

Current Positions

Investigator, Howard Hughes Medical Institute
Professor, Genome Sciences, University of Washington
Scientific Director, Brotman Baty Institute for Precision Medicine
Lead Scientific Director, Seattle Hub for Synthetic Biology (Allen-CZI-UW)

Contact Information

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Education and Training

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|---------------|--|--------------------------|
| • 1992 – 1996 | Undergraduate studies | Princeton University |
| • 1996 | A.B., <i>summa cum laude</i> (advisor: Lee Silver) | Princeton University |
| • 1996 – 1997 | Fulbright Scholar to India (advisor: Mrudula Phadke) | Sassoon General Hospital |
| • 1997 – 1998 | Research Scientist, Vaccine Division | Merck Research Labs |
| • 1998 – 2007 | Medical Scientist Training Program (MSTP) Candidate | Harvard Medical School |
| • 2005 | Ph.D. (advisor: George Church; Dept. of Genetics) | Harvard University |
| • 2007 | M.D. | Harvard Medical School |

Post-Training Positions

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| • 2024 – present | Lead Scientific Director | Seattle Hub for Synthetic Biology (Allen-CZI-UW) |
| • 2017 – present | Scientific Director | Allen Discovery Center for Cell Lineage Tracing |
| • 2017 – present | Scientific Director | Brotman Baty Institute for Precision Medicine |
| • 2015 – present | Investigator | Howard Hughes Medical Institute |
| • 2015 – present | Full Professor w/ tenure | Dept. of Genome Sciences, University of Washington |
| • 2010 – present | Affiliate Professor | Div. of Human Biology, Fred Hutch Cancer Res. Center |
| • 2011 – 2015 | Associate Professor | Dept. of Genome Sciences, University of Washington |
| • 2007 – 2011 | Assistant Professor | Dept. of Genome Sciences, University of Washington |

Honors, Awards, Named Lectures

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| • 2025 | Election to Membership | National Academy of Medicine |
| • 2022 | Mendel Lecture | European Society of Human Genetics |
| • 2022 | Election to Membership | National Academy of Sciences |
| • 2022 | Election to Membership | National Academy of Inventors |
| • 2022 | Election to Membership | Washington Academy of Sciences |
| • 2019 | Richard Lounsbery Award | National Academy of Sciences |
| • 2019 | AAAS Fellow | American Assc. Advancement of Science |
| • 2019 | Jeffrey M. Trent Lectureship in Cancer Research | National Human Genome Research Institute |
| • 2019 | Paul D. Gottlieb Distinguished Lectureship | University of Texas, Austin |

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| • 2018 | Allan C. Wilson Memorial Lectureship | University of California, Berkeley |
| • 2018 | Richard and Carol Hertzberg Prize | University of California, San Diego |
| • 2018 | Nancy Andrews Physician-Scientist Lectureship | Duke University |
| • 2017 | British Society of Genetic Medicine Lectureship | British Society of Genetic Medicine |
| • 2014 | Cell “40 under 40”, Cell 40th Anniversary | Cell Press |
| • 2014 | 7th Annual Scripps Genomic Medicine Award | Scripps Health |
| • 2014 | HudsonAlpha Prize for Life Sciences | HudsonAlpha Institute for Biotechnology |
| • 2013 | FEDERAprijs | Fed. of Dutch Medical Scientific Societies |
| • 2013 | NIH Director’s Pioneer Award | National Institutes of Health |
| • 2012 | Curt Stern Award | American Society of Human Genetics |
| • 2010 | Lowell Milken Young Investigator | Prostate Cancer Foundation |
| • 2008 | Science in Medicine New Investigator Lecture | University of Washington |
| • 2008 | 3rd Annual Tomorrow’s Pls | Genome Technology Magazine |
| • 2007 | James Tolbert Shipley Prize | Harvard Medical School |
| • 2006 | TR35 Young Innovator Award | M.I.T. Technology Review |
| • 1998 | Medical Science Training Program Fellowship | National Institutes of Health |
| • 1996 | Fulbright Scholarship | U.S. State Department |
| • 1996 | <i>summa cum laude</i> | Princeton University |
| • 1996 | Honorary Major in Anthropology | Princeton University |
| • 1996 | Sigma Chi Thesis Award for Molecular Biology | Princeton University |
| • 1996 | Senior Prize for Best Thesis in Anthropology | Princeton University |

Service

Pandemic Response

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|---------------|----------------------|---|
| • 2020 – 2022 | Co-Lead Investigator | Seattle Coronavirus Assessment Network (SCAN) |
| • 2018 – 2022 | Co-Lead Investigator | Seattle Flu Study (SFS) |

