Brandon M. Lind, Ph.D.

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EDUCATION

Virginia Commonwealth University

Richmond, Virginia May 2018

Ph.D. Integrative Life Sciences

Advisor: Dr. Andrew Eckert

Dissertation title: Understanding Natural and Anthropogenic Drivers of Tree Evolutionary Dynamics

Iowa State University

Ames, Iowa

Graduate coursework in ecology, genetics, and plant physiology August 2010-May 2011

University of Northern Iowa Cedar Falls, Iowa

Bachelor's in Chemistry
Bachelor's in Biology
May 2010

PROFESSIONAL EXPERIENCE

Research Fellow – Ecological and Conservation Genomics, Biodiversity Informatics

April 2024 - Present

Department of Ecology & Evolutionary Biology

University of Connecticut, Connecticut, USA

Research Supervisor: Dr. Jill Wegrzyn

Research Fellow – Artificial Intelligence, Adaptation to Complex Environments January 2022 – March 2024

Department of Marine & Environmental Sciences

Northeastern University, Massachusetts, USA

Research Supervisor: Dr. Katie Lotterhos

Research Fellow – Ecological Genomics of Climate Adaptation in Conifers

June 2018 – December 2021

Centre for Forest Conservation Genetics, Department of Forest and Conservation Sciences

University of British Columbia, British Columbia, Canada

Research Supervisors: Drs. Sally Aitken, Sam Yeaman

PUBLICATIONS

- 15. (2024)# **BM Lind**, KE Lotterhos. The accuracy of predicting maladaptation to new climates with genomic data. *In revision* at *Molecular Ecology Resources*. First draft available on *bioRxiv* https://doi.org/10.1101/2024.01.30.577973
- 14. (2023) J Whiting, TR Booker, **BM Lind**, Genomics Consortium, S Yeaman. Core genes drive climate adaptation in plants. In revision at *Nature Ecology & Evolution*.
- 13. (2023)# M Lu, N Feau, **BM Lind**, P Singh, D Vidakovic, R Hamelin, SN Aitken, S Yeaman. Genetic architecture underlying response to the fungal pathogen *Dothistroma septosporum* in *Pinus contorta*, *Pinus banksiana*, and their hybrids. In revision at *Heredity*.
- 12. (2024)# **BM Lind**, R Candido-Ribeiro, P Singh, M Lu, D Vidakovic, T Booker, M Whitlock, N Isabel, S Yeaman, SN Aitken. How useful is genomic data for predicting maladaptation to future climate? *Global Change Biology*. DOI: https://doi.org/10.1111/gcb.17227
- 11. (2024)# P Singh, B St Clair, **BM Lind**, R Cronn, NP Wilhelmi, M Lu, D Vidakovic, R Hamelin, D Shaw, SN Aitken, S Yeaman Genomic architecture of resistance and tolerance to Swiss needle cast and Rhabdocline needle cast diseases in Douglas-fir. New Phytologist. DOI: https://doi.org/10.1111/nph.19797

