Ilias Tagkopoulos

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Education	 Princeton University, Princeton, NJ. Postdoctoral Research Fellow, Lewis-Sigler Institute, 2009 Ph.D. in Electrical Engineering, October 2008. M.A. in Electrical Engineering, May 2005. PEI-STEP Certificate in Environmental and Public Policy, May 2009. Advisors: Saeed Tavazoie (MolBio) and Sun-Yuan Kung (ECE)
	 Columbia University, New York, NY. M.Sc. in Electrical Engineering, February 2003. Concentration: Microelectronics & Wireless Networks. Advisors: Charles Zuckowski (EE) and Dimitrios Anastassiou (EE)
	◊ University of Patras, Patras, Greece. Diploma in Electrical and Computer Engineering, June 2001.
WORK EXPERIENCE	 ◇ Associate Professor (with tenure) University of California, Davis Department of Computer Science and UC Davis Genome Center. (2015 - now)
	 Advisor MARS Inc., McLean, VA (2015 - now) Advisor to the Chief Science Officer of MARS Inc. Member of the Computational Science Advisory Board.
	 ◇ Assistant Professor University of California, Davis (2009 - 2015) Department of Computer Science and UC Davis Genome Center.
	◇ Relationship Manager, DIT - Sales & Research Analytics, Credit Suisse (2008 - 2009) Liaison between modeling, programming and business teams. Supervised the development of quantitative financial models and solutions within the investment bank. Member of the 2008 Quantitative Associate class and the Fixed Income Derivatives LOCUS team.
	 ◇ Research Assistant, Lewis-Sigler Institute for Integrative Genomics (2003 - 2008) Developed a multi-scale microbial evolution simulator and machine learning algorithms for biological data integration.
	 ◇ Research Assistant, Columbia Integrated Systems Laboratory (CISL). (2001 - 2003) Designed and fabricated a biomimetic mixed signal VLSI chip.
Teaching experience	 ECS 188: Ethics in the Era of Technology UC Davis, 2012-2015, Participation: 24-31, Evaluation: 4.9/5.0 ECS 124: Theory and Practice of Bioinformatics UC Davis, 2010-2015, Participation: 46-50, Evaluation: 4.7/5.0 ECS 289K: Computational Systems and Synthetic Biology
	UC Davis, $2009-2013$ Participation: 5-21, Evaluation: $9.0/10.0$

♦ ECS 171: Machine Learning

UC Davis, 2014-2015, Participation: 80-125, Evaluation: 4.2/5.0

UPCOMING \diamond M. Kim, **I. Tagkopoulos**, "DeepPep: Deep proteome reconstruction from short peptide PUBLICATIONS profiles", submitted, 2016

- ◊ X. Wang, V. Zorraquino-Salvo, M. Kim, A. Tsoukalas, I. Tagkopoulos, "Predicting evolution from genome-scale mutation data", submitted, 2016
- M. Kim, V. Zorraquino-Salvo, N. Rai, I. Tagkopoulos, "Ecomics: A multi-omics data resource for *Escherichia coli*", submitted, 2016
- ◊ V. Zorraquino-Salvo, M. Kim, N. Rai, I. Tagkopoulos, "Cross-stress protection and the dynamics of complex behaviors under adaptation in combination of stresses", under review, bioRxiv, doi: http://dx.doi.org/10.1101/010595, 2016
- ◊ M. Meisner, J. Rosenheim, I. Tagkopoulos, "A data-driven, machine learning framework for optimal pest management in cotton", under revision, 2016

JOURNAL \diamond L. Huynh, **I. Tagkopoulos**. "Consensus parameter estimation for synthetic circuit design", PUBLICATIONS accepted, ACS Synthetic Biology, 2016

