(949) 751-7015

Qualification summary

- Over ten years experience with large-scale biological data (genome sequence, population variation, epigenetic and TF ChIP-chip/seq, gene expression RNA-seq/microarrays)
- o Expertise in machine learning, statistics, computational modeling, high-throughput computing, experiment design
- Experience with C/C++, Java, R, MATLAB, Python, HTML, Bash
- Member of several NIH consortia (12 Drosophila, 29 Mammals, ENCODE, modENCODE, Epigenomics Roadmap, GTEx)

Education

Massachusetts Institute of Technology at Cambridge, MA

August 2005-February 2012

Ph.D. in Computer Science Advisor: Manolis Kellis

Thesis title: Computational Regulatory Genomics: Motifs, Networks, and Dynamics

University of Illinois at Urbana-Champaign

August 2001-May 2005

B.S. in Computer Science M.S. in Computer Science

Advisors: ChengXiang Zhai, Gene Robinson, and Charles Whitfield

Thesis title: Computational Methods for Modeling Eukaryotic Gene Regulation

Research Experience

Verily Life Sciences—Staff Scientist
 Verily Life Sciences—Senior Scientist
 Verily Life Sciences—Scientist
 Google[x]—Quantitative Analyst
 Computational biology research and development

February 2012-July 2015

November 2015-April 2017

July 2015-November 2015

March 2020-Present

April 2017-March 2020

Designed and analyzed high-throughput enhancer experiments; characterized RNA edited sites across hundreds of individuals and dozens of tissues; predicted links between enhancers and genes; developed model to distinguish individuals from NGS data

o MIT CSAIL—Graduate Research Assistant

o MIT CSAIL—Postdoctoral Associate

August 2005-February 2012

Developed and applied computational algorithms for understanding regulatory dynamics using regulatory motifs and epigenetic data

o **UIUC Entomology**—Research Assistant

February 2003-August 2005

Applied computational methods for microarray and sequence analysis of honey bee and related organisms

• Argonne National Laboratory—Research Intern

Summer 2003

Designed genetic and statistical algorithms for identifying functional sites in BRCA1 proteins

Teaching Experience

• Linguam Connect LLC—Instructor

Summer 2014

- CCP11/CCP13: Designed and taught online course on JavaScript programming
- MIT Electrical Engineering and Computer Science—Graduate Teaching Assistant
 6.095/6.047: Undergraduate/graduate computational biology course; wrote and graded homework and exam problems; led recitation sessions; delivered five guest lectures between 2008-2012
- UIUC Computer Science—Graduate Teaching Assistant
 Spring 2004, Fall 2004, Spring 2005
 CS101: Undergraduate introductory computer science course in MATLAB and C; taught two laboratory sections weekly and helped write and grade exam and laboratory problems

Publications

Evidence of reduced recombination rate in human regulatory domains

Yaping Liu, Abhishek Sarkar, Pouya Kheradpour, Jason Ernst, and Manolis Kellis *Genome Biology*, 2017 October 20, doi:10.1186/s13059-017-1308-x

The Genotype-Tissue Expression (GTEx) pilot analysis: multitissue gene regulation in humans GTEx Consortium