- 10. (2022) R Jasper, TK McDonald, P Singh, M Lu, C Rougeux, **BM Lind**, S Yeaman. Evaluating the accuracy of variant calling methods using the frequency of parent-offspring genotype mismatch. Molecular Ecology Resources (22) 2254-2533. DOI: https://doi.org/10.1111/1755-0998.13628
- 09. (2022)# **BM Lind***, M Lu*, D Vidakovic, P Singh, T Booker, SN Aitken, S Yeaman. Haploid, diploid, and pooled exome capture recapitulate features of biology and paralogy in two non-model tree species. Molecular Ecology Resources (00) 1-14. DOI: https://doi.org/10.1111/1755-0998.13474
- 08. (2021) IR MacLachlan, TK McDonald, **BM Lind**, LH Rieseberg, S Yeaman, SN Aitken. Genome-wide shifts in climate-related variation underpin responses to selective breeding in a widespread conifer. Proceedings of the National Academy of Science 18(10) e2016900118. https://doi.org/10.1073/pnas.2016900118
- 07. (2020) CR Mahoney, IR MacLachlan, **BM Lind**, JB Yoder, T Wang, SN Aitken. Evaluating genomic data for management of local adaptation in a changing climate: A lodgepole pine case study. Evolutionary Applications 00:1-16. DOI: https://doi.org/10.1111/eva.12871
- 06. (2019) CJ Friedline, TM Faske, **BM Lind**, EM Hobson†, P Dylan†, R Dyer, D Johnson, L Thompson, K Grayson, AJ Eckert. Evolutionary genomics of gypsy moth populations sampled along a latitudinal gradient. Molecular Ecology 28:2206-2223. DOI: https://doi.org/10.1111/mec.15069
- 05. (2019)# **BM Lind**, MP North, PE Maloney, AJ Eckert. Effect of fire and thinning on fine-scale genetic structure and gene flow in fire-suppressed populations of sugar pine (Pinus lambertiana Douglas). Forest Ecology and Management 447:115-129. DOI: https://doi.org/10.1101/448522
- 04. INVITED (2018)# **BM Lind**, M Mennon, CE Bolte, TM Faske, AJ Eckert. The genomics of local adaptation in trees: Are we out of the woods yet? Tree Genetics & Genomes 14: 29. DOI: https://doi.org/10.1007/s11295-017-1224-y
- 03. (2017)# **BM Lind**, CJ Friedline, JL Wegrzyn, PE Maloney, DR Vogler, DB Neale, AJ Eckert. Water availability drives signatures of local adaptation in whitebark pine (*Pinus albicaulis*) across fine spatial scales of the Lake Tahoe Basin, USA. Molecular Ecology 26: 3168-3185. DOI: http://dx.doi.org/10.1111/mec.14106
- 02. (2016) AJ Eckert, DE Harwood, **BM Lind**, EM Hobson†, A Mix, PE Maloney, CJ Friedline. The genetic architecture of local adaptation II: The QTL landscape of water-use efficiency for foxtail pine (*Pinus balfouriana* Grev. & Balf.). https://doi.org/10.1101/038240
- 01. (2015) CJ Friedline, **BM Lind**, EM Hobson†, DE Harwood, A Mix, PE Maloney, AJ. The genetic architecture of local adaptation I: The genomic landscape of foxtail pine (*Pinus balfouriana* Grev. & Balf.) as revealed from a high-density linkage map. Tree Genetics & Genomes 11:1-15. DOI: http://dx.doi.org/10.1007/s11295-015-0866-x

PRESENTATIONS

Keynote Speaker

18. (Sept 2023) **Lind BM** The limits of predicting maladaptation to future climates with genomic data. EVOLTREE conference. Transilvania University of Braşov, Braşov, Romania.

Invited

- 17. (Jan 2024) **Lind BM** How useful is genetic data for species management? Lessons from Forest Trees. Department seminar, Nicholas School of the Environment, Duke University.
- 16. (Dec 2022) **Lind BM** Leveraging ecological genomics to inform species management. Department seminar, Department of Marine & Environmental Sciences, Northeastern University.
- 15. (Sept 2021) **Lind BM**, R Candido-Ribeiro, P Singh, M Lu, D Vidakovic, T Booker, M Whitlock, N Isabel, S Yeaman, SN Aitken. How off are genetic offset predictions? Lessons from common gardens and three clades of conifers. EVOLTREE conference. Virtual.
- 14. (Jan 2020) **Lind BM**; P Singh, S Yeaman, SN Aitken. Convergence of Polygenic Climate Adaptation in Conifers. Tree Genetics Workshop, Plant & Animal Genome conference. San Diego, California.

- 13. (Jan 2019) Lind BM; M Lu, D Vidakovic, S Yeaman, SN Aitken. An efficient method to genotype species with large and complex genomes. Tree Genetics Workshop, Plant & Animal Genome conference. San Diego, California.
- 12. (Oct 2016) **Lind BM**; CJ Friedline, JL Wegrzyn, PE Maloney, DR Vogler, CE Jensen, AD Mix, DB Neale, and AJ Eckert. Local adaptation of three white pine species across fine spatial scales of the Lake Tahoe Basin, USA: Implications to sustainable management and conservation of conifer species. Natural Areas conference. Davis, California.

