Claudia R. Solís-Lemus

Curriculum Vitae

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Education

- 2010–2015 Ph.D., Statistics, University of Wisconsin-Madison, Madison.
 - 2013 M.A., Mathematics, University of Wisconsin-Madison, Madison.
- 2003–2008 **B.S., Applied Mathematics**, *Instituto Tecnológico Autónomo de México (ITAM)*, Mexico City, Summa cum laude.
- 2003–2008 **B.S., Actuarial Sciences**, *Instituto Tecnológico Autónomo de México (ITAM)*, Mexico City, Summa cum laude.

Publications

Submitted

- 2018 Fast reconstruction of phylogenetic networks via outlier detection, C. Solís-Lemus.
- 2018 **Prediction of functional markers of mass cytometry data via deep learning**, *C. Solís-Lemus*, *X. Ma, M. Hostetter II, S. Kundu, P. Qiu, D. Pimentel-Alarcón*.
 - Journal papers
- 2018 Greatly reduced phylogenetic structure in the cultivated potato clade (*Solanum* section *Petota* pro parte), *Spooner*, *D.M.*, *H. Ruess*, *C. Arbizu*, *F. Rodríguez*, and *C. Solís-Lemus*, American Journal of Botany, doi:10.1002/ajb2.1008.
- 2017 Phylogenetic comparative methods on phylogenetic networks with reticulations, Bastide, P., C. Solís-Lemus, R. Kriebel, K.W. Sparks, and C. Ané, Systematic Biology, doi: 10.1093/sysbio/syy033.
- 2017 **PhyloNetworks:** a package for phylogenetic networks, *Solís-Lemus, C., P. Bastide and C. Ané*, Molecular Biology and Evolution, doi: 10.1093/molbev/msx235.
- 2016 Inferring phylogenetic networks with maximum pseudolikelihood under incomplete lineage sorting, Solís-Lemus, C. and C. Ané, PLoS Genetics 12(3): e1005896, doi:10.1371/journal.pgen.1005896.
- 2016 Inconsistency of species-tree methods under gene flow, *Solís-Lemus, C., M. Yang and C. Ané*, Systematic Biology, doi: 10.1093/sysbio/syw030.
- 2016 Statistical evidence for common ancestry: application to primates, Baum, D., C. Ané, B. Larget, C. Solís-Lemus, L.S.T. Ho, P. Boone, C. Drummond, M. Bontrager, S. Hunter, B. Saucier, Evolution, doi: 10.1111/evo.12934.
- Bayesian species delimitation combining multiple genes and traits in a unified framework, *Solís-Lemus, C., L.L. Knowles and C. Ané*, Evolution, 2:69,492–507.
 - Conference papers
- 2018 **Breaking the Limits of Subspace Inference**, *Solís-Lemus, C., D. Pimentel-Alarcón*, 56th Annual Allerton Conference on Communication, Control, and Computing.
- 2017 **Processus d'évolution réticulée: tests de signal phylogénétique**, *Ané, C., P. Bastide, M. Mariadassou, S., and C. Solís-Lemus*, Journées de Statistique.
- 2017 Adversarial Principal Component Analysis, Pimentel-Alarcón, D., A. Biswas and C. Solís-Lemus, IEEE International Symposium on Information Theory (ISIT).
- 2016 **Crime detection via crowdsourcing**, *Pimentel-Alarcón*, *D. and C. Solís-Lemus*, 8th Mexican Conference on Pattern Recognition, Springer International.

In preparation

- 2018 Enhanced genome-wide association studies leveraging family history, with M. Epstein, Emory U.
- 2018 **Genome-wide association study of delta toxin production in** *S. aureus*, *with T. Read and M. Su*, Emory U.
- 2018 Bayesian importance sampling for phylogenetic inference, with B. Larget, UW-Madison.
- 2018 Modeling and inference of endogenous retrovirus dynamics, with F. Nascimiento, U. of Oxford.
- 2018 A new robust measure of relative dispersion, with V. Guerrero, ITAM.

Grants

Pending

2018 NIH K99/R00 Path to Independence, Enhanced Penalized Regression Methods for Detecting Trait Loci in GWAS, Role: Pl.

Score: 36

2018 **NIH R01**, Advanced Deep Learning Methods to Predict Disease Risk from a Variety of Health Data, PI: Pimentel-Alarcón (GSU), Role: Significant Contributor.

