CHUN SU

Bioinformatics Scientist | Computational Biologist

I have 10+ years of hands-on experience in bioinformatics. Skilled in both molecular cloning and computational analysis of next generation sequencing (NGS) data. Proficient in a variety of programming languages, HPC systems, cutting-edge bioinformatic software, and multivariate statistical analysis. Knowledgeable about molecular cell biology and immunology. Proven experience in troubleshooting and meeting challenging deadlines in a fast-paced work environment. Excellent professional communication and leadership skills developed through community service and extensive teaching experiences.



EDUCATION

2017 2011

Ph.D, Biology

University of Virginia

Charlottesville, VA

• Thesis: Uncovering the molecular mechanism underlying the virulence of Striga aesnerioides



B.S Biology

Huazhong Agricultural University

Wuhan, China



PROFESSIONAL EXPERIENCE

Scientist, Computational Biology

Current 2022

2022

2018

Amgen, Inc

Remote

Bioinformatics Scientist

Children's Hospital of Philadelphia

Philadelphia, PA

- Established in-house bioinformatics analysis pipelines and SOP for the various types of NGS data including RNA-seq, ChIP-seq, ATAC-seq, Capture-C and Hi-C.
- Predicted candidate gene targets for drug repurposing by integrating human genetics (eg. GWAS, rare variants) with functional genomics data in immune, neurological, and metabolic contexts
- Innovated computational workflows to analyze data from high-throughput screening, including CRISPR screen, MPRA (Massive Parallel Reporter Assay) and HiDRA (High-resolution Dissection of Regulatory Activity).
- Designed and maintained Laboratory Information Management System (LIMS) using relationship database application FileMaker
- Developed and maintained web applications using R Shiny.
- Authored 14 scientific publications of which 4 are first-authored and 10 are peer-
- Presented research at national and international scientific conferences
- Recruited and trained bioinformatics fellows and graduate students.

CONTACT INFO

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- learniningwithsckinta
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- G chun-su

SKILL **HIGHLIGHTS**

- NGS analysis
- Single cell genomics
- Programming with R, Perl, Python, Shell and Matlab
- Database management with SQL and FileMaker
- Web application development with R shiny
- HPC system with PBS, SLURM and SGE
- Version control with Git and GitHub

Last updated on 2022-08-31.

Prepared by {Pagedown}

Graduate Researcher 2017 Charlottesville, VA University of Virginia (Timko Lab) 2013 • Initiated and led a research project characterizing the genetic basis for virulence variance of a major African crop pest • Implemented large-scale data mining and developed custom bioinformatic scripts to identify expression pattern differences under different biological conditions. · Collaborated with external research groups on comparative functional genomics of parasitic plants • Designed and performed experiments testing putative parasitism genes • Authored scientific publications and assisted in writing government-funded grants • Mentored and trained undergraduate students **Graduate Researcher** 2013 University of Virginia (Li Lab) Charlottesville, VA 2011 • Standardized computational pipeline to predict microRNA on multiple crop species using small RNA sequencing analysis on HPC clusters · Modified and optimized microRNA prediction Perl package for mining Expression Short Tags (ESTs) data • Identified novel microRNAs using Northern Blotting and qPCR 2010 **Summer Intern** Beijing Genomics Institute (BGI) Shenzhen, China 2010 • Designed custom code to parse small RNA sequencing and degradome data to predict small RNA and their targets in Arabidopsis · Assisted with alignment algorithm development for Short Oligonucleotide Analysis Package (SOAP) I LEADERSHIP EXPERIENCE Co-organizer Current R ladies Philly Philadelphia, PA 2019 • Organized monthly meetup https://www.meetup.com/rladies philly/ Authored blogs to summarize events. • Designed an outreach brochure and developed the mentorship/speaker application shiny dashboard https://rladiesphilly .shinyapps.io/mentorship/ • Organized and led data hackathon projects to help local non-profit organizations **Teaching Assistant** 2017 University of Virginia, Department of Biology Charlottesville, VA 2012 • Taught Introduction to Biology Lab (Head TA), Genetics Lecture, Plant Pathology Lab · Explained challenging biological concepts and demonstrated laboratory techniques to undergraduate students. · Led and coordinated graduate teaching assistants to maximize learning opportunities for students. T AWARDS AND HONORS **Poster Winner** 2020 Penn joint epigenetics and immunology symposium Philadelphia, PA **Reviewers' Choice Abstract** 2019 American Society of Human Genetics (ASHG) OHouston, Texa **Best Poster Award** 2015 The 13th World Congress on Parasitic Plants Kunming, China

