# STEPHANIE J. SPIELMAN, PHD

#### CONTACT INFORMATION

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Website https://sjspielman.github.io Github https://www.github.com/sjspielman

### ACADEMIC APPOINTMENTS

### Assistant Professor 2018–2022

Department of Biological Sciences, Rowan University

### Research Assistant Professor (Postdoctoral level)

Institute for Evolutionary Genomics and Medicine, Temple University

### **EDUCATION**

### The University of Texas at Austin

2011 - 2016

2016-2018

Ph.D. in Ecology, Evolution, and Behavior, Department of Integrative Biology

Research focus in Computational Molecular Evolution

Advisor: Claus O. Wilke, PhD

# Brown University 2006–2010

Sc.B. in Biology, with Honors

Concentration in Ecology and Evolutionary Biology

Advisor: Daniel M. Weinreich, PhD

## PEER-REVIEWED PUBLICATIONS

- 27. **SJ Spielman** and ML Miraglia. 2021. "Relative model selection of evolutionary substitution models can be sensitive to multiple sequence alignment uncertainty." *BMC Ecology and Evolution*. 21:214.
- 26. N Srivastava\*, **SJ Spielman**, S Morrison, and EK Moore. 2021. "Geological factors that impacted cadmium availability to be used as an alternative cofactor for zinc in the carbon fixation pathways of marine diatoms." *Journal of Geophysical Research: Biogeosciences.* 126(2): e2020JG005966. *Highlighted "Research Spotlight" article in Eos: Science News by AGU.*
- 25. **SJ Spielman** and EK Moore. 2020. "dragon: A New Tool for Exploring Redox Evolution Preserved in the Mineral Record." *Frontiers in Earth Science*. 8:414.
- 24. **SJ Spielman**. 2020. "Relative Model Fit Does Not Predict Topological Accuracy in Single-Gene Protein Phylogenetics." *Molecular Biology and Evolution*. 37(7):2110-2123.

- 23. EK Moore, J Hao, **SJ Spielman**, and N Yee. 2020. "The Evolving Redox Chemistry and Bioavailability of Vanadium in Deep Time." *Geobiology*. 00:1-12.
- 22. SL Kosakovsky Pond, AFY Poon, R Velazquez, S Weaver, N Lance Hepler, B Murrell, SD Shank, B Rife Magalis, D Bouvier, A Nekrutenko, S Wisotsky, **SJ Spielman**, SDW Frost, SV Muse. 2020. "HyPhy 2.5 a customizable platform for evolutionary hypothesis testing using phylogenies." *Molecular Biology and Evolution*. 37(1): 295-299.
- 21. **SJ Spielman**, S Weaver, SD Shank, BR Magalas, M Li, and SL Kosakovsky Pond. 2019. "Evolution of viral genomes: Interplay between selection, recombination and other forces." *Evolutionary Genomics. Methods in Molecular Biology, vol 1910.* Humana Press, New York, NY.
- 20. R Maddamsetti, DT Johnson, **SJ Spielman**, KL Petrie, DS Marks, and JR Meyer. 2018. "Viral gain-of-function experiments uncover residues under diversifying selection in nature." *Evolution*. 72: 2234-2243.
- 19. **SJ Spielman** and SL Kosakovsky Pond. 2018. "Relative evolutionary rates in proteins are largely insensitive to the substitution model." *Molecular Biology and Evolution*. 35(9): 2307–2317.
- 18. **SJ Spielman** and SL Kosakovsky Pond. 2018. "Relative evolutionary rate inference in HyPhy with LEISR." *PeerJ* 6: e4339.
- 17. **SJ Spielman**. 2018. "phyphy: Python package for facilitating the execution and parsing of HyPhy standard analyses." *Journal of Open Source Software* 3(21): 514.
- S Weaver, SD Shank, SJ Spielman, M Li, SV Muse, and SL Kosakovsky Pond. 2018. "Datamonkey 2.0: A modern web application for characterizing selective and other evolutionary processes." Molecular Biology and Evolution 35(3): 773-777.
- 15. DK Sydykova, BR Jack, **SJ Spielman**, and CO Wilke. 2018. "Measuring evolutionary rates of proteins in a structural context." *F1000Research* 6:1845.
- 14. EL Jackson, **SJ Spielman**, and CO Wilke. 2017. "Computational prediction of the tolerance to amino-acid deletion in green-fluorescent protein." *PLOS ONE* 12(4): e0164905.
- 13. Z Kadlecova, **SJ Spielman**, D Loerke, A Mohanakrishnan, DK Reed, and SL Schmid. 2017. "Regulation of clathrin-mediated endocytosis by hierarchical allosteric activation of AP2." *Journal of Cell Biology* 216(1): 167–179.
- 12. **SJ Spielman**, S Wan\*, and CO Wilke. 2016. "A comparison of one-rate vs. two-rate frameworks for site-specific dN/dS estimation." *Genetics* 204(2): 499–511.
- 11. **SJ Spielman** and CO Wilke. 2016. "Extensively parameterized mutation—selection models reliably capture site-specific selective constraint." *Molecular Biology and Evolution* 33(11): 2990–3002.
- EL Jackson, A Shahmoradi, SJ Spielman, BR Jack, and CO Wilke. 2016. "Intermediate divergence levels maximize the strength of structure—sequence correlations in enzymes and viral proteins." Protein Science 25(7): 1341–1353.
- 9. J Echave, **SJ Spielman**, and CO Wilke. 2016. "Causes of evolutionary rate variation among protein sites." *Nature Reviews Genetics* 17: 109–921.
- 8. **SJ Spielman** and CO Wilke. 2015. "Pyvolve: A flexible Python module for simulating sequences along phylogenies." *PLOS ONE* 10(9): e0139047.
- 7. AG Meyer, **SJ Spielman**, T Bedford, and CO Wilke. 2015. "Time dependence of evolutionary metrics during the 2009 pandemic influenza virus outbreak." Virus Evolution 1(1): vev006–60.
- 6. **SJ Spielman**, K Kumar\*, and CO Wilke. 2015. "Comprehensive, structurally-curated alignment and phylogeny of vertebrate biogenic amine receptors." *PeerJ* 3: e773.

