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Current position

Postdoctoral Scholar Department of Psychiatry University of California, Los Angeles

Education

2022	Ph.D. Integrative Genetics and Genomics University of California Davis, Davis, CA, USA <i>Mentor: Megan Y. Dennis</i>
2015	M.S. Chemical Engineering Pontificia Universidad Católica de Chile, Santiago, Chile <i>Mentor: Loreto M. Valenzuela</i>
2012	B.S. Biological Engineering Pontificia Universidad Católica de Chile, Santiago, Chile Summa cum laude

Employment

2023 - present	Postdoctoral Scholar - University of California Los Angeles, Los Angeles, CA, USA Mentor: Jonathan Flint
2023	Postdoctoral Scholar - University of California Davis, Davis, CA, USA Mentor: Megan Y. Dennis
2018 - 2022	Graduate Student Researcher - University of California Davis, Davis, CA, USA. Mentor: Megan Y. Dennis
2015 - 2017	Bioinformatics Analyst - Pontificia Universidad Católica de Chile, Santiago, Chile <i>Mentor: Rodrigo A. Gutiérrez</i>
2012 - 2014	Research Assistant - Pontificia Universidad Católica de Chile, Santiago, Chile Mentor: Loreto M. Valenzuela
2013	Part-Time R&D - Biofiltro SPA, Santiago, Chile

Honors and awards

2023	ASHG Trainee Research Excellence Award (co-first author).
2021	UC Davis Summer Graduate Student Researcher Award.
2017 - 2021	Chilean Government scholarship "BecasChile" (Ph.D.)
2017 - 2021	Fulbright fellowship (Ph.D.)
2014 - 2015	Chilean Government scholarship (M.Sc.)

Publications

* These authors contributed equally to this work.

Peer-reviewed

- Gutierrez Fugón OJ, Sharifi O, Heath N, <u>Soto DC</u>, Gomez JA, Yasui DH, et al. Integration of CTCF loops, methylome, and transcriptome in differentiating LUHMES as a model for imprinting dynamics of the 15q11-q13 locus in human neurons. Hum Mol Genet. 2024; ddae111.
- Chen PB, Chen R, LaPierre N, Chen Z, Mefford J, Marcus E, et al. Complementation testing identifies genes mediating effects at quantitative trait loci underlying fear-related behavior. Cell Genom. 2024; 100545.
- 3. Libé-Philippot B, Lejeune A, Wierda K, Louros N, Erkol E, Vlaeminck I, et al. LRRC37B is a human modifier of voltage-gated sodium channels and axon excitability in cortical neurons. Cell. 2023;186: 5766-5783.e25.
- 4. Behera S, LeFaive J, Orchard P, Mahmoud M, Paulin LF, Farek J, **Soto DC**, et al. FixItFelix: improving genomic analysis by fixing reference errors. Genome Biol. 2023;24: 31.
- 5. **Soto DC***, Uribe-Salazar JM*, Shew CJ*, Sekar A, McGinty SP, Dennis MY. Genomic structural variation: A complex but important driver of human evolution. Am J Biol Anthropol. 2023. doi:10.1002/ajpa.24713
- 6. Nurk S, Koren S, Rhie A, Rautiainen M, Bzikadze AV, Mikheenko A, et al. The complete sequence of a human genome. Science. 2022;376: 44–53.
- 7. Aganezov S*, Yan SM*, **Soto DC***, Kirsche M*, Zarate S*, Avdeyev P, et al. A complete reference genome improves analysis of human genetic variation. Science. 2022;376: eabl3533.
- 8. McCartney AM, Shafin K, Alonge M, Bzikadze AV, Formenti G, Fungtammasan A, et al. Chasing perfection: validation and polishing strategies for telomere-to-telomere genome assemblies. Nat Methods. 2022.
- 9. Zhu Y, Gomez JA, Laufer BI, Mordaunt CE, Mouat JS, **Soto DC**, et al. Placental methylome reveals a 22q13.33 brain regulatory gene locus associated with autism. Genome Biol. 2022;23: 1–32.
- 10. Eshel G*, Araus V*, Undurraga S, **Soto DC**, Moraga C, Montecinos A, et al. Plant ecological genomics at the limits of life in the Atacama Desert. Proc Natl Acad Sci U S A. 2021:118.
- 11. Shew CJ, Carmona-Mora P, <u>Soto DC</u>, Mastoras M, Roberts E, Rosas J, et al. Diverse molecular mechanisms contribute to differential expression of human duplicated genes. Mol Biol Evol. 2021.
- 12. Carrasco-Puga G, Díaz FP, **Soto DC**, Hernández-Castro C, Contreras-López O, Maldonado A, et al. Revealing hidden plant diversity in arid environments. Ecography. 2020;75: 55.
- 13. Miga KH, Koren S, Rhie A, Vollger MR, Gershman A, Bzikadze A, et al. Telomere-to-telomere assembly of a complete human X chromosome. Nature. 2020.
- 14. **Soto DC***, Shew C*, Mastoras M, Schmidt JM, Sahasrabudhe R, Kaya G, et al. Identification of Structural Variation in Chimpanzees Using Optical Mapping and Nanopore Sequencing. Genes. 2020;11: 276.
- 15. Díaz FP, Latorre C, Carrasco-Puga G, Wood JR, Wilmshurst JM, **Soto DC**, et al. Multiscale climate change impacts on plant diversity in the Atacama Desert. Glob Chang Biol. 2019;25: 1733–1745.
- 16. Vidal EA, Moyano TC, Bustos BI, Pérez-Palma E, Moraga C, Riveras E, et al. Whole Genome Sequence, Variant Discovery and Annotation in Mapuche-Huilliche Native South Americans. Sci Rep. 2019;9: 2132.
- Sánchez BJ, <u>Soto DC</u>, Jorquera H, Gelmi CA, Pérez-Correa JR. HIPPO: An Iterative Reparametrization Method for Identification and Calibration of Dynamic Bioreactor Models of Complex Processes. Ind Eng Chem Res. 2014;53: 18514–18525.