- N. Rai, A. Ferreiro, A. Neckelmann, A. Soon, A. Yao, J. Siegel, M.T. Facciotti, I. Tagkopou- los, "RiboTALE: a versatile and modular tool for the multi-dimensional control of gene expression", 5, Scientific Reports, 2015
- L. Huynh, I. Tagkopoulos. "Fast and Accurate Circuit Design Automation through Bounded Search and Hierarchical Model Switching", Apr 28, 25916918, ACS Synthetic Bi-ology, 2015
- M. Kim, V. Zorraquino-Salvo, I. Tagkopoulos, "Prediction of phenotypic characteristics from large-scale gene expression profiles", 11(3): e1004127. doi:10.1371/journal.pcbi.1004127, PLoS Comput Biol, 2015
- Taylor-Teeples M, Lin Lc, Trabucco G, de Lucas M, Turco G, Doherty C, Toal T, Gaudinier A, Young NF, Xiong G, Corwin J, Tsoukalas A, Pauly M, Kliebenstein DJ, I. Tagkopoulos, Breton G, Ahnert S, Kay SA, Brady SM, Hazen SP. "Environmental, developmental and genotype-dependent regulation of Xylem cell specification and secondary cell wall biosynthesis in *Arabidopsis thaliana*", 517, 571575, *Nature*, 2015
- ◊ A. Tsoukalas, T. Albertson, I. Tagkopoulos, "From data to optimal decision making: A data-driven, probabilistic machine learning approach to decision support for patients with Sepsis", 3(1), Jour. Med. Int. Res., 2015
- Hnin-Hnin Ma, Nasos Tsoukalas, John Rutledge, I. Tagkopoulos. "A systems biology analysis of brain microvascular endothelial cell lipotoxicity", BMC Systems Biology, 8:80, 2014
- J. Carrera, R. Estrela, J. Luo, N. Rai, A. Tsoukalas, I. Tagkopoulos. "An integrative, genome-scale model reveals the phenotypic landscape of *Escherichia coli*", 10(7):735, *Molec- ular Systems Biology*, 2014
- ◊ L. Huynh, I. Tagkopoulos. "Optimal part and module selection for synthetic gene circuit design automation", 3(8):556, doi: 10.1021/sb400139h, ACS Synthetic Biology, 2014
- E. Gultepe, J. Green, H. Nguyen, J. Adams, T. Albertson, I. Tagkopoulos, "From vital signs to clinical outcomes for patients with sepsis: A machine learning basis for a clinical decision support system", 21:315-325, doi: 10.1136/amiajnl-2013-001815, Journal of the American Medical Informatics Association (JAMIA), 2014

- A. Pavlogiannis, V. Mozhayskiy, I. Tagkopoulos, "A flood-based information flow analysis and network minimization method for bacterial systems", 14:137 DOI:10.1186/1471-2105-14-137, BMC Bioinformatics, 2013
- ◊ M. Dragosits, V. Mozhayskiy, S. Quinones-Soto, I. Tagkopoulos, "Evolutionary potential, cross-stress dependencies, and the genetic basis of acquired stress resistance in *E. coli*", doi:10.1038/msb.2012.76, 9:643, *Molecular Systems Biology*, 2013
- A. Yao, T. Fenton, K. Owsley, P. Seitzer, D. Larsen, H. Lam, J. Lau, A. Nair, J. Tantiongloc, I. Tagkopoulos, M. Facciotti, "Promoter activity arising from the fusion of standard BioBrick parts", 2(2), pp 111120, DOI: 10.1021/sb300114d, ACS Synthetic Biology, 2013
- L. Huynh, M. Köppe, I. Tagkopoulos, "SBROME: A scalable optimization and module matching framework for automated biosystem design", DOI: 10.1021/sb300095m2, pp 263-273, ACS Synthetic Biology, 2013
- V. Mozhayskiy, I. Tagkopoulos, "Microbial evolution in vivo and in silico: methods and applications", DOI:10.1039/C2IB20095C, 5(2):26277, Integrative Biology, 2013
- ◊ I. Tagkopoulos, "Microbial factories under control: Auto-regulatory control through engineered stress-induced feedback", 4:1, 1-4, *Bioengineered*, 2013
- Y. Liang, H. Wu, R. Lei, RA. Chong, Y. Wei, X. Lu, I. Tagkopoulos, SY. Kung, Q. Yang, G. Hu, Y. Kang, "Transcriptional Network Analysis Identififies BACH1 as a Master Regula- tor of Breast Cancer Bone Metastasis", 287(40):33533-44, *Journal of Biological Chemistry*, 2012
 2012
- ◊ L. Huynh, J. Kececioglu, M. Köppe, I. Tagkopoulos, "Automated Design of Synthetic Gene Circuits through Linear Approximation and Mixed Integer Optimization", 7(4):e35529, *PLoS ONE*, 2012
- M. Dragosits, D. Nicklas, I. Tagkopoulos, "A synthetic biology approach to self-regulatory recombinant protein production in *Escherichia coli*", 6:2, *Journal of Biological Engineering*, 2012
- ◊ V.Mozhayskiy, I. Tagkopoulos, "Guided evolution of *in silico* microbial populations in complex environments accelerates evolutionary rates through a step-wise adaptation", 13:S10, *BMC Bioinformatics*, 2012
- ◊ V.Mozhayskiy, I. Tagkopoulos, "Horizontal gene transfer dynamics and distribution of fitness effects during microbial In silico Evolution", 13:S13, BMC Bioinformatics, 2012
- I. Tagkopoulos, Y.Liu, S. Tavazoie, "Predictive Behavior Within Microbial Genetic Networks", Science, 320:1313-7, 2008
- S.Y.Kung, M.W. Mak, and I. Tagkopoulos, "Symmetric and Asymmetric Multi-modality Biclustering Analysis for Microarray Data Matrix", *Journal of Bioinformatics and Computational Biology*, vol 4(2), pp. 275-298, 2006

