Jordi Abante

Stanford University jabante@stanford.edu Stanford School of Medicine https://jordiabante.github.io Department of Biomedical Data Science Skype: jordi.abante1 1265 Welch Rd, Stanford CA 94305, USA Phone: +1 (979) 739-2083RESEARCH Machine Learning, Computational Genomics, Epigenetics. **INTERESTS** LANGUAGES Catalan (native), Spanish (native), English (proficient), French (basic). PROGRAMMING julia, python, R, bash, perl, C++, Matlab, IATFX. **EDUCATION** Stanford University 9/2021 - Present & TRAINING Stanford, California, U.S. • Postdoctoral research fellow at the Salzman lab • Department of Biomedical Data Science • Center for Computational, Evolutionary & Human Genomics Johns Hopkins University Baltimore, Maryland, U.S. • Ph.D., Electrical and Computer Engineering 8/2016 - 8/2021 • M.S.E., Applied Mathematics and Statistics 8/2016 - 12/2018 • Teaching Institute Certificate 6/2020 - 3/2021Texas A&M University College Station, TX, U.S. • M.S., Electrical and Computer Engineering 8/2014 - 5/2016 5/2015 - 6/2015 • Business Management Certificate Polytechnic University of Catalonia Barcelona, Catalonia, Spain • B.S., Industrial Engineering 9/2008 - 5/2014

• Major: Signal Processing

RESEARCH EXPERIENCE

Research Assistant

9/2016 - 6/2021

Whitaker Biomedical Engineering Institute (CIS) Johns Hopkins University, Baltimore MD, U.S.

- Computational biology
- Genomics & epigenetics data modeling & analysis
- Statistical inference & machine learning

Research Assistant

9/2014 - 1/2015, 6/2015 - 5/2016

Center for Bioinformatics and Genomic Systems Engineering Texas A&M University, College Station, TX, U.S.

- Computational biology
- Genome & transcriptome data modeling & analysis
- Statistical inference & machine learning

Team Captain & Member

8/2012 - 8/2014

ETSEIB Motorsport Formula Student Team Polytechnic University of Catalonia Barcelona, Catalonia, Spain

- Power electronics R&D
- Electric vehicle R&D

PEER REVIEWED JOURNAL ARTICLES

- **J. Abante**, S. Kambhampati, A.P. Feinberg, J. Goutsias. "Estimating DNA methylation potential energy landscapes from nanopore sequencing data". *Scientific Reports* (in press).
- M. Koldobskiy, G. Jenkinson, **J. Abante**, V.A. Rodriguez DiBlasi, W. Zhou, E. Pujadas, A. Idrizi, R. Tryggvadottir, C. Callahan, C. Bonifant, K. Rabin, P.A. Brown, H. Ji, J. Goutsias, and A.P. Feinberg.. "Converging genetic and epigenetic drivers of paediatric acute lymphoblastic leukaemia identified by an information-theoretic analysis". *Nature Biomedical Engineering* 5.4 (2021): 360-376.
- J. Abante, Y. Fang, A.P. Feinberg, J. Goutsias. "Detection of haplotype-dependent allele-specific DNA methylation in WGBS data". *Nature Communications* 11, 5238 (2020). Featured in Nature Communications Editors' Highlights.