Academic, Governmental, or Non-Profit Advisory or Consortium Leadership Roles

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|------------------|-------------------------------------|--|
| • 2024 – present | Scientific & Medical Advisory Board | openRxiv |
| • 2024 – present | Board of Directors | Hypothesis Fund |
| • 2017 – present | Board of Reviewing Editors | Science / AAAS |
| • 2018 – present | Scientific Advisor | Chan Zuckerberg Initiative |
| • 2020 – present | Scientific Advisory Board | New York Genome Center |
| • 2021 – present | Co-Lead, Cancer Basic Biology | Fred Hutch-UW-SCH Cancer Consortium |
| • 2017 – 2022 | Advisory Council | Allen Institute for Cell Science |
| • 2018 – 2022 | Scientific Advisory Board | Allen Institute for Immunology |
| • 2021 – 2022 | Scientific Advisory Board | Open Targets |
| • 2017 – 2020 | Advisory Committee to NIH Director | National Institutes of Health |
| • 2014 – 2018 | National Advisory Council | National Human Genome Research Institute |
| • 2015 | NIH ACD Working Group | AllOfUs / US Precision Medicine Initiative |

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|---------------|---------------------------|--|
| • 2012 – 2014 | Scientific Advisory Board | Joint Genome Institute, DoE |
| • 2012 – 2015 | Steering Committee | NIH/NHGRI Centers for Mendelian Genomics |
| • 2009 – 2012 | Steering Committee | NIH/NHLBI Exome Sequencing Project |

Scientific Meetings & Symposia

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|---------------|--------------|--|
| • 2022 | Co-organizer | 20 th Anniversary Symposium (UW Genome Sciences) |
| • 2019 – 2022 | Co-organizer | Biology of Genomes (Cold Spring Harbor Labs) |
| • 2020 – 2021 | Co-organizer | Hindsight 2020 Series (Allen Institute) |
| • 2015 – 2019 | Co-organizer | Genomics of Rare Diseases (Wellcome / Sanger) |
| • 2018 | Co-organizer | Symposium: The Personal Genome (UW Genome Sciences / BBI) |
| • 2014 | Co-organizer | Symposium: Genetic Networks (UW Genome Sciences) |
| • 2010 | Co-organizer | Symposium: Healthcare Implications of Medical Discoveries (UW) |

Peer or Program Review

Note: my roles as an advisor on the NIH ACD and/or NACHGR precluded NIH CSR service from 2015-2020. The sole exception below was due to an oversight.

- 2024 Reviewer, Hanna H. Gray Fellows Competition, Howard Hughes Medical Institute
- 2023 Chair, NHGRI Multi-Omics for Health and Disease Special Emphasis Panel, NIH
- 2023 Reviewer, Investigator Competition, Howard Hughes Medical Institute
- 2022 Reviewer, Hanna H. Gray Fellows Competition, Howard Hughes Medical Institute
- 2022 Chair, NHGRI Single Molecule Protein Sequencing Special Emphasis Panel, NIH
- 2021 Reviewer, Investigator Competition, Howard Hughes Medical Institute
- 2021 Reviewer, Investigator Competition, Chan Zuckerberg Biohub
- 2020 Reviewer, Wellcome Sanger Quinquennial Review
- 2018 Reviewer, Investigator Competition, Howard Hughes Medical Institute
- 2017 Reviewer, International Scholars Competition, Howard Hughes Medical Institute
- 2017 Reviewer, Advanced Genomic Technology Development Special Emphasis Panel, NIH
- 2016 Reviewer, Faculty Scholars Competition, Howard Hughes Medical Institute
- 2014 Reviewer, Paul G. Allen Family Foundation ADI 2014 Life Science Focus
- 2014 Reviewer, TEDDY Whole Genome Sequencing Lab RFP, NIH
- 2014 Reviewer, NIDDK Special Emphasis Panel, NIH
- 2013 Reviewer, NICHD Special Emphasis Panel, NIH
- 2013 Reviewer, 63th Annual Meeting of American Society of Human Genetics
- 2013 Reviewer, The Wellcome Trust
- 2011 Reviewer, W. M. Keck Foundation
- 2011 Reviewer, Lasker Clinical Research Scholars Program
- 2010 Reviewer, UK Medical Research Council, Molecular and Cellular Medicine Board
- 2009 Reviewer, National Science Foundation
- 2009 Reviewer, NIH ARRA Challenge Grants (Genes, Genomes and Genetics IRG), NIH
- 2009 Reviewer, Ontario Research Fund (GL2 Competition)
- 2008 Reviewer, Genome BritishColumbia

Teaching

- 2025 Lead “Developmental Genomics” (UW undergraduate seminar)
- 2008 – present Co-Lead “Methods and Logic in Genetics” (UW, graduate seminar)
- 2017, 2019, 2021 Co-Lead “Genomics & Proteomics” (UW, undergraduate lecture course)
- 2012 – 2016 Co-Lead “Genetics” (UW, pre-clinical med school requirement)
- 2012 – 2015 Co-Lead “Genetic Anatomy” (UW, pre-clinical med school elective)
- 2010 – 2012 Co-Lead “Genome Informatics” (UW, undergraduate lecture course)
- 2001 – 2003 TA “Principles of Pharmacology” (HMS, pre-clinical med school requirement)