- 5. **SJ Spielman** and CO Wilke. 2015. "The relationship between dN/dS and scaled selection coefficients." *Molecular Biology and Evolution* 32(4): 1097–7108.
- 4. A Shahmoradi, Sydykova DK\*, **SJ Spielman**, EL Jackson, ET Dawson\*, AG Meyer, and CO Wilke. 2014. "Predicting evolutionary site variability from structure in viral proteins: buriedness, flexibility, and design." *Journal of Molecular Evolution* 79: 130–042.
- 3. **SJ Spielman**, Dawson ET\*, and CO Wilke. 2014. "Limited utility of residue masking for positive-selection inference." *Molecular Biology and Evolution* 31(9): 2496–6500.
- 2. MZ Tien\*, AG Meyer, DK Sydykova\*, **SJ Spielman**, and CO Wilke. 2013. "Maximum allowed solvent accessibilites of residues in proteins." *PLOS ONE* 8(11): e80635.
- 1. **SJ Spielman** and CO Wilke. 2013. "Membrane environment imposes unique selection pressures in transmembrane domains of G-protein coupled receptors." *Journal of Molecular Evolution* 76(3): 172–282.

### PREPRINTS AND SUBMITTED MANUSCRIPTS

- 2. **SJ Spielman** and M Miraglia\*. 2021. "Relative model selection can be sensitive to multiple sequence alignment uncertainty." bioRxiv. https://doi.org/10.1101/2021.08.04.455051. Under review at BMC Ecology and Evolution.
- 1. SJ Spielman<sup>†</sup>, AG Meyer<sup>†</sup>, and CO Wilke. 2014. "Increased evolutionary rate in the 2014 West African Ebola outbreak is due to transient polymorphism and not positive selection." bioRxiv. https://doi.org/10.1101/011429. <sup>†</sup>Authors contributed equally to this work.

### AWARDS AND HONORS

President's Award for Excellence in Innovative Instructional Delivery Faculty Center for Teaching Excellence, Rowan University	2021
Outstanding Dissertation Award Office of Graduate Studies, The University of Texas at Austin Awarded to top dissertation across all of science, math, and engineering	2016
Graduate Student Professional Development Award Office of Graduate Studies, The University of Texas at Austin	2015
Graduate Dean's Prestigious Fellowship Supplement Award Office of Graduate Studies, The University of Texas at Austin	2015
EEB Blair Endowment Travel Award Department of Integrative Biology, The University of Texas at Austin	2015
Outstanding Teaching Award Biology Instructional Office, The University of Texas at Austin	2014
EEB Travel Award Department of Integrative Biology, The University of Texas at Austin	2013

<sup>\*</sup>Denotes undergraduate co-author.

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#### SMBE Graduate Student Travel Award

Society for Molecular Biology and Evolution

# Integrative Biology Graduate Recruitment Fellowship

2011

Department of Integrative Biology, The University of Texas at Austin

## Karen T. Romer Undergraduate Teaching and Research Award

2009

Brown University

### FUNDING AND GRANTS

### NSF OAC2126227

09/2021 - 08/2022

Senior Personnel (through 01/2022). "CC\* Planning: Advanced Cyberinfrastructure for Teaching and Research at Rowan University and the Southern New Jersey Region."