 PEER ◇ L.Huynh, N. Rai, I. Tagkopoulos, "Parameter inference for gene circuit models", Pro

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 ceedings of the 7th International Workshop on Bio-design Automation, IWBDA'15, Seattle,

 2015

- PUBLICATIONS L.Huynh, M. Kim, I. Tagkopoulos, "Integration of circuit design automation and genomescale modeling", Proceedings of the 6th International Workshop on Bio-design Automation, IWBDA'14, Boston, 2014
 - ◊ L.Huynh, I. Tagkopoulos, "Optimizing module matching for synthetic gene circuit design automation", Proceedings of the 5th International Workshop on Bio-design Automation, IWBDA'13, London, 2013
 - ◊ E. Gultepe, Hien Nguyen, Tim Albertson, I. Tagkopoulos, "A Bayesian network for early diagnosis of sepsis patients: a basis for a clinical decision support system", 2nd IEEE International Conference on Computational Advances in Bio and Medical Sciences (ICCABS), Las Vegas, NV, pp.1-5, 23-25, 2012

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- L. Huynh, I. Tagkopoulos, "A robust, library-based, optimization-driven method for automatic gene circuit design", 2nd IEEE International Conference on Computational Advances in Bio and Medical Sciences (ICCABS), Las Vegas, NV, pp.1-6, 24-26, 2012
- ◊ L.Huynh, J.Kececioglu, I. Tagkopoulos, "Scaling responsibly, Towards a reusable, modular, automated gene circuit design", Proceedings of the 4rth International Workshop on Bio-design Automation, IWBDA'12, San Francisco, 2012
- V.Mozhayskiy, R. Miller, KL. Ma, I. Tagkopoulos, "A Scalable Multi-scale Framework for Parallel Simulation and Visualization of Microbial Evolution", TeraGrid'11; Salt Lake City, Utah, 2011, DOI:10.1145/2016741.2016749 (Best Paper Award)
- ◊ V.Mozhayskiy, I. Tagkopoulos, "In silico Evolution of Multi-scale Microbial Systems in the Presence of Mobile Genetic Elements and Horizontal Gene Transfer", ISBRA'11, Lecture Notes in Bioinformatics, LNBI 6674, pp.262-273, Springer, 2011
- ◊ L.Huynh, J.Kececioglu, I. Tagkopoulos, "Automated Design of Synthetic Gene Circuits through Linear Approximation and Mixed Integer Optimization", Proceedings of the 3rd International Workshop on Bio-design Automation, IWBDA'11, San Diego, 2011.
- I. Tagkopoulos, D. Serpanos, "Gene Classification and Regulatory Prediction Based on Transcriptional Modeling." Proceedings of the IEEE Symposium on Signal Processing and Information Technology, ISSPIT'05, pp. 29-34, Proceedings of the Firth IEEE International Symposium on Signal Processing and Information Technology, Athens, Greece, 2005.
- ◊ I. Tagkopoulos, "A Transcriptional Approach to Gene Clustering", CIBCB'05, Proceedings of the IEEE Symposium on Computational Intelligence in Bioinformatics and Computational Biology, pp.1-7, San Diego, California, 2005. (student paper award)
- I. Tagkopoulos, N. Slavov, S.Y. Kung, "Multi-class Biclustering and Classification Based on Modeling of Gene Regulatory Networks", BIBE'05, Proceedings of the 5th IEEE Symposium on Bioengineering and Bioinformatics, pp. 89-96, Minneapolis, Minnesota, 2005.
- S.Y.Kung, M.W. Mak, and I. Tagkopoulos, "Multi-Metric and Multi-Substructure Biclustering Analysis for Gene Expression Data", CSB'05, Proceedings of the IEEE Computational Systems Bioinformatics Conference, pp.123-134, Stanford, California, 2005.
- I. Tagkopoulos, C.Zukowski, G.Cavelier, D.Anastassiou, "A Custom FPGA for the Simulation of Gene Regulatory Networks", GLSVLSI'03, Proceedings of the 13th ACM Great Lakes Symposium on VLSI, pp. 135-141, Washington D.C., 2003.
- N.D.Zervas, I. Tagkopoulos, V. Spiliotopoulos, D.Soudris, C.E.Goutis, "Comparison of DWT Scheduling Algorithms Alternatives on Programmable Platforms", ISCAS'01, Pro- ceedings of the International Symposium on Circuits and Systems, , pp. 761-764, vol. 2, Sidney, Australia, 2001.