Fellowships and Awards

2015 Student travel support

Society for Systematic Biologists

2015 Vilas conference presentation funds

UW-Madison

2014 Advanced Computing Infrastructure Fellowship

UW-Madison

2012 College of Letters and Science Teaching Fellowship

UW-Madison

2009 Ex-ITAM Research Award in Actuarial Sciences

ITAM

2008 Social Responsibility Award

ITAM

2003 Academic Excellence Scholarship

ITAM

Software development

- 2015 **PhyloNetworks**, https://github.com/crsl4/PhyloNetworks, Julia package for phylogenetic networks including estimation method SNaQ.
- 2014 iBPP, https://github.com/cecileane/iBPP, Bayesian species delimitation integrating genes and traits.

Research Experience

2017–present **Postdoctoral fellow**, *NIH-supported project*, Epilepsy consortium and 22q11.1 deletion syndrome consortium, PI: Michael Epstein.

Emory University

2016–2017 **Postdoctoral research**, *NSF-supported project*, Improved Bayesian phylogenetic inference based on approximate conditional independence, PI: Bret Larget.

University of Wisconsin-Madison

2014–2015 **Research assistant**, *NSF-supported project*, ATOL: Assembling a taxonomically balanced genome-scale reconstruction of the evolutionary history of the Enterobacteriaceae, PI: Nicole Perna, Bret Larget, Cécile Ané, Colin Dewey.

University of Wisconsin-Madison

2013–2014 **Research assistant**, *NSF-supported project*, Reconciling gene trees: Deciphering the source and extent of genealogical discordance, PI: David Spooner, David Baum, Bret Larget, Cécile Ané, Gregory Thain. University of Wisconsin-Madison

Teaching Experience

- 2016 Guest lecturer, Botany 563: Phylogenetic networks, UW-Madison.
- 2014 Statistics tutorial, PhD qualifying examination summer camp, UW-Madison.
- 2012–2013 **Teaching assistant**, Regression methods for population health, UW-Madison, Professor Ron Gangnon.

- 2011–2012 **Teaching assistant**, *Introduction to Biostatistics for population health*, UW-Madison, Professor Ron Gangnon.
 - 2011 **Teaching assistant**, *Introduction to Statistical methods*, UW-Madison, Professor Kevin Packard (Spring), Kam Wah Tsui (Summer).
 - **Teaching assistant**, *Introduction to theory and methods of mathematical statistics I*, UW-Madison, Professor Zhengjun Zhang.
 - 2010 Instructor in Mathematics, Advanced Algebra II, ITAM.
 - 2010 Instructor in Mathematics, Calculus II, ITAM.
 - 2009 Instructor in Mathematics, Calculus III, ITAM.
- 2009–2010 Instructor in Actuarial Sciences, Actuarial Mathematics I, ITAM.
- 2009–2010 Instructor in Actuarial Sciences, Actuarial Mathematics III, ITAM.

Mentoring Experience

- 2018 Software Development Project, Julia package for GAMuT, Anna Voss Emory University.
- 2018 Statistics Senior Honors Thesis, GWAS for epilepsy, Mengtong Hu Emory University.
- 2016 **Software Development Project**, *Parallelization of SNaQ in PhyloNetworks*, Josh McGrath University of Wisconsin-Madison.
- 2016 **Undergraduate Research Scholar**, *Computing tools for bayesian phylogenetic inference*, Jordan Vonderwell University of Wisconsin-Madison.
- 2016 **Statistics Senior Honors Thesis**, *Computing tools and performance of PhyloNetworks*, Nan Ji University of Wisconsin-Madison.
- 2015 **Statistics Senior Honors Thesis**, *Inconsistency of species-tree methods under gene flow*, Mengyao Yang University of Wisconsin-Madison, published work.
- 2015 **Integrated Biological Sciences Summer Research Program**, *Visualizing inferred phylogenetic networks in Julia*, John Spaw University of Wisconsin-Madison.
- 2014 **Integrated Biological Sciences Summer Research Program**, *Using quartets to estimate phylogenetic networks*, John Malloy University of Wisconsin-Madison.

Presentations

Invited talks

- 2018 **Statistics seminar Creighton University**, *Statistical methods to identify genes associated with disease.*
- 2018 **JSM Advanced statistical inference for stochastic models of evolutionary biology**, *Comparative methods in phylogenetic networks*.
- 2018 **BUGS seminar Georgia Institute of Technology**, Statistical methods to reconstruct phylogenetic networks.
- 2017 **Microbiome group Emory University**, Statistical methods to reconstruct phylogenetic networks.
- 2017 **CIBS Seminar Emory University**, Statistical methods and Julia computings tools for the reconstruction of the tree of life.
- 2017 **Human Genetics Seminar Emory University**, *Phylogenetic inference for big data*.
- 2017 Mathematical Approaches to Evolutionary Trees and Networks BIRS, Likelihood challenges for big trees and networks.
- 2016 **Statistics Seminar UW-Madison**, Phylogenetic inference for Big Data.
- 2016 **Instituto de Biología UNAM Mexico City, Mexico**, *Quartet-based inference of phylogenetic networks*.
- 2016 **Simposio de Inferencia y modelación estadística Guanajuato, Mexico**, *Bayesian phylogenetic inference for big data*.

