Emily Jane McTavish

CONTACT Information School of Natural Sciences University of California, Merced 5200 N. Lake Rd, Merced CA 95343

ejmctavish@ucmerced.edu

EDUCATION

University of Texas, Austin, Texas

Ph.D, Ecology, Evolution and Behavior

May 2013

Advisor: David M. Hillis

Dissertation: Estimating population histories using single-nucleotide polymorphisms sampled through-

out genomes

 \mathbf{McGill} University, Montreal, Quebec, Canada $\mathit{BS},$ Honours Biology

April 2006

Professional Appointments University of California, Merced, Merced, CA

Associate Professor, Life and Environmental Sciences

June 2016 - Present

Assistant Professor, Life and Environmental Sciences

June 2016 - 20

University of Kansas, Lawrence, Kansas

Postdoctoral researcher, Department of Ecology and Evolutionary Biology

Advisor: Mark T. Holder

May 2013 - June 2016

Heidelberg Institute for Theoretical Studies, Heidelberg, Germany

Humboldt Research Fellow, Scientific Computing Group

Advisor: Alexandros Stamatakis

December 2014 - August 2015

Publications

 $^{\wedge}$ indicates a postdoctoral or graduate student advisee

Submitted

Mikryukov, V., Abarenkov, K., Laffan, S., Robertson, T., **McTavish, E.J.**, Stjernegaard Jeppesen, T., Waller, J., Blissett, M., Köljalg, U., Miller, J. *in review*. PhyloNext: a pipeline for phylogenetic diversity analysis of GBIF-mediated data.

^Lopez Fang, L., Ortega-Del Vecchyo, D., **McTavish, E.J.**, Huerta-Sanchez, E. *in review*. Leveraging shared ancestral variation to detect local introgression. Preprint posted on *bioRxiv*https://doi.org/10.1101/2022.03.21.485082

Refereed journal articles

^Sánchez Reyes, L.L., **McTavish, E.J.**, O'Meara, B. *accepted*. DateLife: leveraging databases and analytical tools to reveal the dated Tree of Life. *Systematic Biology*, Preprint posted on *bioRxiv*; doi: https://doi.org/10.1101/782094

^Sánchez Reyes, L.L., **McTavish**, **E.J.** (2022) Approachable case studies support learning and reproducibility in data science: An example from evolutionary biology, *Journal of Statistics and Data Science Education*, 10.1080/26939169.2022.2099487

^Toscani Field, J., Abrams, J., Cartee, J., McTavish, E.J. 2022. Rapid Alignment Updating with Extensiphy. Methods in Ecology and Evolution 13:682-693 GitHub repository: https://github.com/McTavishLab/Extensiphy

^Sánchez Reyes, L.L., ^Kandziora, M., **McTavish, E.J.** 2021. Physcraper: a python package for continual update of evolutionary estimates using the Open Tree of Life. *BMC Bioinformatics* 22:355 Software package reviewed and accepted by *pyOpenSci*. GitHub repository: https://github.com/McTavishLab/Physcraper

Brock, K., McTavish, E.J., Edwards, D. 2022. Color Polymorphism is a Driver of Diversification in the Lizard Family Lacertidae. Systematic Biology 7:24-39

Publications continued

McTavish, E.J., Sánchez Reyes, L.L., Holder, M.T. 2021. OpenTree: A Python package for Accessing and Analyzing data from the Open Tree of Life Systematic Biology 6:1295–1301. GitHub repository: https://github.com/OpenTreeofLife/python-opentree

Lin, M., Simons, A.L., Curd, E.E., Harrigan, R.J., Schneider, F.D., Ruiz-Ramos, D.V., Gold, Z., Osborne, M.G., Shirazi, S., Schweizer, T.M., Moore, T.N., Fox, E.A., Turba, R., Garcia-Vedrenne, A.E., Helman, S.K., Rutledge, K., Mejia, M.P., Munguia Ramos, M.N., Wetzer, R., Pentcheff, D., McTavish, E.J., Dawson, M.N., Shapiro, B., Wayne, R.K., Meyer, R.S., 2021. Landscape analyses using eDNA metabarcoding and Earth observation predict community biodiversity in California. *Ecological Applications* 31(6):e02379

McTavish, E.J., Drew, B.T., Cranston K. 2017. How and why to build a unified tree of life. *BioEssays* 39:1700114.