Contributed

- 11. (July 2024) **Lind BM**, KE Lotterhos. Prospects and limitations of predicting maladaptation to changing environments. Evolution Conference.
- 10. (June 2024) **Lind BM**, L Rettenbacher, SN Aitken. Quantitative and population genetics of North American and European lineages of coastal Douglas-fir. North American Forest Genetics Association Conference. Oaxaca, Mexico.
- 09. (June 2024) **Lind BM**, M Myles, I Harding, J Gamer, R Wolther, JL Wegrzyn, I Cobo-Simon, R Ramanth, E Grau, V Savitsky, G Barrett, N Herndon, M Staton. North American Forest Genetics Association Conference. Oaxaca, Mexico. Poster.
- 08. (May 2021) **Lind BM**, S Yeaman, SN Aitken. Patterns and climate drivers of adaptive variation in Douglas-fir. Western Forest Genetics Association conference. Virtual.
- 07. (June 2019) **Lind BM**, M Lu, D Obreht Vidakovic, S Yeaman, SN Aitken. Combining exome capture and poolseq: Lessons from three conifer species. Evolution conference. Providence, RI.
- 06. (June 2017) **Lind BM**, M North, AJ Eckert. Time to tree-think the genetics effects of forest management: Assessing patterns and processes beyond heterozygosity indices. Evolution conference. Portland, OR
- 05. (Feb 2017) **Lind BM**; M Menon, C Bolte, T Faske, AJ Eckert. The genomics of local adaptation in trees: Are we out of the woods yet? ILS Showcase. Richmond, Virginia.
- 04. (June 2016) **Lind BM**; CJ Friedline, JL Wegrzyn, PE Maloney, DR Vogler, DB Neale, and AJ Eckert. When local means local: polygenic signatures of local adaptation within whitebark pine (*Pinus albicaulis* Engelm.) across the Lake Tahoe Basin, USA. Evolution conference. Austin, Tx.
- 03. (June 2015) **Lind BM**; PE Maloney, DR Vogler, DB Neale, and AJ Eckert. Genetic architecture of fitness-related traits in whitebark pine (*Pinus albicaulis* Englm.). 35th New Phytologist Symposium The Genomes of Forest Trees: New Frontiers in Forest Biology. Harvard University. Boston, MA. Poster.
- 02. (Nov 2014) Friedline CJ; **BM Lind**, EM Hobson, DE Harwood, AD Mix, PE Maloney, and AJ Eckert. Local adaptation and linkage maps a first step towards the genomic architecture of fitness-related phenotypes in natural populations. VCU Integrative Life Science Ph.D. Showcase. Richmond, VA. Poster.
- 01. (June 2014) Lind BM; EM Hobson, MP North, AJ Eckert. Proposal: Genetic effects of factorial fire and thinning in fire-suppressed populations of Jeffrey and sugar pine. Evolution conference. Raleigh, NC. Poster.

TEACHING

Instructor of Record

2024 Biodiversity and Conservation Genomics Training program

Storrs, CT

Lecture Series

2023	Logistics for Project Informatics: From File Systems to Project API	Nahant, MA
2023	Version control, git, and GitHub	Nahant, MA
2023	Primer in Geographic Information Systems	Nahant, MA
2022	Bioinformagics: Tips, tricks, and best practices for productive and reproducible research	Nahant, MA

Guest Lecturer

2022 Reproducibility and Publication Bias (in: Intro to environmental, social, and biological data) Nahant, MA

2018	Hardy-Weinberg equilibrium (in: Evolution)	Richmond, VA
2017	Hardy-Weinberg equilibrium (in: Evolution)	Richmond, VA
2016	Null hypotheses in evolutionary biology (in: Evolution)	Richmond, VA
2015	Population differentiation/genetics, F_{ST} (in: Evolution)	Richmond, VA

Teaching Assistant

2014-2017 Introduction to Biology Laboratory

Richmond, VA

- When I began teaching the laboratory sections for this course, there were no teaching materials to convey information to students, and there was no link between hypothesis testing and their midterm or final experiments for the course
- When I ended my assistantship, I left with the course organizers Power Point slides for each lab's lecture portion, quizzes for each lab exercise, and developed a drag-and-drop statistical analysis in R so that students could incorporate hypothesis testing (support for null or alternative hypothesis) into their experimental results and discussions
- The drag-and-drop analysis was customizable with custom error messages so that it would prompt students for sample sizes and measurement type to check their data for errors and to ensure the correct statistical test was run consistently across lab sections, without the need for entry-level students to understand how to use R and instead just how to drag and drop the script file into R and hit enter, which would automatically find and run analysis on their data file