Pre-prints

 Soto DC*, Uribe-Salazar JM*, Kaya G, Valdarrago R, Sekar A, Haghani NK, et al. Gene expansions contributing to human brain evolution. bioRxiv. 2024. p. 2024.09.26.615256. doi:10.1101/2024.09.26.615256

- Uribe-Salazar JM, Kaya G, Weyenberg KB, Radke B, Hino KK, <u>Soto DC</u>, et al. Zebrafish models of human-duplicated gene SRGAP2 reveal novel functions in microglia and visual system development. bioRxiv. 2024. p. 2024.09.11.612570. doi:10.1101/2024.09.11.612570.
- 3. Ostridge HJ, Fontsere C, Lizano E, **Soto DC**, Schmidt JM, Saxena V, et al. Local genetic adaptation to habitat in wild chimpanzees. bioRxiv. 2024. p. 2024.07.09.601734. doi:10.1101/2024.07.09.601734.

Book chapters

 Contreras-López O*, Moyano TC*, <u>Soto DC</u>, Gutiérrez RA. Step-by-Step Construction of Gene Co-expression Networks from High-Throughput Arabidopsis RNA Sequencing Data. Root Development. Humana Press, New York, NY; 2018. pp. 275–301.

Other publications

- 1. <u>Soto DC</u>, Kirsche M, Yan SM, Zarate S. The human reference genome is finally complete. TheScienceBreaker. 2023;9. doi:10.25250/thescbr.brk721
- Walker K, Kalra D, Lowdon R, Chen G, Molik D, <u>Soto DC</u>, et al. The third international hackathon for applying insights into large-scale genomic composition to use cases in a wide range of organisms. F1000Res. 2022;11: 530.
- 3. Mc Cartney AM, Mahmoud M, Jochum M, Agustinho DP, Zorman B, Al Khleifat A, et al. An international virtual hackathon to build tools for the analysis of structural variants within species ranging from coronaviruses to vertebrates. F1000Res. 2021;10: 246.