Science, 2015 May 8, doi:10.1126/science.1262110

o Integrative analysis of 111 reference human epigenomes

Roadmap Epigenomics Consortium, Anshul Kundaje*, Wouter Meuleman*, Jason Ernst*, Misha Bilenky*, Angela Yen, Alireza Heravi-Moussavi, Pouya Kheradpour, Zhizhuo Zhang, Jianrong Wang, Michael J. Ziller, Viren Amin, John W. Whitaker, Matthew D. Schultz, Lucas D. Ward, Abhishek Sarkar, Gerald Quon, Richard S. Sandstrom, Matthew L. Eaton, Yi-Chieh Wu, Andreas R. Pfenning, Xinchen Wang, Melina Claussnitzer, Yaping Liu, Cristian Coarfa, R. Alan Harris, Noam Shoresh, Charles B. Epstein, Elizabeta Gjoneska, Danny Leung, Wei Xie, R. David Hawkins, Ryan Lister, Chibo Hong, Philippe Gascard, Andrew J. Mungall, Richard Moore, Eric Chuah, Angela Tam, Theresa K. Canfield, R. Scott Hansen, Rajinder Kaul, Peter J. Sabo, Mukul S. Bansal, Annaick Carles, Jesse R. Dixon, Kai-How Farh, Soheil Feizi, Rosa Karlic, Ah-Ram Kim, Ashwinikumar Kulkarni, Daofeng Li, Rebecca Lowdon, GiNell Elliott, Tim R. Mercer, Shane J. Neph, Vitor Onuchic, Paz Polak, Nisha Rajagopal, Pradipta Ray, Richard C. Sallari, Kyle T. Siebenthall, Nicholas A. Sinnott-Armstrong, Michael Stevens, Robert E. Thurman, Jie Wu, Bo Zhang, Xin Zhou, Arthur E. Beaudet, Laurie A. Boyer, Philip L. De Jager, Peggy J. Farnham, Susan J. Fisher, David Haussler, Steven J. M. Jones, Wei Li, Marco A. Marra, Michael T. McManus, Shamil Sunyaev, James A. Thomson, Thea D. Tlsty, Li-Huei Tsai, Wei Wang, Robert A. Waterland, Michael Q. Zhang, Lisa H. Chadwick, Bradley E. Bernstein, Joseph F. Costello, Joseph R. Ecker, Martin Hirst, Alexander Meissner, Aleksandar Milosavljevic, Bing Ren, John A. Stamatoyannopoulos, Ting Wang, and Manolis Kellis *Nature*, 2015 February 18, doi:10.1038/nature14248

Comparative analysis of regulatory information and circuits across distant species

Alan P. Boyle*, Carlos L. Araya*, Cathleen Brdlik, Philip Cayting, Chao Cheng, Yong Cheng, Kathryn Gardner, LaDeana W. Hillier, Judith Janette, Lixia Jiang, Dionna Kasper, Trupti Kawli, Pouya Kheradpour, Anshul Kundaje, Jingyi Jessica Li, Lijia Ma, Wei Niu, E. Jay Rehm, Joel Rozowsky, Matthew Slattery, Rebecca Spokony, Robert Terrell, Dionne Vafeados, Daifeng Wang, Peter Weisdepp, Yi-Chieh Wu, Dan Xie, Koon-Kiu Yan, Elise A. Feingold, Peter J. Good, Michael J. Pazin, Haiyan Huang, Peter J. Bickel, Steven E. Brenner, Valerie Reinke, Robert H. Waterston, Mark Gerstein, Kevin P. White, Manolis Kellis, and Michael Snyder

Nature, 2014 August 27, doi:10.1038/nature13668

o Diverse patterns of genomic targeting by transcriptional regulators in Drosophila melanogaster

Matthew Slattery*, Lijia Ma*, Rebecca F. Spokony*, Robert K. Arthur, Pouya Kheradpour, Anshul Kundaje, Nicolas Nègre, Alex Crofts, Ryan Ptashkin, Jennifer Zieba, Alexander Ostapenko, Sarah Suchy, Alec Victorsen, Nader Jameel, A. Jason Grundstad, Wenxuan Gao, Jennifer R. Moran, E. Jay Rehm, Robert L. Grossman, Manolis Kellis, and Kevin P. White

Genome Research, 2014 July 1, doi:10.1101/gr.168807.113

Systematic discovery and characterization of regulatory motifs in ENCODE TF binding experiments Pouya Kheradpour and Manolis Kellis

Nucleic Acids Research, 2013 December 13, doi:10.1093/nar/gkt1249

Systematic dissection of regulatory motifs in 2000 predicted human enhancers using a massively parallel reporter assay

Pouya Kheradpour, Jason Ernst, Alexandre Melnikov, Peter Rogov, Li Wang, Xiaolan Zhang, Jessica Alston, Tarjei S. Mikkelsen, and Manolis Kellis

