Alejandro Ochoa

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https://ochoalab.github.io/

Education

2006

2008

2001

2013-2018

2016

PHD in Molecular Biology, Princeton University

Dissertation: Protein domain prediction using context statistics, the false discovery rate, and comparative genomics, with application to Plasmodium falciparum

Advisers: Mona Singh (Computer Science) and Manuel Llinás (Molecular Biology)

BS in Biology and Mathematics, Massachusetts Institute of Technology

Awards & Honors

NSF Graduate Research Fellowship. Computational Biology.

Ford Foundation Diversity Fellowship Predoc. Comp. (Declined, listed as Honorable Mention.)

MIT Department of Biology Merck Prize. Awarded to a graduating senior for outstanding research and academic performance in biophysics or bioinformatics.

Mexican Math Olympiad Gold Medal. Awarded to top 15 competitors nationally.

Employment

Assistant Professor, Duke Center for Statistical Genetics and Genomics, Department of Biostatistics and Bioinformatics, Duke University

Postdoctoral Research Associate at John D Storey's group, Lewis-Sigler Institute for Integrative Genomics, and Center for Statistics and Machine Learning, Princeton University

Publications

JOURNAL ARTICLES

Alejandro Ochoa, Mona Singh. Domain prediction with probabilistic directional context. *Bioinf* 33(16) 2471-2478.

Simon A Cobbold, Joana M Santos, Alejandro Ochoa, David H Perlman, Manuel Llinás. Proteomewide analysis reveals widespread lysine acetylation of major protein complexes in the malaria parasite. *Sci Rep* 2016;6:19722.

- Alejandro Ochoa, John D Storey, Manuel Llinás, Mona Singh. Beyond the *E*-value: stratified statistics for protein domain prediction. *PLoS Comput Biol* 11 e1004509.
- Moriah L Szpara, Derek Gatherer, Alejandro Ochoa, Benjamin Greenbaum, Aidan Dolan, Rory J Bowden, Lynn W Enquist, Matthieu Legendre, Andrew J Davison. Evolution and diversity in human herpes simplex virus genomes. *J Virol* 88:1209-1227.
- Alejandro Ochoa, Manuel Llinás, Mona Singh. Using context to improve protein domain identification. *BMC Bioinformatics* 12:90.
- Gevorg Grigoryan, Alejandro Ochoa, Amy E Keating. Computing van der Waals energies in the context of the rotamer approximation. *Proteins* 68 (4), 863-78.

MANUSCRIPTS IN SUBMISSION

- Alejandro Ochoa, John D Storey. New kinship and $F_{\rm ST}$ estimates reveal higher levels of differentiation in the global human population. Preprint: https://doi.org/10.1101/653279
- Alejandro Ochoa, John D Storey. F_{ST} and kinship for arbitrary population structures II: Method of moments estimators. Preprint: https://doi.org/10.1101/083923
- Alejandro Ochoa, John D Storey. F_{ST} and kinship for arbitrary population structures I: Generalized definitions. Preprint: https://doi.org/10.1101/083915

Distributed software

- human-differentiation-manuscript: Human differentiation analysis. R, bash, markdown.
 - https://github.com/StoreyLab/human-differentiation-manuscript.
- genio: Genetics Input/Output Functions. R.
 - Available on CRAN and https://github.com/OchoaLab/genio.
- simtrait: Simulate Complex Traits from Genotypes. R.
 - https://github.com/OchoaLab/simtrait.
- popkinsuppl: Supplement to "popkin" package. R.
 - https://github.com/OchoaLab/popkinsuppl.
- 2017-2019 popkin: Estimate Kinship and FST under Arbitrary Population Structure. R.
 - Available on CRAN and https://github.com/StoreyLab/popkin.
- bnpsd: Model and Simulate Admixed Populations. R.
 - Available on CRAN and https://github.com/StoreyLab/bnpsd.
- ²⁰¹⁴⁻²⁰¹⁶ dPUC2: Domain Prediction Using Context, Version 2. Perl.
 - https://github.com/alexviiia/dpuc2.
- 2014-2015 DomStratStats: Domain Stratified Statistics (q-values and local FDRs). Perl.
 - https://github.com/alexviiia/DomStratStats.
- 2014-2015 RandProt: High-order Markov random models for protein sequences. Perl.
 - https://github.com/alexviiia/RandProt.

Invited Talks

Computational Biology and Bioinformatics (CBB) seminar. French Family Science Center, Duke University, Durham, NC. 2019-02-04.

- 2019 4th Mexico Population Genomics Meeting. Amoxcalli Complex, Faculty of Sciences, University City, National Autonomous University of Mexico, Mexico City, DF, Mexico. 2019-01-11.
- Population Biology seminar, Department of Biology, Duke University, Durham, NC. 2018-09-27.
- Department of Biostatistics and Bioinformatics, Duke University, Durham, NC. 2018-02-21.
- Department of Biostatistics, Johns Hopkins University, Baltimore, MD. 2018-01-22.
- Department of Biology, University of Richmond, Richmond, VA. 2017-11-10.
- Princeton Research Day. Princeton University, Princeton, NJ. 2017-05-11.
- Department of Genetics, University of North Carolina, Chapel Hill, NC. 2017-02-13.
- New York Area Population Genomics Workshop 2016. Princeton University, Princeton, NJ. 2016-01-21.
- 2015 Probabilistic Modeling in Genomics conference. Cold Spring Harbor Laboratory, Cold Spring Harbor, NY. 2015-10-15.
- Telepresentation for Yun Song's group. UC Berkeley, Berkeley, CA. 2013-04-17.
- Biological sequence analysis and probabilistic models conference. HHMI Janelia Farm, Ashburn, VA. 2013-03-25.
- NCBI, NIH, Bethesda, MD. 2013-02-25.
- Recruiting conference. Department of Computer Science, Princeton University, Princeton, NJ. 2012-03-01.
- Rotary Club Paso del Norte, Ciudad Juarez, CHIH, Mexico. 2004-08-26.