Selected presentations

Talks

- 1. Inaugural Symposium Celebrating Hispanic Heritage Month, Davis, CA. October 2023. "Assessment of duplicated genes in a complete human telomere-to-telomere genome implicates novel paralogs in brain evolution."
- 2. Stanford Genetics Conference on Structural Variants and DNA Repeats SVAR23, Stanford, CA. September 2023. "Long-read Khoe-San genomes reveal structurally divergent loci overlapping genic regions."
- 3. *Invited speaker.* Instituto Nacional de Medicina Genómica, INMIGEN, México. August 2022. "Genetic variant detection in complex genomic regions using long-read sequencing."
- 4. *Invited panelist*. Nanopore Community Meeting [Online]. December 2020. "Identification of Structural Variation in Chimpanzees Using Optical Mapping and Nanopore Sequencing."
- 5. *Invited speaker.* Technology Networks Oxford Nanopore Technologies [Online]. July 2020. "Identification of Structural Variation in Chimpanzees Using Optical Mapping and Nanopore Sequencing."
- 6. 6th Annual UC Davis Health Human Genomics Symposium, Davis, California. November 2019. "Long-read sequencing to assay complex regions of the human genome."
- 7. Best Talk Award. XI Chilean Plant Biology Meeting, Chillán, Chile. November 2016. "Metatranscriptomic approach reveals conserved adaptive processes in Atacama Desert plants."

Posters

- 1. American Society of Human Genetics Meeting, Washington, D.C. November 2023. "Long-read Khoe-San genomes reveal structurally divergent loci overlapping genic regions."
- 2. American Society of Human Genetics Meeting, Los Angeles, CA. October 2022. "Population diversity and selection of recent gene duplications detected using a complete human genome sequence."
- 3. 7th Annual UC Davis Health Human Genomics Symposium. Virtual event. November 2020. "Genomic variant detection within human segmental duplications."
- 4. American Society of Human Genetics Virtual Meeting. Virtual event. October 2020. "Genomic variant detection within human segmental duplications."

- 5. *Best poster award.* Integrative Genetics and Genomics Colloquium, Davis, CA. September 2019. "Genomic variant detection within human-specific segmental duplications."
- 6. Nanopore Community Meeting, San Francisco, CA. November 2018. "Whole-genome sequencing of CHM1 and CHM13 haploid human cell lines for variant discovery in complex genomic regions.

Teaching and mentoring

Courses

University of California, Davis

2024 Winter Guest lecturer. Genomics (1 lecture, 1 bioinformatics lab).
2023 Winter Guest lecturer. Genomics (2 lectures, 1 bioinformatics lab).

2019 Winter Teaching assistant. Genomics.

2018 Summer Volunteer. Data-Intensive Biology Summer Institute: Analyzing High Throughput

Sequencing Data Workshop.

Pontificia Universidad Católica de Chile

2013 Fall Teaching assistant. Mathematics Applied to Process Engineering (MATLAB programming

laboratory).

2013 Spring *Teaching assistant*. Biopolymers.

2012 Fall *Teaching assistant.* Microbial Biotechnology.

2012 Spring *Teaching assistant.* Biopolymers.

Other universities

July 2024 Guest lecturer. Research seminar (1 lecture). Master of Science in Biomolecular

Medicine, School of Medicine, Universidad Autónoma de Sinaloa, Sinaloa, México.

December 2021 Guest lecturer. Central Asia Genomics Workshop: Command-Line Fundamentals.

Undergraduate mentoring

2021 - 2023 Jeffrey Zang. B.S. Computer Science. University of California, Davis.

2019 - 2020 Mira Mastoras. B.S. Cellular and Molecular Biology. University of California, Davis.

Academic service

Reviewer

2024 Ad hoc reviewer Scientific Reports.

2024 Ad hoc reviewer American Society of Human Genetics Annual Meeting.

2024 *Ad hoc* reviewer Science.

2023 Ad hoc reviewer NAR Genomics and Bioinformatics.

University service

2020 - 2021 UC Davis Integrative Genetics and Genomics Graduate Program, Diversity, Equity, and

Inclusion Committee Member.

2019 - 2020 UC Davis Integrative Genetics and Genomics Graduate Program, Student Executive

Committee, Vice-chair.

2018 - 2019 UC Davis Integrative Genetics and Genomics Graduate Program, Student Executive

Committee, Mentoring and Advising Coordinator.

March 2023	UC Davis Postdoctoral Association, 8th Annual Postdoctoral Research Symposium
	Organizing Committee.

Oct 2019 UC Davis Northern California Computational Biology Symposium 2019, Organizing Committee.