Faculty Administrative Responsibilities

- 2025 Member, Admissions Committee, MCB Graduate Program (UW & Fred Hutch)
- 2024 Member, Search Committee for the Director, Office of Strategic Coordination, NIH
- 2022 – 2023 Chair, Faculty Search Committee (UW Genome Sciences)
- 2013 – 2014 Chair, Seminar Series Committee (UW Genome Sciences)
- 2012 – 2013 Co-Chair, UW Scientific Discovery Subcommittee for Curriculum Renewal
- 2009 Member, UW “Two Years to Two Decades” (2y2d) initiative, Discovery focus group
- 2007 – 2021 UW Faculty Search Committees: Genome Sciences (2020-21, 2017-18, 2010-12, 2008-9), Biology (2016-17), Biochemistry (2013-17), Medical Genetics (2008-13)
- 2008 – 2022 UW Genome Sciences Seminar Series Committee (2021-22, 2014-15, 2008-9)

Journal Editorial or Advisory Boards

Current: *Science*, *Cell Genomics*, *Genetics*, *Genome Biology*, *Genome Medicine*, *Genome Research*, *Human Genetics*

Previous: *Biotechniques* (2011-2018), *American Journal of Human Genetics* (2009-2012), *Human Molecular Genetics* (2013-2023), *Molecular Case Studies* (2014-2023)

Commercial SAB & Consulting Roles

- 2025 – present 10x Genomics (Scientific Consultant)
- 2024 – present Somite Therapeutics (Founder; Scientific Advisory Board)
- 2022 – present Prime Medicine (Scientific Advisory Board)
- 2022 – present Pacific Biosciences (Scientific Advisory Board)
- 2016 – present Guardant Health (Scientific Consultant)
- 2018 – present Camp4 Therapeutics (Scientific Advisory Board)
- 2015 – present Phase Genomics (Founder; Scientific Advisory Board)
- 2010 – present Adaptive Biotechnologies (Scientific Advisory Board)
- 2022 – 2025 Sixth Street Partners (Scientific Advisory Board)
- 2021 – 2025 Scale Biosciences (Founder; Scientific Advisory Board)
- 2018 – 2025 Maze Therapeutics (Scientific Advisory Board)
- 2020 – 2024 Cajal Neuroscience (Scientific Advisory Board)
- 2009 – 2020 Stratos Genomics (Scientific Advisory Board)

- 2016 – 2019 Nanostring (Scientific Advisory Board)
- 2016 – 2019 Bellwether Bio (Founder; Scientific Consultant)
- 2016 – 2019 Cambridge Epigenetix (Scientific Advisory Board)
- 2013 – 2018 GenePeeks (Scientific Advisory Board)
- 2009 – 2017 Good Start Genetics (Scientific Advisory Board)
- 2013 – 2017 Gen9 (Scientific Advisory Board)
- 2010 – 2015 Ariosa Diagnostics (Scientific Consultant)
- 2013 – 2015 Ingenuity Systems (Scientific Advisory Board)
- 2013 – 2015 Rubicon Genomics (Scientific Advisory Board)
- 2012 Merck Research Laboratories (Scientific Consultant)
- 2010 – 2011 Halo Genomics (Scientific Advisory Board)
- 2008 – 2009 Complete Genomics (Scientific Consultant)
- 2006 Highland Capital Partners (Scientific Consultant)
- 2004 – 2005 Agencourt Biosciences (Scientific Consultant)

Postdoctoral Fellows Trained

- 2025 – present Kshitij Rai, Ph.D.
- 2025 – present Sophie Seidel, Ph.D.
- 2025 – present Jongbeom Park, Ph.D.
- 2024 – present Mohammed Aljohani, Ph.D. (joint with Sud Pinglay)
- 2024 – present Lijia Li, Ph.D.
- 2024 – present Jonas Koeppel, Ph.D. (joint with Sud Pinglay)
- 2024 – present Zach Stevenson, Ph.D.
- 2024 – present Zukai Liu, Ph.D. (joint with Nobu Hamazaki)
- 2023 – present Yi Fu, Ph.D.
- 2022 – present Haedong Kim, Ph.D.
- 2021 – present Riddhiman Garge, Ph.D. (joint with Lea Starita)
- 2020 – present Eva Nichols, Ph.D. (joint with Brian Beliveau)
- 2020 – 2026 Troy McDiarmid, Ph.D. (Assistant Professor, University of British Columbia)
- 2021 – 2025 Kamen Simeonov, Ph.D. (joint with Chris Lengner; Assistant Professor, Fred Hutch Cancer Center)
- 2022 – 2025 Xiaoyi Li, Ph.D. (Assistant Professor, Westlake University)
- 2020 – 2024 Jean-Benoît Lalanne, Ph.D. (Assistant Professor, University of Montreal)
- 2019 – 2024 Diego Calderon, Ph.D. (joint with Cole Trapnell; Assistant Professor, UCSF)
- 2019 – 2024 Junhong Choi, Ph.D. (Assistant Professor, Memorial Sloan Kettering Cancer Center)
- 2023 – 2024 Sudarshan (Sud) Pinglay Ph.D. (SeaHub Faculty Investigator; Research Assistant Professor, University of Washington)
- 2022 – 2023 Elizabeth Vincent, Ph.D. (joint with David Beier; Molecular Genetics Analysis Assistant, Baylor Genetics)
- 2021 – 2023 Sanjay Srivatsan, Ph.D. (joint with Cole Trapnell; Assistant Professor, Fred Hutch Cancer Center)
- 2020 – 2022 Nobu Hamazaki, Ph.D. (Assistant Professor, University of Washington)