McTavish, E.J., Pettengill, J., Davis, S., Rand, H., Strain, E., Allard, M., Timme R.E. 2017. TreesToReads: Simulating data to test phylogenetic effects of biases in SNP calling pipelines. *BMC Bioinformatics* 18:178. GitHub repository: https://github.com/snacktavish/TreeToReads

Hinchliff, C., Smith, S.A., Allman, J.A., Burleigh, G., Chaudhary, R., Coghill, L.M., Crandall, K.A., Deng, J., Drew, B.T., Gazis, R., Gude, K., Hibbett, D., Katz, L.A., Laughinghouse, H.D., **McTavish**, **E.J.**, Mitford, P., Owen, C., Ree, R., Soltis, D., Williams, T., Cranston, K.A. 2015. Synthesis of phylogeny and taxonomy into a comprehensive tree of life. *Proceedings of the National Academy of Sciences USA*. 112:12764-1276

McTavish, E.J., Holder M.T., Steel, M. 2015. Twisted trees and inconsistency of tree estimation when gaps are treated as missing data – two examples of inconsistency caused by non-linear distance corrections. *Molecular Phylogenetics and Evolution*. 93:289–295

McTavish, E.J., Hinchliff, C., Allman, J.F., Brown, J., Cranston, K.A., Holder M.T., Rees, J. A., Smith, S. A. 2015. Phylesystem: a git-based data store for community curated phylogenetic estimates. *Bioinformatics*. 17:2794-800

McTavish, E.J., Hillis, D.M. 2015. How does ascertainment bias in SNP analyses affect inferences about population history? *BMC Genomics*. 16:266

McTavish, E.J., Hillis, D.M. 2014. A genomic approach for distinguishing between recent and ancient admixture in cattle. *Journal of Heredity*. 105:445–456

McTavish, E.J., Decker, J.E., Schnabel, R.D., Taylor, J.F., Hillis, D.M. 2013. New World cattle show ancestry from multiple independent domestication events. *Proceedings of the National Academy of Sciences USA*. 110:E1398-E1406

Stoltzfus, A., Lapp, H., Matasci, N., Deus, H., Sidlauskas, B., Zmasek, C.M., Vaidya, G., Pontelli, E., Cranston, K., Vos R., Webb, C.O., Harmon, L.J., Pirrung, M., O'Meara, B., Pennell, M.W., Mirarab, S., Rosenberg, M.S., Balhoff, J.P., Bik, H.M., Heath, T., Midford, P., Brown J.W., McTavish E.J., Sukumaran J., Westneat M., Alfaro M.E., Steele A. 2013. Phylotastic! Making Tree-of-Life Knowledge Accessible, Re-usable and Convenient. BMC Bioinformatics. 14:1-1

McTavish*, E.J., Smith*, G., Guerrero, R., Gering, E. 2012. Flight morphology variation in a damselfly with female limited polymorphism. Evolutionary Ecology Research. 14:325-341

Brown, J. Savidge, K. McTavish, E.J. 2010. DIM SUM: Demography and Individual Migration Simulated Using a Markov chain. *Molecular Ecology Resources*. 11:358-363

Smith, G., Gering, E. R. Guerrero, **McTavish, E.J.**, Lydgate, T. 2009. Theobroma cacao L. (Malvaceae) agroecology in Kauai: a case study. Pacific Agriculture and Natural Resources 1:21-26

Book chapters

Holder, M. T. and **McTavish**, **E.J.** 2016. Hypothesis tests for comparing trees. Encyclopedia of Evolution, Elsevier.

Publications Continued Non-refereed publications

Klott, J. S., M. C. Whitfield, M. Cota, **McTavish**, **E.J.** 2007. 2006-2007 wildlife inventory in the Jarbidge field office. BLM ID Tech. Bull. 2007-003.

Refereed conference proceedings

Holder, M.; Cranston, K. A.; **McTavish, E. J.**, 2019 The Need for Semantically Rich Notions of "Taxa" for Interoperability in Biodiversity Informatics. *American Geophysical Union*,

McTavish, E.J. 2019. Linking Biodiversity Data Using Evolutionary History. (Biodiversity_Next). Biodiversity Information Standards,

McTavish, E.J., Holder, M. T., Cranston, K. A. 2018. Nurturing a sustainable Open Tree of Life (TDWG). Biodiversity Information Standards, e25727,