- ABSTRACTS & M. Kim, N. Rai, V. Zorraquinno, L. Huynh, I. Tagkopoulos, "Deep learning and predictive models of a multi-omics dataset for *Escherichia coli*", Oral presentation, 3rd Conference on Quantitative Biology, q-bio'15, Maui, Hawaii, 2015
 - M. Kim, I. Tagkopoulos, "A graph-based integrative learning framework for whole-cell modeling", NetSci'14, Berkeley, 2014
 - ◊ J. Carrera, L. Huynh, I. Tagkopoulos, "A automated synthetic gene circuit design tool based on mathematical optimization", Poster, 1st Conference on Quantitative Biology, qbio'13, Honolulu, Hawaii, 2013
 - ◊ V. Mozhayskiy, M. Dragosits, I. Tagkopoulos, "Guided step-wise adaptation of microbial populations insilico and in vivo". 7th Annual Systems-to-Synthesis Symposium, San Diego, CA, 2012

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- ◊ I. Tagkopoulos, "Self-regulatory circuits for recombinant protein production", Poster, 11th Conference on Microbial Genetics and Ecology, BAGECO'11, Corfu, Greece, 2011
- ◊ I. Tagkopoulos, "Microbial Evolution in Stressful Environments: Theory and Experiments", Poster, 11th Conference on Microbial Genetics and Ecology, BAGECO'11, Corfu, Greece, 2011
- V.Mozhayskiy, I. Tagkopoulos, "Facilitated Variation of In Silico Microbial Populations Affects Evolutionary Rates", Poster and Talk, 7th International Symposium on Bioinformatics Research and Applications, Changsha, China, May 2011
- V.Mozhayskiy, I. Tagkopoulos, "Large-scale Evolutionary Simulations of Complex Microbial Behaviors in Dynamic Environments", Poster, 9th Annual International Conference on Computational Systems Bioinformatics, CSB'10, Stanford, 2010
- ◊ I. Tagkopoulos, "Advancing Broader Impacts of CISE Projects: Scientific Discovery and Public Dissemination", Abstract, Invited Participant in NSF Broader Impacts for Research and Discovery Summit, Washington, D.C., 2010
- V.Mozhayskiy, I. Tagkopoulos, "Emergence of Robust Biological Networks in Petascale Simulations of Bacterial Evolution", Abstract, Extreme Scale I/O and Data Analysis Workshop, NSF/NCSA/TACC, Austin, TX, 2010
- V.Mozhayskiy, I. Tagkopoulos, "Simulations of Microbial Evolution in Fluctuating Environments", Poster, 18th Annual International Conference on Intelligent Systems for Molecular Biology, ISMB'10, Boston, 2010
- A structure of Multi-omics data for predictive analytics, University of California, San Diego, CA, June 2015
 - ◊ Big Data on Small Organisms: Integrative Predictive Modeling with Deep Learning, Simons Foundation, NY, May 2015
 - ◊ An integrative, data-driven whole-cell model for systems and synthetic biology, Mosanto Fellows, St. Louis, April 2015
 - ◊ Multi-scale, genome-wide modeling and simulation, PepTalk15, San Diego, Jan 2015
 - ◊ Integrative modeling for targeted experimentation in Systems and Synthetic Biology, Life Sciences, University of Warwick, July 2014
 - ◊ A genome-scale, integrative model for *E. coli* for targeted experimentation, Biodesign Institute, UIUC, May 2014
 - Petascale simulations in microbial evolution: Challenges and Applications, NCSA, May 2014
 - An integrative E. coli genome-scale model for automated design in synthetic biology, 2nd Conference on Quantitative Biology, q-bio'14, Big Island, Hawaii, February 2014
 - ◊ An integrative genome-scale model for *E. coli*, DOE Joint Genome Institute (JGI), November 2013
 - ◊ The emergence and genetic basis of cross-stress behavior in *E. coli*, Department of Microbiology, UC Davis, April 2013
 - ◊ The Evolution of Cross-stress Protection in *E. coli*, Department of Microbiology and Environmental Toxicology, UCSC, February 2013
 - Self-regulatory Recombinant Protein Production through an Integrative Synthetic Biology Approach, Engineering Genes, Vectors, Constructs and Clones, PepTalk, Palm Springs, January 2013
 - ◊ Multiscale Predictive Models for Synthetic Circuits: Potential and Pitfalls, Biology and Mathematics in the Bay Area (BaMBA VII), UCSF, November 2011