- C. Qiu, H. Jin, I. Vvedenskaya, **J. Abante**, T. Zhao, I. Malik, S. Schwartz, P. Cui, P. Cabart, K. Hoo, R.P. Metz, C.D. Johnson, S. Sze, B.F. Pugh, B.E. Nickels, C.D. Kaplan. "Universal promoter scanning by Pol II during transcription initiation in Saccharomyces cerevisiae". *Genome Biology* (2020), 21:132.
- M. Koldobskiy, **J. Abante**, G. Jenkinson, E. Pujadas, A. Tetens, F. Zhao, R. Tryggvadottir, A. Idrizi, A. Reinisch, R. Majeti, J. Goutsias, and A. P. Feinberg. "A Dysregulated DNA Methylation Landscape Linked to Gene Expression in MLL-Rearranged AML". *Epigenetics* (2020), 1-18.
- M. P. Menden, D. Wang, M. J. Mason, B. Szalai, K. C. Bulusu, Y. Guan, J. Kang, M. Jeon, R. Wolfinger, T. Nguyen, M. Zaslavskiy, **J. Abante**, et al. "Community assessment to advance computational prediction of cancer drug combinations in a pharmacogenomic screen." *Nature Communications* 10, 2674 (2019).
- G. Jenkinson, **J. Abante**, M. Koldobskiy, A. P. Feinberg, J. Goutsias. "Ranking genomic features using an information-theoretic measure of epigenetic discordance". *BMC Bioinformatics* (2019), 20:175.
- G. Jenkinson, **J. Abante**, A. P. Feinberg, J. Goutsias. "An information-theoretic approach to the modeling and analysis of whole-genome bisulfite sequencing data". *BMC Bioinformatics* (2018), 19:87.
- **J. Abante**, N. Ghaffari, C.D. Johnson, A. Datta. "HiMMe: using genetic patterns as a proxy for genome assembly reliability assessment." *BMC Genomics* (2017), 18:694.

PREPRINT PAPERS

- **J. Abante**, S. Kambhampati, A.P. Feinberg, J. Goutsias. "Estimating DNA methylation potential energy landscapes from nanopore sequencing data". *bioRxiv* (2021), 431480.
- **J. Abante**, J. Goutsias. "CpelTdm.jl: a Julia package for targeted differential DNA methylation analysis". bioRxiv (2020), 343020.

C. Qiu, H. Jin, I. Vvedenskaya, **J. Abante**, T. Zhao, I. Malik, A.M. Visbisky, S.L. Schwartz, P. Cui, P. Čabart, K.H. Han, W.K.M. Lai, R.P. Metz, C.D. Johnson, S.H. Sze, B.F. Pugh, B.E. Nickels, C.D. Kaplan. "Promoter scanning during transcription initiation in Saccharomyces cerevisiae: Pol II in the "shooting gallery". *bioRxiv* (2019), 810127.

N. Ghaffari, **J. Abante**, R. Singh, P. D. Blood, L. Pipes, C. Mason, C. D. Johnson. "What are the most influencing factors in reconstructing a reliable transcriptome assembly?". *bioRxiv* (2017), 220269.

PEER REVIEWED CONFERENCE PROCEEDINGS

P. Blood, N. Ghaffari, A. S. Seetharam, L. Pipes, R. Singh, **J. Abante**, A. Severin, C. D. Johnson, C. Mason. "Fast, flexible, and free: enabling large-scale genome assembly and analysis with the Bridges supercomputer". *In Plant and Animal Genome XXVI Conference*, San Diego, CA, U.S., 2018.

N. Ghaffari, **J. Abante**, R. Singh, P. D. Blood, C. D. Johnson. "Computational considerations in transcriptome assemblies and their evaluation, using high quality human RNA-Seq data". *XSEDE16: Extreme Science and Engineering Discovery Environment 2016 Conference*, Miami, FL, U.S., 2016.

INVITED TALKS

On the estimation of epigenetic energy landscapes from nanopore sequencing data @ Machine Learning in Computational and Systems Biology, Intelligent Systems for Molecular Biology and European Conference on Computational Biology 2021, 7/2021.

An information-theoretic approach to allele-specific DNA methylation analysis

- @ Salzman Lab, Stanford University, 11/2020
- @ Yosef Lab, Berkeley University, 11/2020
- @ Gerstein Lab, Yale University, 11/2020
- @ Knowles Lab, Columbia University, 10/2020
- @ Marks Lab, Harvard University, 10/2020

Statistical modeling and analysis of allele-specific DNA methylation at the haplotype level @ Electrical & Computer Engineering Departmental Seminars, Johns Hopkins University, 10/2019.