Invited Talks Seminar October 13, 2023, Cornell Lab of Ornithology, Cornell University, Ithaca, NY Seminar May 19, 2023, EMBO Satellite Workshop on Biodiversity Informatics, Hellenic Center for Marine Research, Heraklion, Crete Seminar October 16, 2020, Cornell Lab of Ornithology, Cornell University, Ithaca, NY Seminar October 16, 2019, Humboldt State University, Arcta, CA Seminar January 4, 2019, Instituto Gulbenkian de Ciência, Oieras, Portugal November 29, 2018, UC Berkeley, Berkeley, CA Integrative Biology seminar Integrative Biology seminar October 31, 2018, University of Minnesota, Minneapolis, MN Computational Biology seminar February 21, 2017, Louisiana State University, Baton Rouge, LA Debate January 10, 2017, Society of Systematic Biologists, Baton Rouge, LA Computational Biology seminar

November 15, 2016, Fred Hutchinson Cancer Research Center, Seattle, WA Biology seminar September 28, 2016, University of California, Los Angeles, CA Frontiers in Phylogenetics Symposium

September 9, 2016, Smithsonian Natural History Museum, Washington, DC
Botany seminar March 23, 2016, Oklahoma State University, Stillwater, OK
A. Watson Armour III Research Seminar October 21, 2015, The Field Museum, Chicago, IL
Population genetics group

September 14, 2015, Ecole Polytechnique Fédérale de Lausanne (EPFL), Lausanne, Switzerland Seminar September 10, 2015, Laboratoire de Biométrie et Biologie Evolutive, Lyon, France Seminar

September 7, 2015, Centre d'Ecologie Fonctionnelle et Evolutive (CEFE), Montpellier, France Early Career Scientists Symposium March 29, 2014, University of Michigan, Ann Arbor, MI NSF BEACON site visit December 8, 2011, Michigan State University, East Lansing, MI IBEST Seminar October 6, 2011, University of Idaho, Moscow, ID

FUNDING AND AWARDS

- 2022 U-RISE at UC Merced (Senior Personnel) NIH URISE
- 2022 Supplemental funding "Cultivating a Sustainable Open Tree of Life" NSF DBI (~\$53,000)
- 2021 GEO-Microsoft Planetary Computer Programme "Phylogenetic Diversity in the cloud" (\$60,000 & Microsoft Azure cloud credits in the amount of \$60,000, with Co-PI's J. Miller, M.T. Holder, S.Laffan)
- 2021 Supplemental funding "Elucidating and Detecting Adaptive Introgression" NSF DEB (~\$60,000)
- 2019 Full proposal invited NSF "Evolutionarily informed eco-inference from large data sets"
- 2019 Invited participant- NSF "Harnessing the Data Revolution" Ideas Lab, May 2019, Tysons Corner, VA.
- 2018 PI "Cultivating a Sustainable Open Tree of Life" NSF ABI (~\$435,000)
- 2017 UC Merced, Committee on Research award.
- 2014 Humboldt Postdoctoral Fellowship, Alexander von Humboldt foundation. Germany.
- 2014 Invited Speaker, Early Career Scientists Symposium. University of Michigan.
- 2014 Travel award, National Evolutionary Synthesis Center (NESCent).
- 2012 Research funding (~\$40,000 co-author with J. Sullivan, B. Sarver and D. M. Hillis), BEACON Center for the study of Evolution in Action.
- 2011 Research funding (~\$150,000, co-author with J. Sullivan, J. Foster and D. M. Hillis), BEACON

 $2011~\mathrm{XSEDE}$ (Extreme Science and Engineering Discovery Environment), $200,\!000~\mathrm{HPC}$ compute time allocations.