Conference posters

- Alejandro Ochoa, John D Storey, Srikanth Gottipati, Shashank Rohatagi, Andrew Forbes, Deborah Profit, Raymond Sanchez, Margaretta Nyilas, William Carson. Mixed Effects Modeling of Placebo Response Across 8 Placebo-controlled Aripiprazole Trials Spanning 20 Years in Acutely Relapsed Schizophrenia Patients. Neuroscience Education Institute Psychopharmacology Congress. Broadmoor Convention Center, Colorado Springs, CO.
- Srikanth Gottipati, Alejandro Ochoa, Shashank Rohatagi, Andrew Forbes, Deborah Profit, Raymond Sanchez, Margaretta Nyilas, William Carson. Canonical Loadings of PANSS Subscales Show Differential Placebo and Aripiprazole Drug Responses in Schizophrenia Patients. Neuroscience Education Institute Psychopharmacology Congress. Broadmoor Convention Center, Colorado Springs, CO.
- Alejandro Ochoa, John D Storey. F_{ST} generalized for arbitrary population structures. John W. Tukey 100th Birthday conf. Center for Statistics and Machine Learning, Princeton, NJ.
- Alejandro Ochoa, John D Storey, Manuel Llinás, Mona Singh. Forget the *E*-value: family-based *q*-values for protein domain prediction, and empirical error detection. Biological sequence analysis and probabilistic models conf. HHMI Janelia Farm, Ashburn, VA.
- Alejandro Ochoa, Manuel Llinás, Mona Singh. Using context to predict protein domains across diverse organisms. Recomb Systems Biology conf. Columbia University, New York, NY.

Teaching

PRINCETON UNIVERSITY

2017	Intro to Genomics and Comp Bio. Quant Comp Bio, Comp Sci. (Co-Lecturer Fall 2017)
2016-2017	Discussion Leader. Summer Undergraduate Research Program (Summer 2016, 2017)
2014-2017	Statistical Treatment of Data, Mol Bio, Quant Comp Bio, (Workshop Fall 2014, 2015, 2016, 201

Intro to Genomics and Comp Mol Bio. Mol Bio, Comp Sci. (TA Fall 2011)

2008 Core Laboratory. Mol Bio. (TA Spring 2008)

Massachusetts Institute of Technology

²⁰⁰⁵ Calculus. OME Project Interphase (TA Summer 2005)

Multivar Calculus and Diff Eqs. OME Seminar XL (Tutor Fall 2003, Spring 2004)

2003-2004 Calculus. MITE2S Program (TA Summer 2003, 2004)

Mentoring

Yiqi Yao (Duke), Master's student, Ochoa lab. Theoretical and empirical evaluation of genetic association study approaches.

Sebastian Nasamu (MIT), summer student, Llinás lab. Cloned DNA of malaria protein domains, purified proteins, used protein-binding microarrays to test for DNA binding.

Jeremy Bigness (Princeton), graduate rotation, Singh lab. Built HMMs of ortholog proteins.

Neo Chung (Princeton), graduate rotation, Llinás lab. Wrote scripts to analyze sequencing data.

Research experience

Postdoctoral Research, Storey Lab, Princeton. Developed new definitions and tools to study arbitrary population structures. Also applied GWAS and mixed effects modeling to longitudinal psychiatric drug clinical trials.

Graduate Research Thesis, Singh Lab and Llinás Lab, Princeton. Developed probabilistic models and statistical methods that improve protein domain prediction. Studied the *Plasmodium falciparum* proteome, focusing on the AP2 transcription factors. Used experimental techniques (cloning, protein purification and protein-binding microarrays) to test predicted AP2 domains for DNA binding.

Graduate Rotation, Tavazoie Lab, Princeton. Cloned an RNA aptamer for a phage display assay.

Graduate Rotation, Singh Lab, Princeton. Analyzed protein interactions predicted from the presence of interacting domain pairs.

Graduate Rotation, Troyanskaya Lab, Princeton. Built and analyzed a potential "gold standard" of Gene Ontology terms integrated from predictions of multiple sources, in *Saccharomyces cerevisiae* and *Homo sapiens*.

Undergraduate Research Assistant, Keating Lab, MIT. Analyzed full-atom computational protein designs using modified van der Waals potentials.

Extracurricular University Activities

Latino Graduate Student Association, Princeton University. Board member: Technology Special-

ist 2007-2010 (handled website, email list, photos, calendar).

2002-2006 Association of Puerto Rican students, MIT. Regular member.

Outreach

Panelist at HISPA Latinos in College Conference. Princeton University.

Ivy-plus recruiting fair at University of Puerto Rico (4 campuses) for Princeton.

²⁰⁰⁸⁻²⁰¹¹ Science and Engineering Expo. HHMI and Princeton University.

Helped local high school students with college personal statements. PUPP, Princeton University.

Languages

2007

English. Native proficiency.

Spanish. Native proficiency.

French. Fluent reading and writing, conversational speaking.