- 2020 – 2022 Alexander Boulgakov, Ph.D. (Senior Bioinformatics Engineer, Twist Bioscience)
- 2018 – 2022 Silvia Domcke Ph.D. (Assistant Professor, University of Zurich)
- 2020 – 2021 Jase Gehring, Ph.D. (Associate Project Scientist, UC Berkeley)
- 2018 – 2021 Jacob Tome, Ph.D. (Senior Scientist, Shape Therapeutics)
- 2018 – 2020 Ronnie Blecher, Ph.D. (Associate Researcher, Weizmann Institute of Science)
- 2016 – 2020 Yi Yin, Ph.D. (Assistant Professor, Human Genetics, UCLA)
- 2015 – 2019 Vikram Agarwal, Ph.D. (Head of mRNA Platform Design Data Science, Sanofi)
- 2014 – 2019 Bridget Kulasekara, Ph.D. (Research Scientist, University of Washington)
- 2014 – 2018 Jes Alexander, Ph.D. (joint with Eric Holland; Assistant Professor, UCSF)
- 2016 – 2018 Malte Spielmann, M.D. (Professor and Chair of Human Genetics, Charité – Universitätsmedizin Berlin, Germany)
- 2014 – 2018 Darren Cusanovich, Ph.D. (Assistant Professor, University of Arizona)
- 2015 – 2017 Lea Starita, Ph.D. (Associate Professor, University of Washington)
- 2012 – 2017 Martin Kircher, Ph.D. (Professor, University of Lübeck)
- 2014 – 2016 Ron Hause, Ph.D. (Head of Computational Sciences, NewLimit)
- 2011 – 2015 Stephen Salipante, M.D., Ph.D. (Professor, University of Washington)
- 2009 – 2013 Jerrod Schwartz, Ph.D. (Chief Technology Officer, ChromaCode)
- 2009 – 2013 Brian O’Roak, Ph.D. (joint with Evan Eichler; Professor, OHSU)
- 2007 – 2009 Emily Turner, Ph.D. (Senior Program Officer, Bill & Melinda Gates Foundation)

Graduate Students Trained

- 2025 – Present Carina Biar (Genome Sciences)
- 2025 – Present Ariana Farrell (Molecular & Cellular Biology)
- 2024 – Present Sanjay Kottapalli (Genome Sciences)
- 2024 – Present Abby McGee (Genome Sciences)
- 2023 – Present Qi Yu (Genome Sciences)
- 2022 – Present Jenny Nathans (Medical Scientist Training Program; Genome Sciences)
- 2022 – Present David Lee (Genome Sciences; joint with David Baker)
- 2022 – Present Shruti Jain (Genome Sciences)
- 2022 – Present Connor Kubo (Genome Sciences; joint with Nobu Hamazaki)
- 2021 – Present Hanna Liao (Molecular & Cellular Biology)
- 2021 – Present Tony Li (Genome Sciences)
- 2021 – Present Aidan Keith (Genome Sciences)
- 2020 – 2025 Chase Suiter (Molecular & Cellular Biology; dissertation entitled “Multiplex Methods for Profiling and Programming Protein Degradation”; Postdoc, Shendure Lab)
- 2020 – 2025 Wei Yang (Genome Sciences; dissertation entitled “Modeling temporal dynamics of early embryogenesis and aging”; Scientist, Illumina)
- 2018 – 2024 Sam Regalado (Medical Scientist Training Program; Genome Sciences; joint with Cole Trapnell; dissertation entitled “Scalable methods for genomic analysis of *in vitro* models of mammalian embryogenesis”; Medical Student, UW)
- 2019 – 2025 Chengxiang (CX) Qiu (Genome Sciences; dissertation entitled “Single-cell Analysis Reveals the Molecular Roadmap of Mouse Embryogenesis”; Assistant Professor, Dartmouth College)

