MICHAEL J. LANDIS

http://mlandis.github.io mlandis@gmail.com January 14, 2016

EDUCATION

Ph.D. Integrative Biology

December 2015

Designated Emphasis in Computational & Genomic Biology *University of California*, Berkeley, CA

B.S. Computer Science

July 2005

California State University, Long Beach, CA

Research Interests

- Evolutionary biology
- Statistical phylogenetics
- Bayesian inference

Publications

- **Landis, M. J.** Biogeographic dating of divergence times using paleogeographically informed processes. Systematic Biology. *Accepted*.
- Höhna, S., **Landis**, **M. J.**, and Huelsenbeck, J. P. Parallel power posterior analyses for fast computation of Bayes factors in phylogenetics. Bioinformatics. *Accepted*.
- Höhna, S., Landis, M. J., Heath, T. A., Boussau, B., Lartillot, N., Moore, B. R., Huelsenbeck, J. P, and Ronquist, F. RevBayes: Bayesian Phylogenetic Inference Using Graphical Models and an Interactive Model-Specification Language. Systematic Biology. Accepted.
- Schraiber, J. G. and Landis, M. J.. (2015) Sensitivity of quantitative traits to mutational effects and number of loci. Theoretical Population Biology. doi:10.1016/j.tpb.2015.03.005
- Höhna, S., Heath, T. A., Boussau, B., **Landis, M. J.**, Ronquist, F., and Huelsenbeck, J. P. (2014). Probabilistic graphical model representation in phylogenetics. Systematic Biology, 63(5), 753-771. doi: 10.1093/sysbio/syu039
- **Landis, M. J.** and Bedford, T. (2014). Phylowood: interactive web-based animations of biogeographic and phylogeographic histories. Bioinformatics, 30(1), 123124. doi:10.1093/bioinformatics/btt635
- Landis, M. J., Matzke, N. J., Moore, B. R., and Huelsenbeck, J. P. (2014). Bayesian Analysis of Biogeography when the Number of Areas is Large. Systematic Biology, 62(6), 789804. doi:10.1093/sysbio/syt040
- Landis*, M. J., Schraiber*, J. G., and Liang, M. (2013). Phylogenetic analysis using Lévy processes: finding jumps in the evolution of continuous traits. Systematic Biology, 62(2), 193204. doi:10.1093/sysbio/sys086

 * authors contributed equally

Presentations

- Landis, M. J. 2015. Divergence time estimation using biogeographic processes. Evolution Conference, Guarujá, Brazil.
- Landis, M. J., Matzke, N. J., Moore, B. R., Huelsenbeck, J. P. 2013. Bayesian inference of species range evolution for hundreds of discrete areas. Phylogenetics & Evolutionary Biology Seminar, North Carolina State University.
- Landis, M. J., Schraiber, J. G.. 2013. Lévy processes as models of continuous trait evolution. Workshop on Mathematics for an Evolving Biodiversity. University of Montréal.
- Landis, M. J., Schraiber, J. G., and Liang, M. 2013. Continuous trait evolution with fat tails: Lévy processes and mutations of large effect. Center for Population Genomics, UC Davis.
- Landis, M. J., Matzke, N. J., Moore, B. R., and Huelsenbeck, J. P. 2013. Bayesian inference of biogeographical histories for hundreds of discrete areas. Evolution Conference, Snowbird, Utah.

- Landis, M. J., Matzke, N. J., Moore, B. R., and Huelsenbeck, J. P. 2013. Bayesian inference of biogeographical histories for hundreds of discrete areas. Center for Theoretical Evolutionary Genomics, UC Berkeley.
- Landis, M. J., Schraiber, J. G., and Liang, M. 2012. Lévy process models of continuous character evolution. Center for Theoretical Evolutionary Genomics, UC Berkeley.
- Landis, M. J., Matzke, N. J., Moore, B. R., Huelsenbeck, J. P. 2012. Statistical Analysis of Biogeography when the Number of Areas is Large. Evolution Conference, Ottawa, Canada.

Professional reviews

Journals: Systematic Biology, Molecular Biology & Evolution, Journal of Biogeography, Bioinformatics, Methods in Evolution & Ecology, Molecular Phylogenetics & Evolution, BMC Evolutionary Biology, Genome Biology & Evolution

RESEARCH EXPERIENCE

Huelsenbeck Lab, University of California, Berkeley, CA

Jul 2009-Present

• Implemented numerous phylogenetic methods in C++, Python, R, and RevBayes with particular experience designing Bayesian methods using MCMC.

Dean Lab, Children's Hospital of Oakland, Oakland, CA

Jan 2009-Jun 2010

- Investigated the role of lateral gene transfer and recombination in the genetic diversity of *Chlamydia trachomatis* strains.
- Designed and configured a bioinformatics server to host clinical sequences and analysis tools.

TEACHING EXPERIENCE

Teaching Assistant, Molecular Evolution Workshop, Woods Hole, MA	Summer 2014, 2015
Instructor, Coordinator, RevBayes Workshop, UC Berkeley, CA	Spring 2015
Instructor, Bodega Phylogenetics Workshop, UC Davis, CA	Spring 2014, 2015
Instructor, Applied Phylogenetics Workshop, NESCent, NC	Summer 2014
Invited Lecturer, IB87, Bioinformatics, UC Berkeley, CA	Summer 2014
Invited Lecturer, EEB101, Macroevolution, UC Davis, CA	Spring 2013
Teaching Assistant, IB200A: Principles of Phylogenetics, UC Berkeley, CA	Spring 2012
Teaching Assistant, IB164: Human Genetics and Genomics, UC Berkeley, CA	Fall 2011

Professional Experience

Systems Engineer, Varsity Technologies

July 2005-October 2008

- Technical lead for Managed Services department, serving non-profits
- Designed, implemented, configured, and maintained diverse infrastructures, including: server domains, mail servers, networks, virtualization, disaster recovery.

SKILLS

Programming: C/C++, Python, R, Java, Javascript (incl. D3.js), MPI, bash, git

IT: Linux, Mac OS X, Windows, Cisco, VMWare

LEADERSHIP ACTIVITIES

Co-chair, Integrative Biology Graduate Student Assembly	Sep 2012–May 2013
Organizer, Paleostatistics Reading Group	Fall 2012
Co-organizer, Evolutionary Theory and Museum Graduate Student Group	Fall 2012
Co-organizer, PhD Student Recruitment	$Mar\ 2011 ext{-}Sep\ 2011$
Organizer, Classic Papers in Evolutionary Biology	Fall 2010

MENTORING & OUTREACH

Mentor, UC Undergraduate Research Apprentice Program

 Jaya Narasimhan, Computer Science. Comparative transcriptome evolution. Bryan Wang, Pure Mathematics. Partition distances. 	2012–2014 2013–2015
Guest Lecturer, Berkeley High School Outreach (coordinated by Rori Rohlfs)	2014-2015
Guest Lecturer, Bay Area Scientists in Schools	Spring 2014
Tutor, 826 Valencia volunteer for English Language Learners	Summer~2009–2010

RESEARCH SOFTWARE

RevBayes, phylogenetic inference using graphical models
Phylowood, biogeographic animations
mlandis.github.io/phylowood
BayArea, biogeographic inference
creepy-jerk, inference of evolutionary jumps in traits
quant_trait_coalescent, mutation-coalescent QTs
github.com/Schraiber/quant_trait_coalescent

Fellowships

Graduate Student Fellowship, National Evolutionary Synthesis CenterFall 2013Google Summer of Code, National Evolutionary Synthesis CenterSummer 2012