Other professional activities

2023 - present	Genetics Society of America Early Career Scientists Program, Communication and Outreach Subcommittee.
2022 - 2023	Data Science Affiliate, UC Davis DataLab.
2022 - 2023	Genetics Society of America Peer Review Training Program, Genome & Systems Biology Group.
Nov 2022	Chilean Society of Bioinformatics Annual Meeting, Organizing Committee.
Oct 2021	Baylor College of Medicine Virtual Structural Variation Hackathon.
Feb 2021	California Undergraduate Bioinformatics Virtual Conference Graduate Student Panel, Invited Panelist.
Jan 2021	UC Davis CompBio Virtual Symposium Graduate Student Panel, Invited Panelist.
Nov 2020	Baylor College of Medicine Virtual Structural Variation Hackathon.
Nov 2016	International Plant Biology Course, Santiago, Chile, Organizing Committee.
July 2016	Chilean Society of Plant Biologists, Logotype Design.

Outreach and public engagement

May 2023	Molecular Biology Laboratory, Sacramento Charter High School, Co-Instructor.
Summer 2022	UC Davis Biochemistry & Molecular Medicine-Sacramento Charter High School Summer Research Program, Co-Mentor.
Apr 2022	Molecular Biology Laboratory, Sacramento Charter High School, Co-Instructor.
Jan - Jun 2022	UC Davis Graduate Students of Color Mentoring Program, Member.
2018 - 2021	UC Davis Picnic Day K-12 science activities, Co-Facilitator.
Oct 2018	SACNAS Student Chapter Panel Discussion, Sacramento State University, Invited Panelist.
2016 - 2017	Chilean Chapter of Girls in Tech, Member.

Media coverage

- [Podcast] Duran, E., & Venegas, L. (May 9, 2022). "Descubriendo los secretos del genoma." En un Mar de Ciencia. https://open.spotify.com/episode/16ket90clPlfcfyOOEBC74?si=xMjj3r32Q92U1VyBHs4ikg
- [Podcast] Aravena, F. (April 25, 2022). "La importancia de la secuenciación completa del genoma humano." El Café Diario, La Tercera.
 https://www.latercera.com/podcast/noticia/el-cafe-diario-la-importancia-de-la-secuenciacion-completa-del-genoma-humano/WFRPQUY4C5F4FNP5T4BV5G227M/
- [Radio Show] Reyes, M. (April 5, 2022). "Las principales conclusiones del último informe del IPCC." Congreso Futuro, Radio Cooperativa.
 - https://www.cooperativa.cl/noticias/sociedad/ciencia/congreso-futuro/congreso-futuro-las-principales-conclusiones-del-ultimo-informe-del-ipcc/2022-04-10/125605.html
- [TV News] Romero, M. (April 5, 2022). "El ADN al descubierto: hablamos con una de las bioinformáticas que ayudó a secuenciar el genoma humano." France 24. https://www.france24.com/es/programas/salud/20220405-adn-secuencia-genoma-humano-daniela-soto

- [Newspaper] Ansede, M. (March 31, 2022). "El primer genoma completo de un ser humano abre una nueva era en la ciencia." El País.
 - https://elpais.com/ciencia/2022-03-31/el-primer-genoma-completo-de-un-ser-humano-abre-una-nueva-era-en-la-ciencia.html
- [University Press Release] Fell, A. (March 31, 2022). "New Human Reference Genome Opens Unexplored Regions." UC Davis News.
 - https://www.ucdavis.edu/health/news/new-human-reference-genome-opens-unexplored-regions
- [University Press Release] Yehya, N. A. (February 15, 2022). "Placenta may hold clues for early autism diagnosis and intervention." UC Davis Health News.
 - https://health.ucdavis.edu/news/headlines/placenta-may-hold-clues-for-early-autism-diagnosis-and-intervention/2022/02

Skills and qualifications

- Operating systems: Linux/Unix.
- Job schedulers: Slurm, Sun Grid Engine.
- Programming languages: Python, R, shell scripting.
- Bioinformatics toolbox: Git/GitHub, Conda/Bioconda, Snakemake, Markdown.
- Web applications: R Shiny, Python Dash.
- Graphic design and illustration: Adobe Illustrator, iPad Procreate.
- Languages: Spanish (native), English (full professional proficiency).