Selected Invited Talks

- ♦ The Future of Science, Institute for the Future, Gordon and Betty Moore Foundation, Palo Alto, September 2011
- \diamond A model-based, synthetic biology approach to recombinant protein production, 11th Conference on Microbial Genetics and Ecology, BAGECO'11, Corfu, Greece, 2011
- ◊ An Integrated Approach to Synthetic Biology and Evolution, Progress in Biotechnology Seminar, UC Davis Biotechnology Seminar Series, Davis, May 2011
- ♦ Large-scale Evolutionary Simulations of Complex Microbial Behaviors in Dynamic Environments, 9th Annual International Conference on Computational Systems Bioinformatics, Stanford, 2010
- ♦ Microbial Evolution in Stressful Environments: Theory and Experiments, Math Biology Seminar, Department of Mathematics, UC Davis, Davis, April 2011
- ♦ Emergence of Robust Biological Networks in Petascale Simulations of Bacterial Evolution, Extreme Scale I/O and Data Analysis Workshop, NSF/NCSA/TACC, Austin, TX, 2010
- ◊ Extrapolating in Petascale: Dynamic optimization of simulation environments through process migration, National Center for Supercomputing Applications Workshop, NCSA, Urbana-Champaign, 2010
- ♦ Multiscale models of microbial evolution, Center for Population Biology, UC Davis, Davis, April 2010
- ♦ Emergence of Predictive Capacity within Microbial Networks, School of Computing, Georgia Institute of Technology, Atlanta, Georgia, May 2009
- ◊ LOCUS: An integrative tool for Fixed Income, FX, and other investment instruments, Credit Suisse, New York, April 2009