- 2018 – 2023 Florence Chardon (Genome Sciences; joint with Lea Starita; dissertation entitled “CRISPR-based functional genomics to study gene regulatory architecture and functional consequences of genetic variation”; Scientist, Seattle Hub for Synthetic Biology)
- 2018 – 2023 Xingfang (Fanny) Huang (Computer Science & Engineering; dissertation entitled “Computational methods of high-dimensional datasets derived from molecular profiling of biological systems”; Computational Biologist, Calico Life Sciences)
- 2017 – 2022 Anna Minkina (Genome Sciences; dissertation entitled “Tethering distinct molecular profiles of single cells by their lineage histories to investigate sources of cell state heterogeneity”; Computational Biologist, Stergachis Lab)
- 2016 – 2022 Wei (Will) Chen (Molecular Engineering; dissertation entitled “Multiplex Molecular Recording of Biological Signals and Events”; Postdoctoral Fellow, Baker Lab)
- 2015 – 2019 Molly Gasperini (Genome Sciences; dissertation entitled “Efficiently searching for enhancers and their target genes in the human genome”; Associate Director, Seattle Hub for Synthetic Biology)
- 2015 – 2019 Andrew Hill (Genome Sciences; joint with Cole Trapnell; dissertation entitled “Expanding the scope and utility of single-cell genomic technologies”; Scientist, Infinimmune)
- 2014 – 2019 Seungsoo Kim (Genome Sciences; dissertation entitled “Maps and mechanisms of three-dimensional genome organization”; Assistant Professor, UC Irvine)
- 2016 – 2019 Junyue Cao (Molecular & Cellular Biology; dissertation entitled “Cell state and fate characterization by high-throughput single cell genomics”; Associate Professor, Rockefeller University)
- 2015 – 2019 Hannah Pliner (Genome Sciences; joint with Cole Trapnell; dissertation entitled “Algorithms for modeling gene regulation and determining cell type using single-cell molecular profiles”; Senior Principal Scientist, Bristol Myers Squibb)
- 2015 – 2018 Jason Klein (Medical Scientist Training Program; Genome Sciences; dissertation entitled “Massively Parallel Characterization of Enhancers in Evolution and Disease”; Assistant Professor, University of Colorado Anschutz)
- 2015 – 2018 Greg Findlay (Medical Scientist Training Program; Genome Sciences; dissertation entitled “High-throughput interrogation of genome function and cellular lineage”; Group Leader, Crick Institute)
- 2014 – 2017 Vijay Ramani (Genome Sciences; dissertation entitled “Massively parallel analysis of nucleic acid structure”; Associate Professor, Gladstone Institutes & UCSF)
- 2013 – 2017 Aaron McKenna (Genome Sciences; dissertation entitled “Whole-organism lineage tracing by combinatorial and cumulative genome editing”; Assistant Professor, Dartmouth College)
- 2012 – 2016 Matthew Snyder (Genome Sciences; dissertation entitled “Expanding the accuracy, resolution, and breadth of cell-free DNA investigation”; Research Scientist, Brotman Baty Institute for Precision Medicine)
- 2011 – 2014 Joshua Burton (Genome Sciences; dissertation entitled “New methods for de novo assembly of genomes and metagenomes”; Associate Director, Natera)
- 2010 – 2014 Akash Kumar (Medical Scientist Training Program; Genome Sciences; dissertation entitled “Mutational Heterogeneity in Cancer: Lessons from the Brain and Prostate”; Chief Medical & Scientific Officer, MyOme).
- 2010 – 2014 Andrew Adey (Molecular & Cellular Biology; dissertation entitled “Comprehensive, precision genomics”; Professor, OHSU)
- 2009 – 2013 Jacob Kitzman (Genome Sciences; dissertation entitled “New technologies for

- 2009 – 2012 sequencing and interpreting genomes”; Associate Professor, University of Michigan)
Joseph Hiatt (Medical Scientist Training Program; Genome Sciences; dissertation entitled “Molecular tagging to overcome limitations of massively parallel sequencing”; Director, Bristol Myers Squibb)
- 2007 – 2012 Sarah Ng (Genome Sciences; dissertation entitled “Next Generation Mendelian Genetics”; Faculty, Genome Institute of Singapore)
- 2007 – 2012 Rupali Patwardhan (Genome Sciences; dissertation entitled “Massively parallel functional dissection of regulatory elements”; Volunteer, Shendure Lab)

