# **Silas Tittes**

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## **EDUCATION University of Colorado at Boulder**

■ PhD. – Ecology and Evolutionary Biology

Aug 2014 – May 2019

• Dissertation: Predicting evolution and inferring its consequences

■ Bachelor of Arts – Ecology and Evolutionary Biology

May 2008 - Dec 2012

• Honors Thesis (*Magna Cum Laude*): Flea genetic diversity in Gunnison's Prairie Dog colonies and its implications for flea transmitted diseases

#### RESEARCH EXPERIENCE

## **University of Oregon**

Population Geneticist, Institute of Ecology and Evolution

May 2022 - present

#### **University of California Davis**

Postdoctoral Researcher, Department of Evolution and Ecology

May 2019 – May 2022

#### **University of Colorado at Boulder**

Professional Research Assistant, Ecology and Evolutionary Biology

Aug 2014 – Aug 2013

Undergraduate Research Student, Ecology and Evolutionary Biology

May 2009 - Aug 2012

#### **SKILLS**

R, R Markdown, R Shiny, Bash, Python, HPC, Snakemake, Nextflow, SLiM, Stan, Keras, Docker, Pandoc, Late, Stan, GitHub, and Googling Stackoverflow.

## AWARDS & SCHOLARSHIPS

Excellence in Teaching Award
 CU Boulder, Ecology and Evolutionary Biology

2019

■ Ling-Ju Harn Fellowship

2014

\$18,000

• Undergraduate Research Opportunities Program

2010

\$1,000

2008

 Edith Scates Memorial Scholarship \$1,000

2008

 Lion's Club International Scholarship \$500

#### **SOFTWARE**

Python package: mop

Simple tool for capturing alignment regions with sufficient quality for genotyping.

R package: rdmc

Implements convergent adaptation models of Lee and Coop (2017).

■ R package: performr

A probablistic Hierarchical Bayesian model to predict performance curves across multiple groups (i.e. lines, populations, taxa, etc.).

R package: pomodoror
 A writing productivity application.

#### **TEACHING**

#### **INSTRUCTOR**

Apple Genomics, CU Boulder

Spring 2018

I designed and taught Apple Genomics as an upper-division undergraduate elective course focused on the assessment of genetic diversity and classification of Boulder county apples trees. Starting from leaf samples students learned DNA extraction and QC; the development of custom pipelines for DNA sequence alignment, genotyping, and calling variants; and the use of several software packages to assess population structure and genetic diversity. This project is ongoing and is now led by several of the students that took the course.

### TEACHING ASSISTANT

Phylogenetics / Comparative Biology, CU Boulder

Spring 2019

Evolutionary Biology, CU Boulder

Fall 2016 - 2018, Spring 2017, Summer 2017

Genetics, CU Boulder

Spring 2015, Summer 2015, Spring 2016

#### Genomics, CU Boulder

#### COURSE DEVELOPMENT

Population Genetics web applications
 I developed a series of R shiny based web applications used to teach undergraduate fundamental concepts in Population Genetics. These are free to use and available on <u>GitHub</u>, and are used regularly in several population genetics courses in several countries.

## HACKATHONS, WORKSHOPS, AND MEETUPS

#### **HACKATHONS**

- Ross-Ibarra Lab Snakefest (organizer), UC Davis
   I organized a one day hackathon to develop a Snakemake based pipeline for aligning short reads, calling and filtering SNPs.
- PanAnd (attendant), Cornell University June 2019 I attended a five day hackathon hosted by the Bucker lab at Cornell University. I explored methods to identify conserved non-coding regions using multi-species alignments of DNA shape statistics.

#### WORKSHOPS

#### MEETUPS

Meetup: R-ladies Boulder Meetup (attendant), Boulder, CO Monthly (starting September 2019)
 I attend the monthly R-ladies Boulder Meetup to practice, learn, and develop new R language skills in an inclusive environment.

#### PUBLICATIONS

#### **PREPRINTS**

- [3] M Elise Lauterbur, Mariz Izabel A Cavassim, ..., S Tittes, ..., Ilan Gronau. Expanding the stdpopsim species catalog, and lessons learned for realistic genome simulations (2022) *bioRxiv*.
- [2] CCR Smith, <u>S Tittes</u>, PL Ralph, AD Kern. Dispersal inference from population genetic variation using a convolutional neural network (2022). *bioRxiv*.
- [1] S Tittes, A Lorant, S McGinty, J Doebley, J Holland, J Sanchez-Gonzalez, A Seetharam, M Tenaillon, J Ross-Ibarra. Not so local: the population genetics of convergent adaptation in maize and teosinte (2021). *bioRxiv*.