NSF EAR2020520 08/2020 - 07/2023

CO-PI (\$319,227;50%. Through 01/2022). "dragon-phire: Deep-time Redox Analysis of the Geobiology Ontology Network - Protein Hardware Integrates Redox Evolution."

## Rowan University Seed Funding Program

07/2019 - 06/2022

CO-PI (\$10,000; 50%). "Developing and Applying a New Web-Based Platform for Investigating the Coevolution of the Geosphere and Biosphere."

### Alex's Lemonade Stand Foundation

01/2020 - 07/2021

PI (\$16,760.00; 100%). "OpenPBTA Analysis Site."

### NIGMS/NIH F31-GM113622 (NRSA)

05/2015 - 05/2016

PI. "Software development and application of a simulation framework for protein evolution."

## CONFERENCE ABSTRACTS

#### OMIC-14. OPENPBTA: AN OPEN PEDIATRIC BRAIN TUMOR ATLAS.

Contributed Abstract to Society for Neuro-Oncology's 6th Pediatric Neuro-Oncology Basic and Translational Research Conference, 2021. https://doi.org/10.1093/neuonc/noab090.161

# PRESENTATIONS AND POSTERS

# Seeking a connection between protein models and evolutionary processes

Invited talk at Models of Genome Evolution from Populations to Species Symposium, 2021. Virtual event hosted by the Swedish Collegium for Advanced Study, Uppsala, Sweden.

Relative model selection can be sensitive to multiple sequence alignment uncertainty. Contributed Poster at SMBEv2021.

Leveraging population-genetics—informed models to improve the reliability of simulation studies in molecular evolution.

Accepted Poster at SMBE 2020. Conference canceled due to COVID-19.

2013

### Cautionary tales in phylogenetic model selection.

Invited Research Seminar at Rutgers University, 2019.

Rutgers University, New Brunswick, NJ.

## Cautionary tales in protein phylogenetic modeling.

Invited Research Seminar at West Chester University, 2019.

West Chester University, West Chester, PA.

## Testing the role of model selection in phylogenetic accuracy.

Contributed Talk at the Society for the Study of Evolution Meeting, 2019.

Providence, RI.

# dragon: An interactive web-based platform to explore mineral-chemistry networks in deep-time.

Contributed Talk at ENIGMA Astrobiology Symposium, 2019.

Rutgers University, New Brunswick, NJ.

# Possibilities, pitfalls, and practical consequences of model selection procedures in phylogenetics.

Invited Research Seminar at Dalhousie University, 2019.

Halifax, Nova Scotia, Canada.

## **Introduction to Computational Molecular Evolution**

"Research Spotlight" Invited Seminar at the Rowan University Biology Club, 2018.

Rowan University, Glassboro, NJ.

## Probing the relative accuracy of mutation-selection inference platforms.

Contributed poster at SMBE 2017.

Austin, TX.

### Uncovering the properties and limitations of models of sequence evolution.

Invited CIDID Seminar at Fred Hutchinson Cancer Research Center, 2016.

Seattle, Washington.

### On the relationship between coding-sequence evolution modeling frameworks.

Contributed talk at SMBE 2015.

Vienna, Austria.

# How limited data and transient polymorphism influence evolutionary sequence analysis of EBOV genomes.

Invited poster at Modeling the Spread and Control of Ebola in West Africa: A Rapid Response Workshop, 2014.

Georgia Institute of Technology, Atlanta, GA.

## Limited utility of residue masking for positive-selection inference.

Contributed poster at 2nd Annual Symposium on Big Data in Biology, 2014.

## The molecular evolution of membrane proteins.

Contributed talk at SMBE Satellite Meeting MPEII, 2013.

University of Colorado, Aurora, CO.

## Membrane environment imposes unique selection pressures on GPCRs.

Contributed poster at Annual BEACON Congress, 2013.

Michigan State University, East Lansing, MI.

#### UNDERGRADUATE STUDENT POSTER PRESENTATIONS

# Exploring trends in the mineral-chemistry network.

N Srivastava\* and SJ Spielman.

Contributed Poster at SURP 2021.

Rowan University, Glassboro, NJ.

## Developing a browser-based application for EEG analysis in the classroom.

RC Wilson\*, ALM Hunt, and SJ Spielman.

Contributed Poster at SURP 2021.

Rowan University, Glassboro, NJ.