Rotation Students Supervised

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|---------------------|--------------------------------|-------------|
| • Shawn Mohammed | MSTP Program | Spring 2025 |
| • Kristian Choate | Molecular & Cellular Biology | Fall 2024 |
| • PeiXi Chen | Molecular & Cellular Biology | Winter 2025 |
| • Yufei Gao | Genome Sciences | Winter 2024 |
| • Sanjay Kottapalli | Genome Sciences | Fall 2023 |
| • Lucas Kerr | Genome Sciences | Summer 2023 |
| • Qi Yu | Genome Sciences | Winter 2023 |
| • Shruti Jain | Genome Sciences | Spring 2022 |
| • Elliott Swanson | Genome Sciences | Spring 2022 |
| • Sydney Sattler | Genome Sciences | Winter 2022 |
| • Connor Kubo | Genome Sciences | Winter 2022 |
| • David Lee | Genome Sciences | Fall 2021 |
| • Jenny Nathans | Genome Sciences | Summer 2021 |
| • Aidan Keith | Genome Sciences | Spring 2021 |
| • Hanna Liao | Molecular & Cellular Biology | Spring 2021 |
| • Tony Li | Genome Sciences | Fall 2020 |
| • Yuzhen Li | Molecular & Cellular Biology | Spring 2020 |
| • Conor Camplisson | Genome Sciences | Winter 2020 |
| • Wei Yang | Genome Sciences | Winter 2020 |
| • Andrew Mullen | MSTP program | Summer 2019 |
| • Chase Suiter | Molecular & Cellular Biology | Summer 2019 |
| • Shawn Fayer | Genome Sciences | Spring 2019 |
| • Chengxiang Qiu | Genome Sciences | Winter 2019 |
| • James Anderson | Molecular & Cellular Biology | Winter 2019 |
| • Eliza Barkan | Molecular & Cellular Biology | Fall 2018 |
| • Michael Goldberg | Genome Sciences | Spring 2018 |
| • Florence Chardon | Genome Sciences | Spring 2018 |
| • Phillip Dishuck | Genome Sciences | Winter 2018 |
| • William DeWitt | Genome Sciences | Fall 2017 |
| • Xingfang Huang | Computer Science & Engineering | Fall 2017 |
| • Bingie Wang | MSTP program | Summer 2017 |
| • Joseph Janizek | MSTP program | Summer 2017 |
| • Sam Regalado | MSTP program | Summer 2017 |
| • Ian Smith | Genome Sciences | Spring 2017 |
| • April Lo | Genome Sciences | Spring 2017 |
| • Anna Minkina | Genome Sciences | Fall 2016 |
| • Wei Chen | Molecular Engineering | Spring 2016 |
| • Eliah Overbey | Genome Sciences | Spring 2016 |
| • Aakash Sur | Biomedical & Informatics | Fall 2015 |

• Junyue Cao	Molecular & Cellular Biology	Summer 2015
• Molly Gasperini	Genome Sciences	Spring 2015
• Serena Liu	Genome Sciences	Spring 2015
• Hannah Pliner	Genome Sciences	Winter 2015
• Damon May	Genome Sciences	Winter 2015
• Andrew Hill	Genome Sciences	Fall 2014
• Vijay Ramani	Genome Sciences	Winter 2014
• Seungsoo Kim	Genome Sciences	Winter 2014
• Jason Klein	MSTP program	Summer 2013
• Hugh Haddox	Molecular & Cellular Biology	Spring 2013
• Aaron McKenna	Genome Sciences	Winter 2013
• Greg Findlay	MSTP program	Summer 2012
• Matthew Snyder	Genome Sciences	Spring 2012
• Jorgen Nelson	Genome Sciences	Winter 2012
• Elyse Hope	Genome Sciences	Winter 2012
• Meara Davies	Molecular & Cellular Biology	Fall 2011
• Josh Burton	Genome Sciences	Winter 2011
• Jenny Wagner	Genome Sciences	Winter 2011
• Andrew Adey	Molecular & Cellular Biology	Fall 2009
• David Young	MSTP program	Summer 2009
• Akash Kumar	MSTP program	Summer 2009
• Jacob Kitzman	Genome Sciences	Spring 2009
• Keisha Carlson	Genome Sciences	Winter 2009
• Jarrett Egerston	Genome Sciences	Winter 2009
• Matthew Maurano	Genome Sciences	Fall 2008
• Joseph Hiatt	MSTP program	Summer 2008
• Sayer Herrin	Genome Sciences	Winter 2008
• Rupali Patwardhan	Genome Sciences	Winter 2008
• Sarah Ng	Genome Sciences	Fall 2007

Graduate Student Committees (in addition to own trainees)

• 2025 –	Yash Sonthalia	Genome Sciences	Advisor: Pinglay & Noble
• 2025 –	Karl Young	Genome Sciences	Advisor: Cole Trapnell
• 2025 –	Lucas Kerr	Genome Sciences	Advisor: Sanjay Srivatsan
• 2024 –	Clara McCurdy	Biochemistry	Advisor: H. Ruohola-Baker
• 2024 –	Morgan Bean	Bioengineering	Advisor: Hao Kueh
• 2024 –	Connor Kelly	Genome Sciences	Advisor: Brian Beliveau
• 2023 –	Ashmita Rajendran	BIME	Advisor: Siobhan Pattwell
• 2023 –	Sahar Attar	Genome Sciences	Advisor: Devin Schweppe
• 2023 –	Conor Camplisson	Genome Sciences	Advisor: Brian Beliveau
• 2022 –	Elliott Swanson	Genome Sciences	Advisor: Andrew Stergachis
• 2022 – 2025	Jack Castelli	LabMed & Pathology	Advisor: Jennifer Adair
• 2021 – 2025	Maddy Duran	Genome Sciences	Advisor: Cole Trapnell
• 2020 – 2024	Eliza Barkan	Genome Sciences	Advisor: Cole Trapnell
• 2021 – 2024	Shangqing Jiang	School of Pharmacy	Advisor: David Veenstra
• 2019 – 2024	Robin Aguilar	Genome Sciences	Advisor: Bill Noble
• 2019 – 2023	Gesine Cauer	Genome Sciences	Advisor: Bill Noble
• 2019 – 2020	Jacob Schreiber	Comp. Science & Engineering	Advisor: Bill Noble
• 2018 – 2022	Andria Ellis	Genome Sciences	Advisor: Cole Trapnell
• 2018 – 2022	Ian Smith	Genome Sciences	Advisor: Judit Villen