CONTRIBUTIONS TO DIVERSITY, EQUITY, INCLUSION, BELONGING, AND MENTORSHIP

Mentorship (ongoing)

†undergraduate

Student mentees

- (2024-present) Andrew Deierleint, Sia Gbondo, Adam Glendeningt, Samira Obbu, Alan Roger
- (2018-2024) Rafa Candido-Ribeiro, Beth Roskilly, Madeline Eppley, Sarit Truskey, Camille Rumberger, Kiran Bajaj†, Elisabeth Leung†, Nicole Mongillo†, Clara Winguth†
- (2012-2018) Britta Austin†; Shelby Bennett†; Ayana Ferrer†; Casey Harless†; Erin Hobson†; Angela Simpson†; Rachel Stevenson†; Alexandrea Stylianou†; Tammy Tran†; Hugo Vargas Cruz†

VCU Graduate Student Mentorship Program (2013 – 2016)

Paired with and mentored three undergraduate students planning on applying to graduate school within a formal
mentoring program at Virginia Commonwealth University. Paired for one academic year. All three of my
mentees have been accepted to graduate school ranging from medical school to academic research in a Ph.D.
program.

Assisted in crafting Land Acknowledgment for Nahant campus of Northeastern University (2023)

UBC Forestry Diversity Crew member (2020 – 2021)

VCU Rice Rivers Center outreach weekend (2012)

 Assisted in and helped organize educational weekend at VCU nature center for the Virginia School for the Deaf and Blind

PHILANTHROPY AND OUTREACH

Skype a Scientist (2017 – 2022, 2024)

• I have been matched with various elementary, middle, and high school classrooms across the United States to help communicate science. These meetings are structured so that students can ask questions about my research as well as about science careers in general. It is very rewarding to engage students in their formative years.

St. Jude Children's Research Hospital's "Up 'til Dawn", University of Northern Iowa (2006-2008) Founder / Director UNI Chapter – Cedar Falls, IA

- Contacted St. Jude regional office and began collaboration for new Up 'til Dawn chapter
- Arranged campus wide informational meetings to recruit, review, and select members of the executive board
- Organized, planned and ran weekly meetings for the executive board as well as organization's budget
- Raised >\$23,000 the inaugural year, and >\$100,000 over the chapter's lifetime

Other Philanthropy (2004-2008)

(Co)Founder – Cedar Falls, IA

- Relay for Life Benefit Concert 'Rock-4-a-Cure' on Main Street, \$1,600 annual average
- American Diabetes Association Dodge Ball Tournament, \$500 annual average
- Valentine's Day registration tables for the Iowa Donor Network 'Truly Give Away Your Heart'
- Organized and volunteered at annual volunteer efforts for Big Brothers Big Sisters, Highway Cleanup, Habitat for Humanity, Special Olympics as well as the flood relief efforts in Black Hawk County, Iowa

SERVICE

Data Editor – The American Naturalist (2021 – 2023)

Responsible for assessing code and data archives for manuscripts accepted to The American Naturalist

Journal Peer Review

Molecular Ecology; Heredity; New Phytologist; Plant Ecology & Diversity; Conservation Genetics;
 Global Change Biology; G3 Genes | Genetics | Genomes; Forest Ecology & Management; Ecology & Evolution; BMC Ecology & Evolution

Society member

 American Genetic Association, Society for the Study of Evolution, Natural Areas Association, Society of Molecular Biology & Evolution

Poster Judge – BIOL 318 University of British Columbia (April 2018)

Virginia Junior Academy of Science judge (2013 – 2014)

Journal club - Assistant to the regional manager (2012 – 2018):

 Helped organize and facilitate Evolution Fridays (EvolFri) weekly reading group at Virginia Commonwealth University

AWARDS & FUNDING

•	Nominated – IUFRO Excellence in Forest Molecular Biology and Genomics	2022
•	VCU Graduate School Dissertation Fellowship - academic year \$25,000	2017-2018
•	VCU College of Humanities and Sciences Outstanding Ph.D. student	2016-2017
•	Integrative Life Science Travel Award \$1,000 USD	2016

