# **David Angeles-Albores**

**L** +1 607 379 4030

☑ davidaalbores@gmail.com

**♥** @DavidAngeles13

dangeles.github.io

https://github.com/dangeles

47 Garden St, Apt 4, Boston, MA, 02114

## **Education**

2013–2018 **Ph.D.**, Biochemistry and Molecular Biophysics,

California Institute of Technology Defense Date: 18 September, 2018

Degree Awarded: 31 October, 2018

2009-2013 B.A., cum laude, Biology

**Cornell University** 

## **Appointments**

11/2019 - Computational Biologist II, eGenesis

One of two founding members of the computational biology unit. Introduced ATAC-seq, scRNA-seq and scATAC-seq to eGenesis.

I collaborated on the development of kidney dissociation protocols and used these protocols to create an atlas of the pig kidney cortex using scRNA-seq.

I used the above methods to generate a compendium of promoters that stably express genes ubiquitously or with high tissue-specificy at desired levels with low burstiness. *Patent pending*.

7/2020 - **Visiting Scholar**, Laboratory of Ilya Ruvinsky,

Northwestern University

I collaborate with the Ruvinsky lab on RNA-seq analysis of C. elegans biology.

01/2019–11/2019 **Postdoctoral Associate**, Laboratory of Eric J. Alm,

Massachusetts Institute of Technology

I developed methods for barcodeless, highly multiplexed RNA-seq of multiple bacterial species using a single library preparation protocol.

My main project centered around recreating the vaginal microbiome *in vitro* and dissecting the causal interactions between microbial species using a mixture of computational and experimental methods. I hoped to apply the algorithms I developed at Caltech (see below) to develop active learning models that combined iterative experimental minibatches with continuous computational analysis.

I terminated my postdoctoral research prematurely due to a major personal loss, unrelated to my work at MIT.

11/2018-01/2019

**Postdoctoral Fellow**, Labs of Paul W. Sternberg and Matt Thomson, California Institute of Technology

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I used this period immediately following my defense to finish projects with my Ph.D. advisor, Paul Sternberg, and a member of my committee, Matt Thomson, before moving on to my postdoctoral position at MIT in January, 2019. I developed active learning algorithms for analysis of CRISPR screens in mammalian cells, using transcriptomes as phenotypes.

## **Research Publications**

† denotes equal contributions.

### **Journal Articles**

- Basta, D. W., Angeles-Albores, D., Spero, M. A., Ciemniecki, J. A., & Newman, D. K. (2020). Heat-shock proteases promote survival of Pseudomonas aeruginosa during growth arrest. *Proceedings of the National Academy of Sciences of the United States of America*. doi:10.1073/pnas.1912082117
- Duncan, E. M., Nowotarski, S. H., Guerrero-Hernández, C., Ross, E. J., D'Orazio, J. A., Clubes de Ciencia México, W. f. D. B., ... Alvarado, A. S. (2020). A new species of planarian flatworm from mexico: Girardia guanajuatiensis. *bioRxiv*. doi:10.1101/2020.07.01.183442. eprint:
  - https://www.biorxiv.org/content/early/2020/07/02/2020.07.01.183442.full.pdf
- Angeles-Albores, D., & Sternberg, P. W. (2018). Using Transcriptomes as Mutant Phenotypes Reveals Functional Regions of a Mediator Subunit in *Caenorhabditis elegans*. *Genetics*, genetics.301133.2018. doi:10.1534/genetics.118.301133
- <sup>†</sup>Angeles-Albores, D., <sup>†</sup>Puckett Robinson, Williams, B. A., Wold, B. J., & Sternberg, P. W. (2018). Reconstructing a metazoan genetic pathway with transcriptome-wide epistasis measurements. *Proceedings of the National Academy of Sciences*, 201712387. doi:10.1073/pnas.1712387115
- <sup>†</sup>**Angeles-Albores**, **D.**, <sup>†</sup>Leighton, D. H. W., Tsou, T., Khaw, T. H., Antoshechkin, I., & Sternberg, P. W. (2017). The *Caenorhabditis elegans* Female-Like State: Decoupling the Transcriptomic Effects of Aging and Sperm Status. *G3* (*Bethesda*, *Md.*) 7(9), 2969–2977. doi:10.1534/g3.117.300080
- Angeles-Albores, D., N. Lee, R. Y., Chan, J., & Sternberg, P. W. (2016). Tissue enrichment analysis for *C. elegans* genomics. *BMC Bioinformatics*, 17(1), 366. doi:10.1186/s12859-016-1229-9
- Albores-Saavedra, J., Chable-Montero, F., **Angeles-Albores**, **D.**, Schwartz, A., Klimstra, D. S., & Henson, D. E. (2011). Early Gallbladder Carcinoma. *American Journal of Clinical Pathology*, 135(4), 637–642. doi:10.1309/AJCPFRKCFEDLV03Y
- Albores-Saavedra, J., Schwartz, A. M., Henson, D. E., Kostun, L., Hart, A., **Angeles-Albores**, **D.**, & Chablé-Montero, F. (2011). Cutaneous angiosarcoma. Analysis of 434 cases from the surveillance, epidemiology, and end results program, 1973-2007. *Annals of Diagnostic Pathology*, 15(2), 93-97. doi:10.1016/j.anndiagpath.2010.07.012

#### **Patents**

eGenesis. (In prep). Cells, tissues, organs, and/or animals having one or more modified genes for enhanced xenograft survival and/or tolerance.