• 2018 – 2021	Sanjay Srivatsan	Genome Sciences	Advisor: Cole Trapnell
• 2018 – 2019	Peter Ney	Comp. Science & Engineering	Advisor: Tadayoshi Kohno
• 2017 – 2021	Robin Kirkpatrick	Genome Sciences	Advisor: Jesse Zalatan
• 2016 – 2021	Nuttada Panpradist	Bioengineering	Advisor: Barry Lutz
• 2016 – 2020	Wei Zhou	Molecular & Cellular Biology	Advisor: Stan Fields
• 2016 – 2020	Clara Amorosi	Genome Sciences	Advisor: Maitreya Dunham
• 2016 – 2018	Rebecca Zaunbrecher	Bioengineering	Advisor: Mike Regnier
• 2016 – 2019	Aaron Wolf	Genome Sciences	Advisor: Josh Akey
• 2017 – 2020	Ian Nova	Molecular Engineering	Advisor: Jens Gundlach
• 2015 – 2020	Melissa Chiasson	Genome Sciences	Advisor: Doug Fowler
• 2015 – 2019	John (Ty) Crowl	Immunology	Advisor: Dan Stetson
• 2015 – 2017	Jocelynn Pearl	Molecular & Cellular Biology	Advisor: Lee Hood
• 2015 – 2016	Alexander Rosenberg	Electrical Engineering	Advisor: Georg Seelig
• 2014 – 2019	Piero Lamelza	Molecular & Cellular Biology	Advisor: Michael Ailion
• 2014 – 2017	Hugh Haddox	Molecular & Cellular Biology	Advisor: Jesse Bloom
• 2013 – 2018	Jorgen Nelson	Genome Sciences	Advisor: David Baker
• 2013 – 2016	David Young	Genome Sciences	Advisor: Stan Fields
• 2012 – 2015	Benjamin Vernot	Genome Sciences	Advisor: Josh Akey
• 2012 – 2014	Andrew Laszlo	Physics	Advisor: Jens Gundlach
• 2012 – 2014	Niklas Krumm	Genome Sciences	Advisor: Evan Eichler
• 2011 – 2017	Jennifer Andrie	Genome Sciences	Advisor: Josh Akey
• 2008 – 2015	Vaughn Iverson	Oceanography	Advisor: Virginia Armbrust
• 2012 – 2012	Lucas Gray	Biochemistry	Advisor: Alan Weiner
• 2011 – 2011	Sung Han	Neurobiology and Behavior	Advisor: William Catterall
• 2010 – 2014	Russell Berg	Molecular & Cellular Biology	Advisor: Lalita Ramakrishnan
• 2010 – 2014	Keisha Carlson	Genome Sciences	Advisor: Christine Queitsch
• 2009 – 2014	Leslie Emery	Genome Sciences	Advisor: Josh Akey
• 2010 – 2013	Peter Sudmant	Genome Sciences	Advisor: Evan Eichler
• 2010 – 2013	Thomas White	Molecular & Cellular Biology	Advisor: Peter Nelson
• 2010 – 2013	Benjamin Whiddon	Genome Sciences	Advisor: Richard Palmiter
• 2010 – 2010	Carlos Araya	Genome Sciences	Advisor: Stanley Fields
• 2009 – 2013	Cailyn Spurrell	Genome Sciences	Advisor: Mary-Claire King
• 2009 – 2012	Joshua Bishop	Electrical Engineering	Advisor: Eric Klavins
• 2009 – 2012	Kyle Minch	Molecular & Cellular Biology	Advisor: David Sherman
• 2008 – 2013	Alan Rubin	Genome Sciences	Advisor: Phil Green
• 2008 – 2010	Steven Josefowicz	Immunology	Advisor: Sasha Rudensky
• 2008 – 2010	Kevin Schutz	Genome Sciences	Advisor: Stan Fields
• 2008 – 2010	Marcia Paddock	Immunology	Advisor: Andy Scharenberg

Issued Patents

- Method for large scale scaffolding of genome assemblies (11,694,764)
- Methods of determining tissues and/or cell types giving rise to cell-free dna, and methods of identifying a disease or disorder using same (11,352,670)
- Massively parallel contiguity mapping (10,457,936; 11,299,730)
- Methods for retrieval of sequence-verified DNA constructs (9,809,904)
- Nanogrid rolling circle DNA sequencing (9,624,538)
- Error detection in sequence tag directed subassemblies of short sequencing reads (8,865,410; 10,577,601; 11,505,795)
- Sequence tag directed subassembly of short sequencing reads into long sequencing reads (8,383,345;

8,846,347; 10,227,585; 12,152,236)