Genome Research, 2013 March 19, doi:10.1101/gr.144899.112

o Analysis of variation at transcription factor binding sites in Drosophila and humans

Mikhail Spivakov, Junaid Akhtar, Pouya Kheradpour, Kathryn Beal, Charles Girardot, Gautier Koscielny, Javier Herrero, Manolis Kellis, Eileen EM Furlong, and Ewan Birney

Genome Biology, 2012 September 5, doi:10.1186/gb-2012-13-9-r49

An integrated encyclopedia of DNA elements in the human genome

The ENCODE Project Consortium

Nature, 2012 September 6, doi:10.1038/nature11247

ChIP-seg guidelines and practices of the ENCODE and modENCODE consortia

Stephen G. Landt*, Georgi K. Marinov*, Anshul Kundaje*, Pouya Kheradpour, Florencia Pauli, Serafim Batzoglou, Bradley E. Bernstein, Peter Bickel, James B. Brown, Philip Cayting, Yiwen Chen, Gilberto DeSalvo, Charles Epstein, Katherine I. Fisher-Aylor, Ghia Euskirchen, Mark Gerstein, Jason Gertz, Alexander J. Hartemink, Michael M. Hoffman, Vishwanath R. Iyer, Youngsook L. Jung, Subhradip Karmakar, Manolis Kellis, Peter V. Kharchenko, Qunhua Li, Tao Liu, X. Shirley Liu, Lijia Ma, Aleksandar Milosavljevic, Richard M. Myers, Peter J. Park, Michael J. Pazin, Marc D. Perry, Debasish Raha, Timothy E. Reddy, Joel Rozowsky, Noam Shoresh, Arend Sidow, Matthew Slattery, John A. Stamatoyannopoulos,

Michael Y. Tolstorukov, Kevin P. White, Simon Xi, Peggy J. Farnham, Jason D. Lieb, Barbara J. Wold, and, Michael Snyder

Genome Research, 2012 September 5, doi:10.1101/gr.136184.111

Computational Regulatory Genomics: Motifs, Networks, and Dynamics Pouya Kheradpour

Ph.D. Thesis, MIT, 2012 February 3, doi:1721.1/70871

o A high-resolution map of human evolutionary constraint using 29 mammals

Kerstin Lindblad-Toh, Manuel Garber*, Or Zuk*, Michael F. Lin*, Brian J. Parker*, Stefan Washietl*, Pouya Kheradpour*, Jason Ernst*, Gregory Jordan*, Evan Mauceli*, Lucas D. Ward*, Craig B. Lowe*, Alisha K. Holloway*, Michele Clamp*, Sante Gnerre*, Jessica Alföldi, Kathryn Beal, Jean Chang, Hiram Clawson, James Cuff, Federica Di Palma, Stephen Fitzgerald, Paul Flicek, Mitchell Guttman, Melissa J. Hubisz, David B. Jaffe, Irwin Jungreis, W. James Kent, Dennis Kostka, Marcia Lara, Andre L. Martins, Tim Massingham, Ida Moltke, Brian J. Raney, Matthew D. Rasmussen, Jim Robinson, Alexander Stark, Albert J. Vilella, Jiayu Wen, Xiaohui Xie, Michael C. Zody, Broad Institute Sequencing Platform Whole Genome Assembly Team, Kim C. Worley, Christie L. Kovar, Donna M. Muzny, Richard A. Gibbs, Baylor College of Medicine Human Genome Sequencing Center Sequencing Team, Wesley C. Warren, Elaine R. Mardis, George M. Weinstock, Richard K. Wilson, Genome Institute at Washington University, Ewan Birney, Elliott H. Margulies, Javier Herrero, Eric D. Green, David Haussler, Adam Siepel, Nick Goldman, Katherine S. Pollard, Jakob S. Pedersen, Eric S. Lander, and Manolis Kellis

Nature, 2011 October 27, doi:10.1038/nature10530

Locating protein-coding sequences under selection for additional, overlapping functions in 29 mammalian genomes