Honors & Awards	 IGEM Grand Prize, First Place, World Championship, Boston, MA (2014 - 2015) NSF CAREER award (\$600,000), National Science Foundation, DC. (2013 - 2018) Science, Technology& Env. Policy Award (\$60,000), Princeton, NJ. (2006 - 2008) Burroghs Wellcome Fellowship (\$130,000), Princeton, NJ. (2004 - 2006) Princeton Graduate Fellowship (\$54,000), Princeton, NJ. (2003 - 2004) Stanley J. Seeger Fellowship (\$5,000), Princeton, NJ. (2003 - 2008) Paul Nichoplas Excellence Award (\$5,000), Columbia University, NY. (2002 - 2003) Geroundelis Foundation Scholarship (\$4,000), Boston, MA. (2001 - 2002) Teaching Assistant Award (\$500), Columbia University, NY. (March 2002) Honorary Distinction Award (\$1,000), Technical Chamber of Greece. (July 2001) Erasmus Scholarship (\$5,000), University of Aachen (RWTH), Germany. (Fall 1998)
Research Funding	 Petascale Simulations of Complex Biological Behavior in Fluctuating Environments National Science Foundation, NSF-OCI 0941360, PI: Tagkopoulos Dates: 10/2009 - 09/2014, Funds: \$115,000 and allocation on the Blue Waters Supercomputer A Model-Based System for the Automated Design of Synthetic Genetic Circuits by Mathematical Optimization National Science Foundation, NSF-CCF 1146926, PI: Tagkopoulos, co-PI: Matthias Koepper (UC Davis, Math), John Kececioglu (University of Arizona, CS) Dates: 10/2011 - 09/2014, Funds: \$300,000 Modeling and Simulation of Microbial Evolution
	 NSF Teragrid, TG-ASC110007, PI: Tagkopoulos, co-PI: Vadim Mozhayskiy Dates: 03/2011 - 12/2012, 1,500,000 SU in supercomputing systems An integrative framework for MD-based protein-DNA binding prediction NSF XSEDE, DEB130003, PI: Tagkopoulos, co-PI: Vadim Mozhayskiy Dates: 01/2013 - 12/2013, 600,000 SU in supercomputing systems

 An EMR-based, probabilistic clinical support system for the diagnosis and treatment of sepsis patients
 CITRIS seed funding, PI: Tagkopoulos, co-PI: Tim Albertson (UCDMC), Hien Nguyen

(UCDMC), Shawn Newsam (UC Merced) Dates: 06/2012 - 06/2014, Funds: \$50,000

- ◊ Guiding clinical decisions for sepsis patients: A data-driven, probabilistic approach to EHR data mining CTSC seed funding, PI: Tagkopoulos, co-PI: Tim Albertson (UCDMC) Dates: 06/2013 -06/2014, Funds: \$20,000
- ◊ Evolution and dynamics of associative memory in bacterial populations Department of Defense, PI: Tagkopoulos Dates: 07/2012 - 06/2015, Funds: \$495,000
- ◊ Instrumentation for Adaptive Evolution in Bacteria Department of Defense, PI: Tagkopoulos Dates: 06/2013 - 05/2014, Funds: \$250,000
- CAREER: Integrative Synthetic Biology: A Scalable Framework for Modular Multilevel Design
 National Science Foundation, PI: Tagkopoulos
 Dates: 02/2013 - 02/2018, Funds: \$600,000
- Elucidating the Genetic Basis and Evolutionary Potential of Cross-stress Behavior in Escherichia coli
 National Science Foundation, PI: Tagkopoulos
 Dates: 07/2013 06/2015, Funds: \$212,000
- An integrated systems biology approach to elucidate viral resistance signaling networks in tomato
 National Science Foundation, PI: Dinesh-Kumar Savithramma, Co-PIs: Gilbertson, Ullman, Tagkopoulos
 Dates: 07/2014 - 06/2017, Funds: \$1,624,000
- ◊ EVOPROG: General-purpose programmable evolution machine on-a-chip CORDIS FP7-ICT 610730, PI: Alfonso Jaramillo (UoWarwick), Co-PIs: Isalan, Rodriguez, de Lorenzo, Cronin, Tagkopoulos Dates: 10/2013 - 09/2016, Funds: \$3,821,000
- ◊ FIGARO:Flexible and Precise Irrigation Platform to Improve Farm Scale Water Productivity

CORDIS FP7-ICT 610730, PI: Netafim LTD, Israel, 17 Universities Dates: 10/2012 - 09/2017, Role: Participant as an external expert in Decision Support Systems (DSS)

◊ Big Data on Small Organisms: Petascale Simulations of Data-driven Whole-cell Microbial Models

National Science Foundation, NSF-ACI, PI: Tagkopoulos Dates: 06/2015 - 06/2018, Funds: \$40,000 and allocation on the Blue Waters Supercomputer

 Predictive omics profiling in cocoa MARS Inc. PI: Tagkopoulos
 Proprietary algorithms and models for multi-omics profiling. Undisclosed data and amounts (R01 level funding).

