis, MO. (January 9, 2017).
- 12. Academia Sinica Institute of Cellular and Organismic Biology. Taipei, Taiwan. (December 21, 2016).
- 13. Carnegie Institution of Science Department of Plant Biology. Stanford, CA. (December 2, 2016).
- 14. Salk Featured Fellow Series. The Salk Institute for Biological Studies. La Jolla, CA. (August 10, 2016).
- 15. 22nd International Conference on Plant Growth Substances. Toronto, Canada. (June 22, 2016).
- 16. UCLA Department of Molecular, Cell, and Developmental Biology and the Institute for Quantitative and Computational Biosciences. Los Angeles, CA. (April 25, 2016).
- 17. Cold Spring Harbor Conference on Systems Biology: Global Regulation of Gene Expression. Cold Spring Harbor, NY. (March 19, 2016).
- 18. University of Toronto Department of Cell and Systems Biology. Toronto, Canada. (March 14, 2016).
- 19. University of New Hampshire Department of Molecular, Cellular, and Biomedical Sciences. Durham, NH. (July 15, 2015).
- 20. San Diego Center for Systems Biology: Cytoscape and Network Analysis Workshop. San Diego, CA. (April 17, 2015).
- 21. Oregon State University Center for Genome Research and Biocomputing, Corvalis, OR. (April 15, 2015).

Students and postdoctoral scholars mentored

Postdoctoral scholars

2020-present Abbas Shojaee 2018-present Miaomiao Li 2020 Yichun Qian

Graduate students

2021-present	Will Hinckley, Ph.D. in Biology, NYU. (joint with Gloria Coruzzi)
2021-present	Wanru Lin, M.S. in Biology, NYU.
2020-2021	Michelle Saavedra, M.S. in Biology, NYU.
2020-2021	Nabilla Zaman, M.S. in Biology, NYU.
2018-2019	Ziyan Lin, M.S. in Biology, NYU.
2018-2019	Ruxin Dai, M.S. in Biology, NYU.
2018-2019	Diogo Mesquita, M.S. in Data Science, NYU.
2018-2019	Yuhan Wang, M.S. in Biology, NYU.

High school intern

2021-present Hannah Will, The Packer Collegiate Institute, Brooklyn, NY.

Teaching

2018-present	Module instructor, Developmental and Stem Cell Systems II (Course DSCSII), NYU School of
	Medicine.
2018-present	Co-instructor, Bio Core III: Molecules and Cells (Course BIOL-GA 2003), NYU Department of
	Biology.
2020-2021	Instructor, Fundamentals of Bioinformatics (Course BIOL-UA 124), NYU Department of Biology.
2020	Co-instructor, Plant Science and Biotechnology (Course BIOL-UA 560), NYU Department of
	Biology.
2019	Co-instructor, Fundamentals of Bioinformatics (Course BIOL-UA 124), NYU Department of
	Biology.

Professional activities

➤ Computational/Modeling/Al Committee Lead, The Plant Cell Atlas Research Coordination Network, 2021.

- ➤ Reviewer for National Sciences and Engineering Research Council of Canada Discovery Grant, 2021.
- ➤ Reviewer for BMC Biology, BMC Bioinformatics, Current Opinion in Plant Biology, eLife, Genome Research, IEEE Transactions on Biomedical Engineering, iScience, Molecular Cell, Nature, Nature Communications, Nature Plants, npj Systems Biology and Applications, Nucleic Acids Research, Plant J, The Plant Cell, PLoS ONE, PNAS.
- ➤ Member, International Society for Computational Biology.
- ➤ Member, American Society of Plant Biologists.

Committees

NYU Biology Department

- ➤ Organizer, Biology Faculty Exchange Seminar Series, AY 2021.
- ➤ Co-chair, PhD Admissions Committee, AY 2021.
- ➤ Member, PhD Admissions Committee, AY 2018-2020.
- ➤ Organizer, Biology Postdoc Seminar Series, AY 2019-2020.
- ➤ PhD Student Thesis Committees
 - Yingzhen Pei, committee chair, 2021-present.
 - Ornob Alam, committee member, 2021-present.
 - Jose Antonio Salome correa, committee member, 2020-present.
 - Xinyi (Cathy) Guo, committee member, 2020-present.
 - Xinhe Xue, committee member, 2020-present.
 - Yuhan Hao, committee member, 2020-present.
 - Cassandra Buzby, committee member, 2020-present.
 - Akash Sookdeo, committee member, 2019-present.
 - Victoria Le, committee member, 2019-present.
 - Chang Wang, committee member, 2019-2021.

Honors and awards

2014	Pioneer Fund Fellow, Salk Institute for Biological Studies
2007-2011	Graduate Fellowship, NCI Center for Cancer Systems Biology at MIT
2011	Scholar-in-Training Award, AACR-NCI Conference on Systems Biology
2005-2009	Post-graduate Scholarship at Foreign Universities, Natural Sciences and Engineering Research
	Council of Canada (NSERC)
2005	Amy E. Sauder Scholarship and Jean Craig Smith Scholarship, UBC
2004, 2005	Undergraduate Student Research Award in Industry and University, NSERC
2004	Wesbrook Scholar, UBC
2004	AstraZeneca Undergraduate Scholarship, UBC
2002-2003	Women in Engineering and Science Program, National Research Council of Canada
2000-2004	Chancellor's Entrance Scholarship, UBC
2000-2004	IBM Canada Pacific Development Center Scholarship