Michael F. Lin, Pouya Kheradpour, Stefan Washietl, Brian J. Parker, Jakob S. Pedersen, and Manolis Kellis Genome Research, 2011 October 12, doi:10.1101/gr.108753.110

o An Epigenetic Signature for Monoallelic Olfactory Receptor Expression

Angeliki Magklara, Angela Yen*, Bradley M. Colquitt*, E. Josephine Clowney*, William Allen, Eirene Markenscoff-Papadimitriou, Zoe A. Evans, <u>Pouya Kheradpour</u>, George Mountoufaris, Catriona Carey, Gilad Barnea, Manolis Kellis, and Stavros Lomvardas

Cell, 2011 May 13, doi:10.1016/j.cell.2011.03.040

A User's Guide to the Encyclopedia of DNA Elements (ENCODE)

The ENCODE Project Consortium

PLoS Biology, 2011 April 19, doi:10.1371/journal.pbio.1001046

Mapping and analysis of chromatin state dynamics in nine human cell types

Jason Ernst, Pouya Kheradpour, Tarjei S. Mikkelsen, Noam Shoresh, Lucas D. Ward, Charles B. Epstein, Xiaolan Zhang, Li Wang, Robbyn Issner, Michael Coyne, Manching Ku, Timothy Durham, Manolis Kellis*, and Bradley E. Bernstein* *Nature*, 2011 May 5, doi:10.1038/nature09906

• A cis-regulatory map of the *Drosophila* genome

Nicolas Nègre*, Christopher D. Brown*, Lijia Ma*, Christopher Aaron Bristow*, Steven W. Miller*, Ulrich Wagner*, Pouya Kheradpour, Matthew L. Eaton, Paul Loriaux, Rachel Sealfon, Zirong Li, Haruhiko Ishii, Rebecca F. Spokony, Jia Chen, Lindsay Hwang, Chao Cheng, Richard P. Auburn, Melissa B. Davis, Marc Domanus, Parantu K. Shah, Carolyn A. Morrison, Jennifer Zieba, Sarah Suchy, Lionel Senderowicz, Alec Victorsen, Nicholas A. Bild, A. Jason Grundstad, David Hanley, David M. MacAlpine, Mattias Mannervik, Koen Venken, Hugo Bellen, Robert White, Mark Gerstein, Steven Russell, Robert L. Grossman, Bing Ren, James W. Posakony, Manolis Kellis, and Kevin P. White *Nature*, 2011 March 24, doi:10.1038/nature09990

o Identification of functional elements and regulatory circuits by *Drosophila* modENCODE

The modENCODE Consortium, Sushmita Roy*, Jason Ernst*, Peter V. Kharchenko*, Pouya Kheradpour*, Nicolas Nègre*, Matthew L. Eaton*, Jane M. Landolin*, Christopher A. Bristow*, Lijia Ma*, Michael F. Lin*, Stefan Washietl*, Bradley I. Arshinoff*, Ferhat Ay*, Patrick E. Meyer*, Nicolas Robine*, Nicole L. Washington*, Luisa Di Stefano*, Eugene Berezikov, Christopher D. Brown, Rogerio Candeias, Joseph W. Carlson, Adrian Carr, Irwin Jungreis, Daniel Marbach, Rachel Sealfon, Michael Y. Tolstorukov, Sebastian Will, Artyom A. Alekseyenko, Carlo Artieri, Benjamin W. Booth, Angela N. Brooks, Qi Dai, Carrie A. Davis, Michael O. Duff, Xin Feng, Andrey A. Gorchakov, Tingting Gu, Jorja G. Henikoff, Philipp Kapranov, Renhua Li, Heather K. MacAlpine, John Malone, Aki Minoda, Jared Nordman, Katsutomo Okamura, Marc Perry, Sara K. Powell, Nicole C. Riddle, Akiko Sakai, Anastasia Samsonova, Jeremy E. Sandler, Yuri B. Schwartz, Noa Sher, Rebecca Spokony, David Sturgill, Marijke van Baren, Kenneth H. Wan, Li Yang, Charles Yu, Elise Feingold, Peter Good, Mark Guyer, Rebecca Lowdon, Kami Ahmad, Justen Andrews, Bonnie Berger, Steven E. Brenner, Michael R. Brent, Lucy Cherbas, Sarah C. R. Elgin, Thomas R. Gingeras, Robert Grossman, Roger A. Hoskins, Thomas C. Kaufman, William Kent, Mitzi I. Kuroda, Terry Orr-Weaver, Norbert Perrimon, Vincenzo Pirrotta, James W. Posakony, Bing Ren, Steven Russell, Peter Cherbas, Brenton R. Graveley, Suzanna Lewis, Gos Micklem, Brian Oliver, Peter J. Park, Susan