### $\mu$ Publications

Angeles-Albores, D., N. Lee, R. Y., Chan, J., & Sternberg, P. W. (2018). Two new functions in the WormBase Enrichment Suite. *microPublication Biology*. doi:10.17912/W25Q2N

## Scientific Talks

2019	Probabilistic 1	Modeling in	Genomics

Genetics is an active learning algorithm for causal reconstruction of biological networks

### Hanna H Gray Semifinalist Symposium

Phenotypes, epistasis, and probability theory

ASBMB Special Symposium: Evolution and Core Processes in Gene Expression

Transcriptomes as phenotypes

2018 Bay Area Worm Meeting

Allelic series analyses using transcriptomic phenotypes

2017 21st C. elegans International Meeting, WormBase: Tools, Content and Community Annotation, Workshop

Gene Set Analysis tool for Gene Ontology (GO), Phenotype, and Tissue Enrichment

Annual Departmental Retreat, California Institute of Technology

Genome-wide, unbiased experimental genetics

Biochemistry and Molecular Biophysics Seminar Series, California Institute of Technology

Transcriptomic Genetics: A new way to use RNA-sequencing data

**Center for Environmental Microbial Interactions**, California Institute of Technology Genome-wide unbiased experimental genetics

Annual Biochemistry and Molecular Biophysics Program Retreat, California Institute of Technology

Reconstruction of a genetic pathway using whole-organism expression profiles

Graduate Biology Seminar, California Institute of Technology

Transcriptome-wide epistasis in mRNA expression profiles

### **Awards**

2020	eGenesis Leadership Award
2019	HHMI Hanna Gray Fellow Finalist
2015	Florence C. Rose and S. Meryl Rose Endowed Scholarship
2014	Amgen Graduate Student Fellowship
2012	EXROP Capstone Award
2011	EXROP HHMI Summer Fellowship

## **Scientific Courses**

2015 Embryology, Marine Biological Laboratory at Woods Hole

# **Teaching and Mentoring Experience**

### **Teaching**

2017-2018	Systems Genetics, Teaching Assistant, California Institute of Technology
2016	Introduction to Biology, Teaching Assistant, California Institute of Technology
2015	Order of Magnitude Biology, Teaching Assistant, California Institute of Technology

# Teaching and Mentoring Experience (continued)

Advanced Experimental Methods in Bioorganic Chemistry, Teaching Assistant, California Institute of Technology

# Mentoring

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2016–2019	<b>Kyung Hoi Min</b> , Caltech undergraduate, experimental and computational student, <i>California Institute of Technology</i>
Summer 2018	<b>Vladimir Molchanov</b> , Saint Petersburg Bioinformatics Institute undergraduate, experimental student, <i>California Institute of Technology</i>
Summer 2015	<b>Tiffany Tsou</b> , UCSB undergraduate, experimental student, <i>California Institute of Technology</i>
2014-2015	Isabelle Phinney, Polytechnic School, computational student, California Institute of Technology

# **Outreach**

2020	Invited speaker, Clubeando en casa, el Podcast, Clubes de Ciencia México
2017	Speaker, What is Clubes de Ciencias Mexico?, SACNAS
2016	Co-instructor, De Planarias y Derivas, Clubes de Ciencia México, Guanajuato
	Co-instructor, De Genes y Animales, Clubes de Ciencia México, Ensenada
2015	Student selection committee, Clubes de Ciencia México
2014	Guest instructor, Biología a través de los números, Clubes de Ciencia México, Ensenada

# **Scientific Societies**

2014-Present Genetics Society of America

2015–Present Society for Developmental Biology

## References

## Toby Bloom, Ph.D.

VP of Information Technology, eGenesis, 300 Technology Square Suite 301, Cambridge, MA 02139 ☑ tbloom@alum.mit.edu

### Professor Eric J Alm

Massachusetts Institute of Technology, 300 Technology Square
Suite 301, Cambridge, MA 02139

☑ burnss@mit.edu

### Professor Dianne K. Newman

California Institute of Technology, MC 147-75, Pasadena CA, 91125 ☑ dkn@caltech.edu

## **Assistant Research Professor**

Cornell University, Biotechnology 351, Cornell University, Ithaca,

Erich M. Schwarz

NY 14853-2703

ems394@cornell.edu

### Ranjith Anand, Ph.D.

Associate Director, Genome Editing, eGenesis, NE47-379, Cambridge, MA 02139

☑ rapanand@gmail.com

## Professor Paul W. Sternberg

California Institute of Technology, MC 156–29, Pasadena, CA 91125 

☑ pws@caltech.edu

### **Assistant Professor Matthew Thomson**

California Institute of Technology, 216–76, Pasadena CA, 91125

mthomson@caltech.edu