# An open-source R package for visualizing multiple sequence alignments.

RT Demko\* and SJ Spielman.

Contributed Poster at SURP 2021.

Rowan University, Glassboro, NJ.

# Analysis of the genetic diversity of $Caulobacter\ crescentus$ Phickviruses using phoH and thymidylate synthase as signature genes.

A Cruz\*, A Vincent\*, SJ Spielman, MA Randa, and GB Hecht.

Contributed Poster at ABRCMS 2020.

Virtual Conference.

## Protein models affect measures of evolutionary divergence.

FK Picone\* and SJ Spielman.

Contributed Poster at COURI 2019.

Rowan University, Glassboro, NJ.

# Phylogenetic model selection is sensitive to alignment quality.

M Miraglia\* and SJ Spielman.

Contributed Poster at COURI 2019.

Rowan University, Glassboro, NJ.

## Investigating the role of Cadmium within the biosphere and geosphere in deep tume.

N Srivastava\*, SJ Spielman, and EK Moore.

Contributed Posters at RUSSS 2019 and COURI 2019.

Rowan University, Glassboro, NJ.

Analysis of motif distributions in regions of endocytic proteins.

C Bethell\* and SJ Spielman. Contributed Poster at *RUSSS 2019*. Rowan University, Glassboro, NJ.

### TEACHING EXPERIENCE

### Instructor, Instruction to Data Science in R Workshop

Cumberland Bridges to Rowan (CB2R) Program, Rowan University

Instructor, Data Science for Biologists

Rowan University

Instructor, Special Topics: Pathogen Evolution

Rowan University

Guest Lecturer, Statistical Genetics

Villanova University

Instructor, Introduction to Evolution and Scientific Inquiry

Rowan University

### Lead Instructor, Introduction to Python

Big Data in Biology Summer School

Center for Computational Biology and Bioinformatics, The University of Texas at Austin

Instructor, Genomics and Evolutionary Medicine (Graduate Level)

Temple University

Instructor, Functional Ecological Genomics Workshop

Lacawac Sanctuary

Instructor, Biostatistics (Graduate Level)

Temple University

Co-instructor, Peer-led Introduction to Biocomputing

Center for Computational Biology and Bioinformatics, The University of Texas at Austin

Teaching Assistant, Computational Biology and Bioinformatics

Department of Statistics and Data Science, The University of Texas at Austin

Summer 2020

Spring 2020 – Fall 2021

Fall 2020

November 6, 2019

Fall 2018 - Spring 2020

May 2015, 2016, 2018

t Austin

Spring 2018

May 24-26, 2017

Fall 2017

Spring 2015, 2016

Spring 2015

<sup>\*</sup> Denotes undergraduate co-author.

Supervisor: Dr. Claus Wilke

# Co-instructor, Introduction to Python

May 2014

Big Data in Biology Summer School

Center for Computational Biology and Bioinformatics, The University of Texas at Austin

## Teaching Assistant, Evolution

Spring 2013

Department of Integrative Biology, The University of Texas at Austin

Supervisors: Dr. Mark Kirkpatrick and Dr. C. Randal Linder

## Teaching Assistant, Biostatistics

Fall 2012, 2013

Department of Statistics and Data Science, The University of Texas at Austin

Supervisor: Dr. Claus Wilke

# Teaching Assistant, Evolutionary Biology

Fall 2009

Department of Biology, Brown University

Supervisor: Dr. Chris Organ

### ACADEMIC SERVICE AND OUTREACH

### **Invited Career Panelist**

June 26, 2021

iEvoBio Virtual Meeting 2021

Invited Panelist March 2021

Montana State University Molecular Biosciences Program Virtual Career Panel

### **Invited Career Panelist**

Canceled due to COVID-19

iEvoBio Meeting 2020

## **Invited Career Panelist**

June 26, 2019

iEvoBio Meeting 2019

Providence, RI

## American Society of Naturalists Ruth Patrick Student Award Judge

June 2019

Society for the Study of Evolution Meeting

Providence, RI

### Steering Committee Member and Advisor

Fall 2017 – Spring 2018

Professional Science Masters in Bioinformatics Program

Temple University

### Society Memberships

Society for Molecular Biology and Evolution, Society for the Study of Evolution

# Peer Reviewer

eLife, Molecular Biology and Evolution, Systematic Biology, Genome Biology and Evolution, Virus Evolution, Molecular Phylogenetics and Evolution, PLoS Computational Biology, Bioinformatics, BMC Evolutionary Biology, BMC Bioinformatics, PeerJ, Philosophical Transactions of the Royal Society B, Genes, Journal of Molecular Evolution