E. Celniker, Steven Henikoff, Gary H. Karpen, Eric C. Lai, David M. MacAlpine, Lincoln D. Stein, Kevin P. White, and Manolis Kellis

Science, 2010 December 24, doi:10.1126/science.1198374

o A comprehensive map of insulator elements for the *Drosophila* genome

Nicolas Nègre*, Christopher D. Brown*, Parantu K. Shah, Pouya Kheradpour, Carolyn A. Morrison, Jorja G. Henikoff, Xin Feng, Kami Ahmad, Steven Russell, Robert A. H. White, Lincoln Stein, Steven Henikoff, Manolis Kellis, and Kevin P. White

PLoS Genetics, 2010 January 15, doi:10.1371/journal.pgen.1000814

o The Tasmanian devil transcriptome reveals Schwann cell origins of a clonally transmissible cancer

Elizabeth P. Murchison, Cesar Tovar, Arthur Hsu, Hannah S. Bender, Pouya Kheradpour, Clare A. Rebbeck, David Obendorf, Carly Conlan, Melanie Bahlo, Catherine A. Blizzard, Stephen Pyecroft, Alexandre Kreiss, Manolis Kellis, Alexander Stark, Timothy T. Harkins, Jennifer A. Marshall Graves, Gregory M. Woods, Gregory J. Hannon, and Anthony T. Papenfuss *Science*, 2010 January 1, doi:10.1126/science.1180616

o Histone modifications at human enhancers reflect global cell-type-specific gene expression

Nathaniel D. Heintzman*, Gary C. Hon*, R. David Hawkins*, <u>Pouya Kheradpour</u>, Alexander Stark, Lindsey F. Harp, Zhen Ye, Leonard K. Lee, Rhona K. Stuart, Christina W. Ching, <u>Keith A. Ching</u>, <u>Jessica E. Antosiewicz-Bourget</u>, Hui Liu, Xinmin Zhang, Roland D. Green, Victor V. Lobanenkov, Ron Stewart, James A. Thomson, Gregory E. Crawford, Manolis Kellis, and Bing Ren *Nature*, 2009 March 18, doi:10.1038/nature07829

o Genome analysis of the platypus reveals unique signatures of evolution

Nature, 2008 May 8, doi:10.1038/nature06936

o Conservation of small RNA pathways in platypus

Elizabeth P. Murchison, Pouya Kheradpour, Ravi Sachidanandam, Carly Smith, Emily Hodges, Zhenyu Xuan, Manolis Kellis, Frank Grützner, Alexander Stark, and Gregory J. Hannon Genome Research, 2008 May 7, doi:10.1101/gr.073056.107

o A single Hox locus in *Drosophila* produces functional microRNAs from opposite DNA strands

Alexander Stark*, Natascha Bushati*, Calvin H. Jan, <u>Pouya Kheradpour</u>, Emily Hodges, Julius Brennecke, David P. Bartel, Stephen M. Cohen, and Manolis Kellis Genes and Development, 2008 January 1, doi:10.1101/gad.1613108

Genes and Development, 2006 January 1, doi:10.1101/gad.1015106

$\circ~$ Discovery of functional elements in 12 Drosophila genomes using evolutionary signatures