New Phytologist Trust Student Travel Award \$500 USD
 Integrative Life Science Travel Award \$1,000 USD

VCU Graduate School Research Fellowship - 12 month
 VCU Graduate School Research Fellowship - 12 month
 2013-2014
 2012-2013

PUBLIC CODING AND GENETIC RESOURCES

Analysis code from previous and current research:

- 12. Lind BM. (2024a) GitHub.com/brandonlind/offset_validation: Publication release (Version 1.1.0). Zenodo. https://doi.org/10.5281/zenodo.7641225 (see Publication 12)
- 11. Lind BM. (2024b) GitHub.com/ModelValidationProgram/MVP-offsets: Revision release (Version 1.0.1). Zenodo. https://zenodo.org/doi/10.5281/zenodo.10611682 (see Publication 15)
- 10. Lind BM. (2023a). GitHub.com/brandonlind/jack_pine_natural_populations: Offset Revision 1 (v1.0.0). Zenodo. https://doi.org/10.5281/zenodo.8018892 (see Publication 12)
- 09. Lind BM. (2023b). GitHub.com/brandonlind/douglas_fir_natural_populations: Offset Revision 1 (v1.0.0). Zenodo. https://doi.org/10.5281/zenodo.8018894 (see Publication 12)
- 08. Lind BM. (2021a) GitHub.com/CoAdapTree/gatk_pipeline: Publication release (Version 1.0.0). Zenodo. http://doi.org/10.5281/zenodo.5083321 (see Publication 09)
- 07. Lind BM. (2021b) GitHub.com/CoAdaptree/varscan_pipeline: Publication release (Version 1.0.0). Zenodo. http://doi.org/10.5281/zenodo.5083302 (see Publication 09)
- 06. Lind BM. (2021c) GitHub.com/CoAdaptree/testdata_validation: Publication release (Version 1.0.0). Zenodo. http://doi.org/10.5281/zenodo.5083292 (see Publication 09)
- 05. Lind BM. (2021d) GitHub.com/CoAdaptree/cmh_test: Preprint release (Version 1.0.0). Zenodo. http://doi.org/10.5281/zenodo.5083798 (see Publications 11, 13)
- 04. Lind BM. (2019) GitHub.com/brandonlind/teakettle (see Publication 05)
- 03. Lind BM. (2018) GitHub.com/brandonlind/out_of_the_woods (see Publication 04)
- 02. Lind BM (2017) GitHub.com/brandonlind/whitebark_pine (see Publication 03)

Foxtail pine genetic linkage map:

01. Tree Genes Comparative Map Database, Accession Number: TG151

TECHNICAL EXPERIENCE

Teakettle Experimental Forest

Field Technician

Sierra National Forest, CA June 2011 – August 2011

- Working unsupervised, measured ecological attributes for a project examining trade-offs of management practices' impact on forest dynamics via the inclusion and exclusion of Rx fire as well as thinning treatments
- Collected and individual-level data (DBH, species, decay/disease class)
- Used knowledge of flora for identification and records of forest plants, using dichotomous key-outs as needed
- Surveyed forest spatial inventory using offset GPS (laser station/Trimble)

Spatially Maximized Arrays of DNA Nanofibers, Iowa State University

Ames, IA

National Science Foundation Research Experiences for Undergraduates (NSF REU)

June 2010-August 2010

- Independently addressed objective to geometrically align DNA nanofibers for future use as a scaffolding for the spontaneous alignment of organized arrays of metallic nanowires for bottoms-up fabrication of electronics
- Through personally devised original methods based upon confined evaporative self-assembly (CESA), spatial
 area over which arrays of parallel DNA nanofibers was increased by a full order of magnitude over that of
 previous research in the literature, resulting in millimeter-scale nanofibers
- Formulated methods reduced cost, effort and timescale of fabrication
- Senior PI acknowledges I earned authorship for a subsequent publication in ACS Nano

OTHER WORK EXPERIENCE

Crop Genome Informatics Laboratory (CGIL), Iowa State Research Reference Librarian

Ames, IA October 2010–December 2010