- 2016 SSB symposium: Advances in the analysis of reticulate population networks Evolution meeting, *Quartet-based inference of phylogenetic networks*.
- 2016 **Phylogenomics symposium and software school Evolution meeting**, *Quartet-based estimation of reticulate evolution*.
- 2015 **System Information Learning Optimization Seminar UW Madison**, *Statistical inference of phylogenetic networks*.
- 2015 **Evolution Seminar Series UW Madison**, Fast reconstruction of hybridization networks from multilocus data.
- 2015 **Networks seminar UW Madison**, Statistical inference on phylogenetic networks.
- 2015 Statistics student seminar UW Madison, Statistical inference on phylogenetic networks.
- 2014 **Seminario Aleatorio ITAM**, The role of Statistics in the inference of the Tree of Life. Contributed talks
- 2018 SACNAS Data Science: The Rise of the Machines, *Identifying genes associated with disease*.
- 2018 SACNAS Postdoc talks, Machine-learning the Tree of Life.
- 2018 **20th IMS New Researchers Conference**, Statistical inference of the Tree of Life.
- 2016 **8th Mexican Conference on Pattern Recognition Guanajuato, Mexico**, *Crime detection via crowdsourcing*.
- 2015 **Evolutionary System Biology and Modeling Workshop UW Madison**, *Statistical inference of phylogenetic networks*.
- 2015 **JSM Bayesian Models and Inference**, Bayesian species delimitation combining multiple genes and traits in a unified framework.
- 2015 **Mathematical and Computational Evolutionary Biology Conference**, *Inferring phylogenetic networks from quartets with maximum pseudolikelihood estimation*.
- 2009 **XXIV Foro Nacional de Estadística**, A parametric measure of dispersion derived from the generalized mean.

Tutorials

- 2017 **Human Genetics Meeting Emory University**, Julia: why do we need another language?.
- 2016 Instituto de Biología UNAM, PhyloNetworks: julia package for phylogenetic networks.
- 2016 **Phylogenomics symposium and software school Evolution meeting**, *PhyloNetworks: julia package for phylogenetic networks.*
- 2016 **Botany 563 UW Madison**, Small tutorial on PhyloNetworks and SNaQ.
- 2015 The hacker within series UW Madison, Short introduction to Julia.
- 2015 Statistics student seminar UW Madison, Short introduction to HTCondor.

Service

since 2016 Systematic Biology, Editorial Board Member.

Symposium organization

- 2018 **SACNAS symposium**, *Data Science: The Rise of the Machines*, Co-organizer: Daniel Pimentel-Alarcón (GSU).
- 2016 **Evolution meeting SSB symposium**, *Advances in the analysis of reticulate population networks*, Co-organizer: David Baum (UW-Madison).

Volunteer work

2005-2008 Sistema de preparatoria abierta para adultos, Math high-school teacher for adults, ITAM, Mexico.

Languages

Spanish Native language

English Proficient level: TOEFL iBT (2009), IELTS (2009)

French Advanced level: DALF C1 (2007)

Portuguese Advanced level: CELPE-BRAS (2008)

German Beginner level: A1 Start Deutsch 1 (2008)

Computer skills

Programming C++, Java, Julia, Perl, Python

Statistics R, SAS, SPSS

Mathematics Matlab, Mathematica, Macaulay2

HTC HTCondor, OSG, SLURM

Certifications

Exam C Society of Actuaries (SOA)

Exam FM Society of Actuaries (SOA)

Exam P Society of Actuaries (SOA)

Construction of Actuarial Models Certificate

Financial Mathematics Certificate

Probability Certificate

Statistical consulting

2016 Analysis of education programs for adults in indigenous populations

INEA. Mexico

References

- o Cécile Ané, Professor of Statistics and Botany, UW-Madison cecile.ane@wisc.edu
- Bret Larget, Professor of Statistics and Botany, UW-Madison bret.larget@wisc.edu
- O Douglas Bates, Emeritus Professor of Statistics, UW-Madison bates@wisc.edu
- o David Baum, Chair of Botany, UW-Madison dbaum@wisc.edu
- Kevin Packard, Statistical consultant, Cornell U. (to address teaching), kcp48@cornell.edu
- Ron Gangnon, Professor of Biostatistics and Population Health (to address teaching) ronald@biostat.wisc.edu
- o Mike Epstein, Professor of Human Genetics, Emory University mpepste@emory.edu

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