TEACHING EXPERIENCE	University of California, Merced Graduate Phylogenetics, Fall 2017, Spring 2020, Spring 2022 Introductory Biology, Fall 2018, Fall 2019 Evolution, Spring 2017, Spring 2019, Fall 2022, Spring 2023, Fall 2023 Phylogenetics, Spring 2018				
	Computational Molecular Evolution	Computational Molecular Evolution Workshop July 19-27, 2022, Hinxton, UK			
	Molecular Evolution Workshop at the Marine Biological Laboratory				
	Faculty Teaching Assistant Teaching Assistant	May 26 to June 5, 2023, Woods Hole, MA May 27 to June 6, 2022, Woods Hole, MA August 1 to August 11, 2019, Woods Hole, MA July 18 to July 28, 2018, Woods Hole, MA July 20 to July 30, 2017, Woods Hole, MA July 17 to July 27, 2016, Woods Hole, MA July 19 to July 29, 2015, Woods Hole, MA July 27 to August 6, 2014, Woods Hole, MA July 21 to July 31, 2013, Woods Hole, MA July 22 to August 1, 2012, Woods Hole, MA			
	Trees in the Desert, Large Scale Ph	ylogenetics Workshop	April 12-14, 2019, Biosphere 2, AZ		
	Workshop on Quantitative Evolutionary Biology				
	Young Scientist Lecturer September 13-21, 2014, Nesin Mathematics Village, Şirince, Turkey				
	Software Carpentry Workshops				
	Organizer, Instructor, University Organizer, University of Californi Instructor, University of Oklahon Instructor, University of Missour Instructor, University of Kansas, Coordinator, Instructor, UT Aus Teaching Assistant, Michigan Sta	ia, Merced na, i, Kansas City, tin,	January 10-11, 2018, Merced, CA August 17-18, 2017, Merced, CA April 4-5, 2014, Norman, OK Feb 15-16, 2014, Kansas City, KS Aug 22-23, 2013, Lawrence, KS Dec 10-11, 2012, Austin, TX May 7-9, 2012, Austin, TX		
	Statistics and Scientific Computing Short Course (Python)				
	Instructor Instructor	, · ,	Feb 26, 2013, Austin, TX Nov 26, 2012, Austin, TX		
	University of Texas				
	Teaching Assistant		Introductory Biology, Fall 2007 on, Spring 2008, Spring 2009, Fall 2009 (Field course), Fall 2008, Spring 2010		
Working Groups	Phyloref/OpenTree/Phenoscape		April 3-5, 2019, Durham, NC		
	Next Generation Phenomics Tools f	or the Tree of Life	May 24-25, 2018, Portland, ME		
	Reproducible Research Hackathon		January 9, 2018, Berkeley, CA		
	Computable Evolutionary Phenotype	oe Knowledge	December 11-14, 2017, Durham, NC		
	Futurephy/Arbor/OpenTree Clade	Workshop	May 4-5, 2017, Chicago, IL		
	Futurephy/Arbor/OpenTree Clade	Workshop	October 21-23 2016, Chicago, IL		
	FuturePhy Interoperability Hackath	non	August 15-18 2016, Lawrence KS		

Phylotastic Hackathon at the National Center for Evolutionary Synthesis (NESCent) June 4-8, 2012, Durham, NC

February 2016, Gainesville, FL

July 23, 2014, Toronto, ON, Canada

Futurephy/Arbor/OpenTree Clade Workshop

Mozilla Science Lab Global Sprint

Symposia Organized Putting 'Dark Matter' Biodiversity on the Map: Crosslinking Metagenomics, Phylogenetics and Specimen Occurrence Data Infrastructures to Advance Scientific Understanding and Policy Relevance of Cryptic Biodiversity (with Kyle Copas)

American Geophysical Union (AGU)

December 9-13 2019, San Francicso, CA

Linking biodiversity data through phylogenetic knowledge (with Guanyang Zhang, Gaurav Vaidya, Hilmar Lapp, Nico Cellinese)

TDWG

September 1-5 2018, Dunedin, New Zealand

Society of Systematic Biologists Mayr Award Symposium (with Tracy Heath)

Evolution meetings

August 22-26, 2018, Montpelier, France

SSB Spotlight Symposium: Uncertainty in the face of strong support (with Dave Weisrock)

Evolution meetings

June 23-27 2017, Portland, OR

INVITED
WORKSHOP
PARTICIPATION

National Workshop on Data Science Education

June 27-30, 2022, Berkeley, CA

Molecular Evolution Workshop at the Marine Biological Laboratory,

July 24 to August 3, 2011, Woods Hole, MA

Quantitative Evolutionary and Comparative Genomics: Linkage and Recombination in Genome Sequences, at the Okinawa Institute of Science and Technology (OIST)

May 16 to June 3, 2011, Okinawa, Japan

Presentations at

Scientific

Meetings

Linking Biodiversity Data Using Evolutionary History

Biodiversity Next

October 2019, Leiden, Netherlands

Nurturing a sustainable Open Tree of Life

TDWG September 2018, Dunedin, New Zealand

Cultivating community collaboration to build a sustainable Open Tree of Life

Evolution Meetings

August 2018, Montpelier, France

Does reference genome choice affect inference?