- Polony fluorescent in situ sequencing beads (7,425,431)

Published Patent Applications

- Multiplex, temporally resolved molecular signal recorder and related methods (20240355418)
- Bacterial DNA cytosine deaminases for mapping DNA methylation sites (20240124867)
- Precise genome deletion and replacement method based on prime editing (20240011055)
- High-throughput single-cell transcriptome libraries and methods of making and using (20210102194)
- Diagnosis of cancer or other physiological condition using circulating nucleic acid fragment sentinel endpoints (20200255905)
- Determining a physiological condition in an individual by analyzing cell-free DNA fragment endpoints in a biological sample (20190309374)
- High-throughput single-cell sequencing with reduced amplification bias (20190382753)
- Multiplex pairwise assembly of DNA oligonucleotides (application; 20180320166)
- A framework for determining the relative effects of genetic mutations (20160357903)
- Methods and systems for large scale scaffolding of genome assemblies (20160239602)
- Multiplex homology-directed repair (20160076093)
- Systems, Algorithms, and Software for Molecular Inversion Probe (MIP) Design (20160055293)
- Highly multiplex single amino acid mutagenesis for massively parallel functional analysis (20160017410)
- Whole genome sequencing of a human fetus (20150105267)
- Multiplex decoding of sequence tags in barcodes (20080269068)
- Wobble sequencing (20070207482)
- Nucleic acid memory device (20030228611)

Preprints & Publications (* denotes equal contributors; # or ^ denotes corresponding or senior authorship; gray numbers denote primary publications, defined as those on which I and/or a member of my lab are a corresponding, senior and/or a first author)

Preprints

22. "Elucidating Neurodevelopmental Trajectories in Cancer with Topic Modeling: Revealing Persistent External Granule Layer Lineages in Medulloblastoma." [*bioRxiv*](#) 2025.11.12.687706 (posted 13-November-2025).
21. "Large-scale discovery of neural enhancers for cis-regulation therapies." [*bioRxiv*](#) 2025.11.04.686611 (posted 5-November-2025).
20. "Single-cell, multi-region profiling of the macaque brain across the lifespan." [*bioRxiv*](#) 2025.10.31.685880 (posted 1-November-2025).
19. "Multiplex design and discovery of proximity handles for programmable proteome editing." [*bioRxiv*](#) 2025.10.13.681693 (posted 13-October-2025).
18. "Dual-patterned pluripotent stem cells self-organize into a human embryo model with extended anterior-posterior patterning." [*bioRxiv*](#) 2025.09.25.678678 (posted 25-September-2025).
17. "Capture-C MPRA: A high-throughput method to simultaneously characterize promoter interactions and regulatory activity." [*bioRxiv*](#) 2025.06.11.658967 (posted 14-Jun-2025).

16. "Barcoded monoclonal embryoids are a potential solution to confounding bottlenecks in mosaic organoid screens." [*bioRxiv*](#) 2025.05.23.655669 (posted 24-May-2025).
15. "Lineage recording in monoclonal gastruloid reveals heritable modes of early development." [*bioRxiv*](#) 2025.05.23.655664 (posted 24-May-2025).
14. "Technical and biological sources of noise confound multiplexed enhancer AAV screening." [*bioRxiv*](#) 2025.01.14.633018 (posted 16-January-2025).
13. "Extensive length and homology dependent chimerism in pool-packaged AAV libraries." [*bioRxiv*](#) 2025.01.14.632594 (posted 15-January-2025).
12. "Sciphy: A Bayesian phylogenetic framework using sequential genetic lineage tracing data." [*bioRxiv*](#) 2024.10.01.615771 (posted 12-Oct-2024).
11. "The proteomic landscape and temporal dynamics of mammalian gastruloid development." [*bioRxiv*](#) 2024.09.05.609098 (posted 18-Sep-2024).
10. "Leukemia-derived apelin selects endothelial niche clones to promote tumorigenesis." [*bioRxiv*](#) 2024.09.09.612077 (posted 13-Sep-2024).
9. "Optics-free reconstruction of 2D images via DNA barcode proximity graphs." [*bioRxiv*](#) 2024.08.06.606834 (posted 8-Aug-2024).
8. "SARS-CoV-2 diversity and transmission on a university campus across two academic years during the pandemic." [*medRxiv*](#) 2024.02.29.24303285 (posted 2-Mar-2024).
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6. "High Density Domain-Focused CRISPR Screens Reveal Novel Epigenetic Regulators of HOX/MEIS Activation in Acute Myeloid Leukemia." [*bioRxiv*](#) 2022.12.12.519332 (posted 6-Mar-2023)
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4. "Interactions among 17 respiratory pathogens: a cross-sectional study using clinical and community surveillance data." [*medRxiv*](#) 2022.02.04.22270474 (posted 6-Feb-2022).
3. "Viral genome sequencing places White House COVID-19 outbreak into phylogenetic context." [*medRxiv*](#) 2020.10.31.20223925 (posted 31-Oct-2020).
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