2015		Graduate Student and Postdoc Association Travel Award University of Virginia, Department of Biology	• Charlottesville, VA	
2011		First Year Fellowship University of Virginia, Department of Biology	• Charlottesville, VA	
2011		Outstanding Undergraduate Thesis Award Huazhong Agricultural University	• Wuhan, China	
2011 2010		Monsanto Fellowship Monsanto Inc & Huazhong Agricultural University	🗪 Wuhan, China	
2011 2010	•	National Fund for Fostering Talents of Basic Science National Natural Science Foundation of China	• Wuhan, China	
		POSTERS AND PRESENTATIONS		
2020		promoter architecture reorganization during iPSC-derived neuronal cell differentiation plicates target genes for neurodevelopment disorders (Poster) perican Society of Human Genetics (ASHG)		
2020		Immune cell promoter connectomes suggest mechanistic bases of genetic susceptautoimmune diseases (Poster and Presentation) Penn joint epigenetics and immunology symposium	tibility to Philadelphia, PA	
2019		Network analysis workshop (Presentation) R ladies Philly	Philadelphia, PA	
2019		3D promoter-open chromatin connectomes in human immune cells yield insight into cell-specific gene regulatory architectures and the genetic basis of multiple autoimmune diseases (Poster) American Society of Human Genetics (ASHG)		
2019		Whole Genome Sequencing (WGS) of saliva and osteochondroma-derived DNA from Multiple Exostoses (HME) families reveals new insights into pathogenesis (Poster American Society for Bone and Mineral Research (ASBMR)	HME) families reveals new insights into pathogenesis (Poster)	
2015		Identification of Parasite Effectors Mediating Striga gesnerioides-Host Plant Intelligence 13th World Congress on Parasitic Plants	ractions (Poster) • Kunming, China	
2015	•	Comparison of transcriptome profiling in different races of Striga gesnerioides during its interaction with Vigna unguiculata L. Walp (Presentation)		
		Annual Conference for Parasite Plant Genomics Project	State College, PA	
		SELECTED PUBLICATIONS		
2022		Implicating effector genes at COVID-19 GWAS loci using promoter-focused Capture-C in disease-relevant immune cell types Genome Biology Matthew C Pahl, Carole Le Coz, Chun Su, Prabhat Sharma, Rajan M Thomas, James A Pippin, Emylette Cruz Cabrera, Matthew E Johnson, Michelle E Leonard, Sumei Lu, Alessandra Chesi, Kathleen E Sullivan, Neil Romberg, Struan FA Grant,		
		Andrew D Wells		

The three-dimensional chromatin structure of the major human pancreatic cell types reveals lineage-specific regulatory architecture of T2D risk bioRxiv

• Su Chun, Long Gao, Catherine L. May, James A. Pippin, Keith Boehm, Michelle Lee, Chengyang Liu, Matthew C. Pahl, Maria L. Golson, Ali Naji, the HPAP Consortium, Struan F.A. Grant, Andrew D. Wells, Klaus H. Kaestner.

Restriction enzyme selection dictates detection range sensitivity in chromatin conformation capture-based variant-to-gene mapping approaches

Human Genetics

• Su, C., Pahl, M.C., Grant, S.F.A. Wells, A.D

Projecting genetic associations through gene expression patterns highlights disease etiology and drug mechanisms

bioRxiv

2021

2014

• Milton Pividori, Sumei Lu, Binglan Li, **Chun Su**, Matthew E. Johnson, Wei-Qi Wei, Qiping Feng, Bahram Namjou, Krzysztof Kiryluk, Iftikhar Kullo, Yuan Luo, Blair D. Sullivan, Benjamin F. Voight, Carsten Skarke, Marylyn D. Ritchie, Struan F.A. Grant, Casey S. Greene

2021 Constrained chromatin accessibility in PU.1-mutated agammaglobulinemia patients

Journal of Experimental Medicine

• Le Coz C, Nguyen DN, **Su C**, Nolan BE, Albrecht AV, Xhani S, Sun D, Demaree B, Pillarisetti P, Khanna C, Wright F, Chen PA, Yoon S, Stiegler AL, Maurer K, Garifallou JP, Rymaszewski A, Kroft SH, Olson TS, Seif AE, Wertheim G, Grant SFA, Vo LT, Puck JM, Sullivan KE, Routes JM, Zakharova V, Shcherbina A, Mukhina A, Rudy NL, Hurst ACE, Atkinson TP, Boggon TJ, Hakonarson H, Abate AR, Hajjar J, Nicholas SK, Lupski JR, Verbsky J, Chinn IK, Gonzalez MV, Wells AD, Marson A, Poon GMK, Romberg N.

3D promoter architecture reorganization during iPSC-derived neuronal cell differentiation implicates target genes for neurodevelopmental disorders

Progress in Neurobiology

• Su, C., Argenziano, M., Lu, S., Pippin, J. A., Pahl, M. C., Leonard, M. E., Cousminer, D. L., Johnson, M. E., Lasconi, C., Wells, A. D., Chesi, A., & Grant, S. F.

Mapping effector genes at lupus GWAS loci using promoter Capture-C in follicular helper T cells

Nature Communication

• Chun Su, Matthew E. Johnson, Annabel Torres, Rajan M. Thomas, Elisabetta Manduchi, Prabhat Sharma, Carole Le Coz, Michelle E. Leonard, Sumei Lu, Kenyaita M. Hodge, Alessandra Chesi, James Pippin, Neil Romberg, Struan F. A. Grant and Andrew D. Wells

SHR4z, a novel decoy effector from the haustorium of the parasitic weed Striga gesnerioides, suppresses host plant immunity

New Phytologist

• Chun Su, Hai Liu, Eric K. Wafula, Loren Honaas, Claude W. dePamphilis, Michael P. Timko

 Identification and characterization of a subset of microRNAs in wheat Genomics

• Chun Su, Xiaozeng Yang, Shiqing Gao, Yimiao Tang, Changping Zhao, Lei Li.