#### **JOURNALS**

- [16] MB Hufford, A Seetharam, ..., <u>S Tittes</u>, ..., RK Dawe. De novo assembly, annotation, and comparative analysis of 26 diverse maize genomes (2021). *Science*.
- [15] NB O'Hara, SJ Franks, NC Kane, <u>S Tittes</u>, JS Rest. Evolution of pathogen response genes associated with increased disease susceptibility during adaptation to an extreme drought event in a *Brassica rapa* plant population (2021). *BMC Ecology and Evolution*.
- [14] <u>S Tittes</u>. rdmc: an open source R package implementing convergent adaptation models of Lee and Coop (2017). (2020). *G*3.
- [13] R Wooliver, <u>S Tittes</u>, SN Sheth. A resurrection study reveals limited evolution of thermal performance in response to recent climate change across the geographic range of the scarlet monkeyflower. (2020). *Evolution*.
- [12] AA Comeault, J Wang, <u>S Tittes</u>, K Isbell, S Ingley, AH Hurlbert, DR Matute. Genetic diversity and thermal performance in invasive and native populations of African fig flies. (2019). *Molecular Biology and Evolution*.
- [11] K Carscadden, M Mcdermott S Turbek, <u>S Tittes</u>, AP Martin. Building bridges: An active learning lesson in evolution and collaboration. (2019). *Journal of College Science Teaching*.
- [10] C Weiss-Lehman, <u>S Tittes</u>, NC Kane, R Hufbauer, BA Melbourne. Riding the wave: genomic signatures of gene surfing and selection in experimental range expansions. (2019). *Philosophical Transactions of the Royal Society B*.
- [9] <u>S Tittes</u>, JF Walker, L Torres-Martinez, NC Emery. Grow where you thrive, or where only you can survive? An analysis of tolerance curve evolution in a clade with diverse habitat affinities. (2019). *The American Naturalist*.
- [8] CS Smith, E Scordato, <u>S Tittes</u>, S Taylor, D Vergara. Book Review: Molecular Population Genetics. Matthew Hahn. (2019). *Molecular Ecology*.

- [7] CS Smith, <u>S Tittes</u>, JP Mendieta, E Collier-zans, H Rowe, LH Rieseberg, NC Kane. (2018). Genetics of alternative splicing evolution during sunflower domestication. *Proceedings of the National Academy of Sciences*.
- [6] Q Gao, NC Kane, B Hulke, S Reinert, C Pogoda, <u>S Tittes</u>, J Prasifka. (2017). Genetic architecture of capitate glandular trichome density in florets of domesticated sunflower (*Helianthus annuus L*.). *Frontiers in plant science*.
- [5] DJ Gray, H Baker, K Clancy, RC Clarke, K deCesare, J Fike, MJ Gibbs, F Grotenhermen, NC Kane, KG Keepers, DP Land, RC Lynch, JP Mendieta, M Merlin, K Muller-Vahl, CS Pauli, BJ Pearson, B Rhan, TC Ruthenberg, CJ Schwartz, <u>S Tittes</u>, D Vergara, KH White, RN Trigiano. (2016). Current and future needs and applications for *Cannabis*. *Critical Reviews in Plant Sciences*.
- [4] D Vergara, H Baker, K Clancy, KG Keepers, JP Mendieta, CS Pauli, <u>S Tittes</u>, KH White, NC Kane. (2016). Genetic and genomic tools for *Cannabis sativa*. *Critical Reviews in Plant Sciences*.
- [3] RC Lynch, D Vergara, <u>S Tittes</u>, KH White, CJ Schwartz, MJ Gibbs, TC Ruthenburg, K deCesare, DP Land, NC Kane. (2016). Genomic and chemical diversity in *Cannabis*. *Critical Reviews in Plant Sciences*.
- [2] SJ Franks, NC Kane, NB O'Hara, <u>S Tittes</u>, JS Rest. (2016). Rapid genome-wide evolution in *Brassica rapa* populations following drought revealed by sequencing of ancestral and descendant gene pools. *Molecular Ecology*.
- [1] <u>S Tittes</u>, NC Kane. (2014). The genomics of adaptation, divergence and speciation: a congealing theory. *Molecular Ecology*.

#### CONFERENCES

- [6] <u>S Tittes</u>. (2020). Evolutionary genetics of lots of grass. *UC Davis, Center for Population Biology, Seminar*, remote
- [5] <u>S Tittes</u>. (2020). Local Adaptation in Maize landrace and *parviglumis* populations. *Zeavolution*, remote
- [4] <u>S Tittes</u>, NC Emery. (2018). A novel Bayesian inference method to model tolerance curves. *The American Society of Naturalists*, Montrey, CA
- [3] <u>S Tittes</u> C Weiss-Lehman, NC Kane, R Hufbauer, BA Melbourne. (2017). Surfing in pools of beetles: using replicated landscape experiments to disentangle signatures of selection and drift. *Evolution*, Portland, OR
- [2] NB O'Hara, SJ Franks, NC Kane, <u>S Tittes</u>, Amidi-Abraham G, JS Rest. (2014). Genomic signatures of rapid evolution in drought response and disease susceptibility in an annual plant, *Brassica rapa*. *Society for Molecular Biology and Evolution*, Puerto Rico
- [1] SJ Franks, NC Kane, NB O'Hara, <u>S Tittes</u>, JS Rest. (2014). Genome-wide analysis reveals rapid genetic changes in natural *Brassica rapa* populations following drought. *Evolution*, Raleigh, North Carolina

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