POSTER SESSIONS

- J. Abante, S. Kambhampati, A.F. Feinberg, J. Goutsias. "On the estimation of epigenetic energy landscapes from nanopore sequencing data". *Machine Learning in Computational and Systems Biology, Intelligent Systems for Molecular Biology and European Conference on Computational Biology* 2021, July 2021.
- **J. Abante**, N. Ghaffari, C.D. Johnson, A. Datta. "Using hidden Markov models to analyze next-generation sequencing data". *ENG-LIFE 2016: At the Interface of Engineering and Life Sciences*, College Station, TX, U.S., 2016.

FELLOWSHIPS

Stanford CEHG Postdoctoral Fellowship

10/2021 - 9/2022

Center for Computational, Evolutionary and Human Genomics (CEHG) Stanford University, Stanford, California, U.S.

• Provided funds to launch my first year of postdoctoral research

"la Caixa" Fellowship

8/2016 - 8/2017

"la Caixa" Foundation, Barcelona, Catalonia, Spain

• Provided funds for my first two years in my Ph.D. program

TEACHING

Course Assistant & Guest Lecturer

Fall 2018, 2019, 2020

EN.520.622. Principles of Complex Networked Systems Johns Hopkins University, Baltimore, MD, U.S.

- Graded assignments and exams
- Provided students with one-on-one tutoring and out of class assistance

Teaching Assistant & Guest Lecturer

Spring 2015

ENGR-112 Foundations of Engineering II

Texas A&M University, College Station, TX, U.S.

• Tutored students and managed a team of grading assistants

MENTORING

Stanford University, Stanford, CA, U.S.

• Aasavari Kakne, MS student in ICME	Fall 2021
• Angelika Hirsch, Ph.D Student in Biophysics	Fall 2021
• Ayushi Tandel, MS Student in Computer Science	Fall 2021
• Nathaniel Chien, MS Student in Computer Science	Fall 2021

Johns Hopkins University, Baltimore, MD, U.S.

• Sandeep Kambhampati, Under. Student in BME 2020-2021

AWARDS & HONORS	Best Thesis Award Polytechnic University of Catalonia Barcelona, Catalonia, Spain	5/2015
	Golden Key Honor Society Texas A&M University, College Station, Texas, U.S.	5/2016
	Phi Kappa Phi Texas A&M University, College Station, Texas, U.S.	4/2016
	IEEE-Eta Kappa Nu Honor Society Texas A&M University, College Station, Texas, U.S.	1/2016
INDUSTRY EXPERIENCE	R&D Engineer 5/2016 • Sensory Value Sant Cugat del Vallès, Catalonia, Spain • Machine learning applied to market research	- 8/2016
	~ .	- 5/2014
UNIVERSITY SERVICE	Biomedical Data Science JEDI committee 20 Department of Biomedical Data Science Stanford University, Stanford, CA, U.S. • Member of justice, equity, diversity, and inclusion (JEDI) commit	021-2022 etee.
	ECE Department Head Search Department of Electrical & Computer Engineering (ECE) Johns Hopkins University, Baltimore, Maryland, U.S. • Collaborated in the hiring process of 8 candidates	ing 2021
COMMUNITY SERVICE	Voluntariat per la llengua Fall 2020, Spr. Baltimore, Maryland, U.S. • Language partner for people interested in learning Catalan	ing 2021
	 Our Daily Bread Volunteer Baltimore, Maryland, U.S. Cooked food for families in need during COVID 19 pandemic 	2020

Barclay Hopkins STEM Partnership

2017

Johns Hopkins University, Baltimore, Maryland, U.S.

• Worked with elementary school students to engage them in STEM

The Big Event

2015-2016

Texas A&M University, College Station, Texas, U.S.

• Provided yard work and window washing for elderly residents

Grup de voluntariat ANTAR

2006-2008

Àgora International School, Sant Cugat del Vallès, Catalonia, Spain

• Organized fundraising activities and food drives for people in need

REFERENCES Available upon request.

PROFESSIONAL International Society for Computational Biology (ISCB). AFFILIATIONS