- Dr. Navneet Rai, Postdoctoral Research Associate

Ph.D. in Synthetic Biology, Indian Institute of Technology, Bombay, India

- Dr. Nasos Tsoukalas, Postdoctoral Research Associate
- Ph.D. in Electrical Engineering, University of Patras, Greece
- Dr. Violeta Zorraquino Salvo, Postdoctoral Research Associate

	 Ph.D. in Microbiology, University of Pablona Linh Huynh, 5th year Ph.D. Candidate in Computer Science, UC Davis Minseung Kim, 3rd year Ph.D. Candidate in Computer Science, UC Davis Ameen Eetemadi, 2nd year Ph.D. Candidate in Computer Science, UC Davis Beatriz Pereira, 2nd year Ph.D. Candidate in Microbiology, UC Davis Xiaokang Wang, 3rd year Ph.D. Candidate in Biomedical Engineering, UC Davis
	 Past: Dr. Javier Carrera, Ph.D. in Synthetic Biology, Universite de Evry, France currently: Postdoctoral Research Associate at Stanford University Dr. Semarhy Quinones-Soto, Ph.D. in Microbiology, University of California-Davis currently: Faculty Instructor at Sacramento State University, Sacramento, CA Dr. Jiyeon Park, Ph.D. in Microbiology, University of Illinois, Urbana-Champaign currently: Research staff at the Pacific Northwest National Laboratory, Richland, WA Dr. Martin Dragosits, Ph.D. in Biotechnology, BOKU, Austria Dr. Vadim Mozhayskiy, Ph.D. in Physical Chemistry, University of Southern California currently: Scientist at Life Technologies (Ion Torrent) Dr. Matthew Meisner, Population & Ecology (with S. Schreiber, J. Rosenheim) currently: Head of Data Analytics at Farmer's Business Network, Inc. Eren Gultepe, currently Ph.D. Candidate in Neuroscience, Queen's University Daniel Nicklas, currently Ph.D. Candidate in Computer Science, UC Davis Mohammad Gharehyazie, currently Ph.D. Candidate in Computer Science, IST, Austria Adam Sumner, currently at Dynamic Bio, Inc.
PHD Committees	 William Wright, Ph.D Candidate in Mathematics (Advisor: M. Friedlander) Aubrey Gress, Ph.D. Candidate in Computer Science (Advisor: I. Davidson) Victor Missirian, Ph.D. Candidate in Computer Science (Advisor: V. Filkov) Spencer Mathews, Ph.D. Candidate in Biophysics (Advisor: J. Crutchfield) Christopher Schwarz, Ph.D. Candidate in Computer Science (Advisor: O. Carmichael) Jishang Wei, Ph.D. Candidate in Computer Science (Advisor: KL. Ma) Yu-Hsuan Chan, Ph.D. Candidate in Computer Science (Advisor: KL. Ma) Xiang Wang, Ph.D. Candidate in Computer Science (Advisor: I. Davidson) Kristin Lui, Ph.D. Candidate in Mathematics (Advisor: I. Davidson) Rushell Neches, Ph.D. Candidate in Computer Science (Advisor: J. Eisen) Leonid Shamis, M.Sc. Candidate in Computer Science (Advisor: I. Davidson)
Panel Service	 National Science Foundation, Panelist, Broader Impact Panel (CISE Division) National Science Foundation, Panelist, Proposal Review Panel (BIO Division) National Science Foundation, Panelist, Proposal Review Panel (BIO Division) National Science Foundation, Consortium for Mathematics, Panelist and author National Science Foundation, Panelist, Proposal Review Panel (CISE Division) National Science Foundation, Panelist, Proposal Review Panel (CISE Division) National Science Foundation, Panelist, Proposal Review Panel (CISE Division) National Science Foundation, Panelist, Proposal Review Panel (CISE Division) National Science Foundation, Panelist, Proposal Review Panel (CISE Division) National Institutes of Health, Panelist, Scientific Review Group (NHLBI) National Institutes of Health, Panelist, Scientific Review Group (NHLBI) (2015b)
Activities & Broader Impacts	Reviewer for various journals (NPG, AAAS, BMC, Bioinformatics, AMIA, EMBO, ACM, IEEE Transactions, NAR, ACS, among others). REU advisor (3-4 students per year), UC Davis Biotechnology (1-3 students per year) and IGEM synthetic biology team advising (6-10 students per year, providing space, mentorship, funding). Consulting activities and founder of a data analytics company that specializes in bioinformatics solutions.