Alexander Stark*, Michael F. Lin*, Pouya Kheradpour*, Jakob S. Pedersen*, Leopold Parts, Joseph W. Carlson, Madeline A. Crosby, Matthew D. Rasmussen, Sushmita Roy, Ameya N. Deoras, J. Graham Ruby, Julius Brennecke, Harvard FlyBase curators, Berkeley Drosophila Genome Project, Emily Hodges, Angie S. Hinrichs, Anat Caspi, Benedict Paten, Seung-Won Park, Mira V. Han, Morgan L. Maeder, Benjamin J. Polansky, Bryanne E. Robson, Stein Aerts, Jacques van Helden, Bassem Hassan, Donald G. Gilbert, Deborah A. Eastman, Michael Rice, Michael Weir, Matthew W. Hahn, Yongkyu Park, Colin N. Dewey, Lior Pachter, W. James Kent, David Haussler, Eric C. Lai, David P. Bartel, Gregory J. Hannon, Thomas C. Kaufman, Michael B. Eisen, Andrew G. Clark, Douglas Smith, Susan E. Celniker, William M. Gelbart, and Manolis Kellis

Nature, 2007 November 8, doi:10.1038/nature06340

Evolution of genes and genomes on the *Drosophila* phylogeny

Drosophila 12 Genomes Consortium

Nature, 2007 November 8, doi:10.1038/nature06341

o Reliable prediction of regulator targets using 12 Drosophila genomes

Pouya Kheradpour*, Alexander Stark*, Sushmita Roy, and Manolis Kellis *Genome Research*, 2007 November 7, doi:10.1101/gr.7090407

Systematic discovery and characterization of fly microRNAs using 12 Drosophila genomes

Alexander Stark*, Pouya Kheradpour*, Leopold Parts, Julius Brennecke, Emily Hodges, Gregory J. Hannon, and Manolis Kellis

Genome Research, 2007 November 7, doi:10.1101/gr.6593807

Beyond the Bits: Cooperative Packet Recovery Using Physical Layer Information

Grace R. Woo, <u>Pouya Kheradpour</u>, Dawei Shen, and Dina Katabi *ACM MobiCom*, 2007 September 9, doi:10.1145/1287853.1287871

o Fold-specific substitution matrices for protein classification

Richard B. Vilim, R. M. Cunningham, Bo Lu, Pouya Kheradpour, and F. J. Stevens *Bioinformatics*, 2004 February 5, doi:10.1093/bioinformatics/btg492

Selected presentations

| 0 | Regulatory motif centric validation, dissection, and construction of transcriptional enhancers Selected talk, ENCODE Consortium Meeting, Stanford, CA | May 2013 |
|---|--|---------------|
| 0 | Systematic prediction and validation of enhancer regulatory motifs in ENCODE cell lines Selected talk, RECOMB Satellite on Regulatory Genomics, San Francisco, CA | November 2012 |
| 0 | Computational Regulatory Genomics: Motifs, Networks, and Dynamics Invited talk, UIUC Institute for Genomic Biology, Urbana, IL | May 2011 |
| 0 | Regulatory motifs associated with TF binding and chromatin dynamics Selected talk, RECOMB Satellite on Regulatory Genomics, Cambridge, MA | December 2009 |
| 0 | Target identification in flies and mammals using comparative genomics Selected talk, Systems Biology: Global Regulation of Gene Expression, Cold Spring Harbor, NY | March 2008 |
| 0 | Systematic discovery and characterization of <i>Drosophila</i> miRNAs Selected talk RECOMB Satellite on Regulatory Genomics, Cambridge, MA | October 2007 |

Awards & Honors

- o 2005 National Science Foundation Graduate Research Fellowship Recipient
- o 2005 Hertz Fellowship Finalist
- $\circ~$ 2005 Outstanding Teaching Assistant, Department of Computer Science, UIUC
- o 2004 Excellent Teaching Assistant, Department of Computer Science, UIUC