Bay Area Population Genomics

September 2016, San Francisco State University, CA

Continually updated phylogenies

Evolution meetings

July 2016, Austin, TX

The Open Tree of Life - Curating, synthesizing, and updating phylogenetic information across 1.8 million taxa (poster)

Society for Molecular Biology and Evolution

July 2015, Vienna, Austria

 ${\it Effects}\ of\ ascertainment\ bias\ on\ population\ genetic\ inference$

Workshop on Quantitative Evolutionary Biology

September 2014, Nesin Mathematics Village, Turkey

Phylesystem: a git-based data store for community curated phylogenetic estimates

iEvoBio June 2014, Raleigh, NC

Challenges of synthesizing divergence time information on the tree of life

iEvoBio June 2013, Snowbird, UT

Estimating population histories using single-nucleotide polymorphisms sampled throughout genomes

Evolution meetings

June 2013, Snowbird, UT

Genomic structure of introgression in New World cattle (poster)

Gordon Research Conference Februa

February 2013, Galveston, TX

Inferring ancestry of genomic regions in cattle of hybrid origin

Evolution meetings July 2012, Ottawa, ON, Canada

Tracking introgression between cattle subspecies using genomic SNP data

Evolution meetings

July 2011 Norman, OK

Using SNP data to assess population structure and hybridization in Texas Longhorn cattle

Plant and Animal Genome Conference

January 2011, San Diego, CA

Impact of dispersal and sampling on inference of population structure (poster)

BEACON: Evolution in Action

August 2010, Michigan State University, MI

Distinguishing signal from noise in phylogeographic data

Evolution meetings

July 2010, Portland, OR

Jan 2019 - 2022

Emily Jane M	cTavish	7		
Professional	Society of Systematic Biologists, Treasurer	Nov 2021 - Dec 2024		
SERVICE	UC Merced Faculty Advisory Committee on IT (FACIT) Co-Chair	2019 - present		
AND OUTREACH	UC Committee on Academic Computing and Communications (UCACC)	•		
	UCM representative	2019 - present		
	Society of Systematic Biologists, Awards committee	2017 - present		
	Society of Systematic Biologists, Communications Director	2018 - 2019		
	Society of Systematic Biologists, Elected Council Member	Jan 2016 - Dec 2018		
	JupyterCon, Program committee	March-September, 2018		
	Software Carpentry, Instructor	2011 - present		
	Breaking Bio Podcast, Guest	September 25, 2013		
	UT Forum, Lecturer	March 22, 2013		
	Darwin Day Speaker, Texas Memorial Museum,	February 12, 2012		
	Information Technology advisory committee, Student representative	Sept 2010 - Dec 2011		
	Science Under the Stars, Presenter	December 8, 2011		
	UT Learning Activities for Mature People, Lecturer	November 4, 2011		
	NPR's Science Friday, Guest	July 7, 2011		
	Flying Cloud Institute Young Women In Science Program, Teacher	July, 2010		
	UT Austin Integrative Biology Graduate Student Symposium, Chairperso	· ·		
Advising	Postdoctoral scholars			
ACTIVITIES		August 2019 - August 2023		
		eptember 2017 - May 2019		
	Doctoral student advisees (Quantitative and Systems Biology - QSB) Graduated with PhD			
	Lesly Lopez Fang	August 2023		
	Jasper Toscani-Field	October 2022		
	Active (advanced to candidacy)			
	Lucia Bazan-Williamson	Fall 2020 - present		
	Active (not yet advanced to candidacy)			
	Josue Duque	Fall 2022 - present		
	Graduate student committee membership (QSB unless otherwis	e noted)		
	Active			
	Clarence Le	March 2023 - present		
	Brooke Weinstein	August 2020 - present		
	Karly Higgins-Polling,	March 2017 - present		
	Nattanon Wutthituntisil	August 2020 - present		
	Jordan Collignon (Math)	April 2019 - present		
	Completed	1		
	Yumary Vasquez	August 2023		
	Noelle Anderson	July 2023		
	Megha Suswaram	October 2022		
	Kinsey Brock	May 2021		
	Robert Boria	May 2021		
	Danaan Deneve Weeks	December 2020		
	Travis Lawrence	June 2018		
	Jesse Wilson (ES)	August 2017		
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Editorial	Systematic Biology, Editorial board	2017 - present		
SERVICE	pyOpenSci, Guest editor Proceedings of The Poyel Society P. Associate editor	Fall 2023		
	Unaccoding at The Povel Secretar D. Aggacieto editor	102 9010 9099		

NSF Committee of Visitors (2020) Reviewer

NSF panelist (2014, 2015, 2016, 2018)

French National Research Agency, reviewer

Proceedings of The Royal Society B, Associate editor

Peer reviewer: Animal; BMC Bioinformatics; BMC Genomics; Bioinformatics; Ethology, Ecology & Evolution; Evolutionary Applications; G3:Genes-Genomes-Genetics; GigaScience; Methods in Ecology and Evolution; Molecular Biology and Evolution; Molecular Ecology; Molecular Ecology Resources; PLoSOne; PNAS; pyOpenSci; Biology; Trends in Ecology and Evolution