Jordi Abante Llenas

Universitat de Barcelona jordi.abante@ub.edu

Department of Biomedical Sciences https://jordiabante.github.io

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RESEARCH Machine Learning, Computational Genomics, Epigenetics. INTERESTS

LANGUAGES Catalan (native), Spanish (native), English (proficient), French (basic).

PROGRAMMING julia, python, R, bash, perl, C++, Matlab, LATEX.

EDUCATION Universitat de Barcelona

1/2023 - Present

& TRAINING Barcelona, Catalonia, Spain

- Postdoctoral research fellow
- Department of Biomedical Sciences
- Department of Mathematics and Computer Science

Stanford University

9/2021 - 1/2023

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
\bullet M.S.E., Applied Mathematics and Statistics	8/2016 - 12/2018
• Teaching Institute Certificate	6/2020 - 3/2021

Texas A&M University

College Station, TX, U.S.

• M.S., Electrical and Computer Engineering	8/2014 - 5/2016
• Business Management Certificate	5/2015 - 6/2015

Polytechnic University of Catalonia

Barcelona, Catalonia, Spain

• Llicenciatura, Industrial Engineering

9/2008 - 5/2014

• Major: Power Electronics

RESEARCH EXPERIENCE

Research Assistant

9/2016 - 6/2021

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

- 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.

- 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**, P. L. Wang, J. Salzman. "DIVE: a reference-free statistical approach to diversity-generating and mobile genetic element discovery". *Nature Methods* (under review).
- Y. Fang, Z. Ji[†], W. Zhou[†], **J. Abante**[†], M.A. Koldobskiy, H. Ji, and A.P. Feinberg. "DNA methylation entropy is associated with DNA sequence features and developmental epigenetic divergence". *Nucleic Acids Research* (2023), gkad050. [†] Equal contribution.
- **J. Abante**, S. Kambhampati, A.P. Feinberg, J. Goutsias. "Estimating DNA methylation potential energy landscapes from nanopore sequencing data". *Scientific Reports* (2021), 11-21619.

- M.A. Koldobskiy, G. Jenkinson, **J. Abante**, V.A. Rodriguez DiBlasi, W. Zhou, E. Pujadas, ..., 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). **Nature Comms 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

- Y. Fang, Z. Ji, W. Zhou, **J. Abante**, M. Koldobskiy, H. Ji, A. P. Feinberg. (2022). "DNA methylation entropy is associated with DNA sequence features and developmental epigenetic divergence". *bioRxiv*, 2022-12.
- **J. Abante**, P. L. Wang, J. Salzman. "DIVE: a reference-free statistical approach to diversity-generating and mobile genetic element discovery". *bioRxiv* (2022), 495703.
- **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

Nanopore-derived epigenetic landscapes and reference-free analysis of microbial evolution @ Biomedical Sciences, Universitat de Barcelona, 9/2022.

On the estimation of epigenetic energy landscapes from nanopore sequencing data @ MLCSB + ISMB + ECCB 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 Department, Johns Hopkins University, 10/2019.

POSTER SESSIONS

- **J. Abante**, P. Wang, J. Salzman. "DIVE: a reference free statistical approach to diversity-generating & mobile genetic element discovery". *European Conference on Computational Biology* 2022, Sitges, Barcelona, Spain, September 2022.
- 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.

RESEARCH PROJECTS

• In vivo reprogramming to rescue alterations in Huntington's disease.

PIs: J. M. Canals Coll, P. I. Radeva.

Funding agency: "la Caixa" Foundation (9/19-8/23).

• A Mesoscopic Bottom-up Approach for Understanding and Modulating the Physical Epigenome.

PIs: A. P. Feinberg, M. D. Fallin, J. Goutsias, K. D. Hansen, J. Rashelle, S. Szalay, and W. Timp.

Funding agency: National Science Foundation (9/19-8/23).

• Integration of Genomics and the Environment.

PIs: A. P. Feinberg, Goutsias, T. Ha, and S. D. Taverna.

Funding agency: National Institutes of Health (9/16-8/21).

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 - 8/2018

"la Caixa" Foundation, Barcelona, Catalonia, Spain

• Funded my Ph.D. at Johns Hopkins University

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

THESIS SUPERVISION

Universitat de Barcelona, Barcelona, Catalonia, Spain

• U. Muhammad, PhD - UB Math. & C.S.	Spring 2023
• J. Rosell, MS - UB Biomedical Engineering	Spring 2023
• R. Smeriglio, MS - UB Biomedical Engineering	Spring 2023
• A. Melendez, BS - UPC Data Science	Spring 2023

MENTORING

Stanford University, Stanford, CA, U.S.

• A. Kakne, MS - Stanford ICME	Fall 2021
• A. Hirsch, PhD - Stanford Biophysics	Fall 2021
• A. Tandel, MS - Stanford Computer Science	Fall 2021
• N. Chien, MS - Stanford Computer Science	Fall 2021

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

• S. Kambhampati, BS - Johns Hopkins BME 2020-2021

AWARDS & HONORS	Best Thesis Award 5/2015 Polytechnic University of Catalonia Barcelona, Catalonia, Spain	
	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
REVIEWER	Intelligent Systems in Molecular Biology (ISMB) 2022	2/2022
INDUSTRY EXPERIENCE	 R&D Engineer Sensory Value Sant Cugat del Vallès, Catalonia, Spain Machine learning applied to market research 	5/2016 - 8/2016
	 R&D Engineer Cinergia, Control Intel·ligent de l'Energia Barcelona, Catalonia, Spain Power electronics R&D for EV applications 	3/2013 - 5/2014
UNIVERSITY SERVICE	 Biomedical Data Science JEDI committee Department of Biomedical Data Science Stanford University, Stanford, CA, U.S. Member of justice, equity, diversity, and inclusion () 	2021-2022 JEDI) committee
	 ECE Department Head Search Department of Electrical & Computer Engineering (Electrical & Hopkins University, Baltimore, Maryland, U.S. Collaborated in the hiring process of department head 	,

COMMUNITY SERVICE

Voluntariat per la llengua

Fall 2020, Spring 2021

Baltimore, Maryland, U.S.

• Language partner for people interested in learning Catalan

Our Daily Bread Volunteer

2020

Baltimore, Maryland, U.S.

• Cooked food for families in need during COVID 19 pandemic

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