Contact	Department of Biology Tufts University Robinson Hall Medford, MA 02155 (310) - 498 - 8608, Lawrence. Uricchio@tufts.edu		https://scholar.google.com/citations?user=XyVUpZYAAAAJ https://uricchio.github.io/	
Academic	2021 -	Assistant Professor, Tufts Universi	2, 2	
positions	2019 - 2021	Postdoc, UC Berkeley, Department of Integrative Biology		
	2015 - 2018	Postdoc, Stanford University, Depa	rtment of Biology	
Education	2014	PhD, Bioinformatics, University of California, San Francisco		
Education	2011	MS, Computer Science, University	•	
	2009	MS, Biophysical Sciences, University	9	
	2005	BA, Physics, Carleton College	ty of Chicago	
	2000	DA, I hysics, Carleton Conege		
Honors &	2016 - 2018	NIH IRACDA Postdoctoral Fellow	Stanford University	
Fellowships	2015 - 2016	CEHG Postdoctoral Fellow, Stanfo	rd University	
	2014	Discovery Fellow, UCSF		
	2014	ASHG trainee award for excellence	in human genetics research, semi-finalist	
	2013	Teaching Assistant Excellence Awa	rd, UCSF	
	2012 - 2014	Achievement Rewards for College S	Scientists Fellow, UCSF	
	2005	Distinction in the physics major, C	arleton College	
	2005	Phi Beta Kappa (academic honor s	ociety), Carleton College	
	2005	Sigma Xi (scientific research honor	society), Carleton College	
	2005	Magna Cum Laude, Carleton Colle	ge	
	2002	Dean's list, Carleton College		

Publications

Articles & editorials

- URICCHIO LH[†]. Evolutionary perspectives on polygenic selection, missing heritability, and GWAS. Human Genetics, 139(1):5-21, Jan 2020.
- 2. Hernandez RD, Uricchio LH, Hartman K, Ye J, Dahl A, Zaitlen N. Ultrarare variants drive substantial cisheritability of human gene expression. *Nature Genetics*, 51:1349–1355, Sep 2019.
- 3. URICCHIO LH[†], Petrov DA, Enard D[†]. Exploiting selection at linked stites to infer the rate and strength of adaptation.

 Nature Ecology & Evolution, 3:977–984, June 2019.
- 4. Severson AL[‡], Uricchio LH[‡], Arbisser IM[‡], Glassberg EC, Rosenberg NA. Analysis of author gender in TPB, 1991–2018. Theoretical Population Biology, 127:1–6, June 2019.
- 5. Gignoux CR, Torgerson DG, Pino-Yanes M, Uricchio LH, Galanter J et al. An admixture mapping metaanalysis implicates genetic variation at 18q21 with asthma susceptibility in latinos. *Journal of Allergy and Clinical Immunol*ogy, 143(3):957–969, Mar 2019.
- 6. URICCHIO LH^{‡†}, KITANO HC[‡], GUSEV A, ZAITLEN NA[†]. An evolutionary compass for detecting polygenic selection and mutational bias. *Evolution Letters*, 3(1):69–79, Feb 2019.
- 7. URICCHIO LH[†], DAWS SC, SPEAR ER, MORDECAI EA[†]. Priority effects and non-hierarchical competition shape species composition in a complex grassland community. *The American Naturalist*, 193(2):213–226, Feb 2019.
- 8. Goldberg A, Uricchio LH, Rosenberg NA. Natural selection in human populations. Oxford Bibliographies in Evolutionary Biology, Aug 2018.
- 9. URICCHIO LH, WARNOW T, ROSENBERG NA. An analytical upper bound on the number of loci required for all splits of a species tree to appear in a set of gene trees. *BMC Bioinformatics*, 17(14):241–250, Nov 2016.
- 10. URICCHIO LH[†], ZAITLEN NA, YE CJ, WITTE JS, HERNANDEZ RD[†]. Selection and explosive growth alter genetic architecture and hamper the detection of causal rare variants. *Genome Research*, 26:863–873, July 2016.
- 11. URICCHIO LH, TORRES R, WITTE JS, HERNANDEZ RD. Population genetic simulations of complex phenotypes with implications for rare variant association tests. *Genetic Epidemiology*, 39(1):35–44, Jan 2015.

- 12. URICCHIO LH, HERNANDEZ RD. Robust forward simulations of recurrent hitchhiking. Genetics, 197(1):221–236, May 2014.
- 13. Maher MC[‡], **Uricchio LH**[‡], Torgerson DG, Hernandez RD. Population genetics of rare variants and complex diseases. *Human Heredity*, 74(3-4):118–128, Apr 2013.
- 14. URICCHIO LH, CHONG JX, ROSS KD, OBER C, NICOLAE DL. Accurate imputation of rare and common variants in a founder population from a small number of sequenced individuals. *Genetic Epidemiology*, 36(4):312–319, May 2012.
- 15. ÇALIŞKAN M, CHONG JX, **URICCHIO L**, ANDERSON R, CHEN P *et al.* Exome sequencing reveals a novel mutation for autosomal recessive nonsyndromic mental retardation in the TECR gene on chromosome 19p13. *Human Molecular Genetics*, 20(7):1285–1289, Apr 2011.
- 16. MILLER LS, PIETRAS EM, **URICCHIO LH**, HIRANO K, RAO S et al. Inflammasome-mediated production of IL-1 β is required for neutrophil recruitment against staphylococcus aureus in vivo. *Journal of Immunology*, 179(10):6933–6942, Nov 2007.
- 17. PATTANAYAK AK, BROOKS DWC, DE LA FUENTE A, URICCHIO L, HOLBY E et al. Coarse-grained entropy decrease and phase-space focusing in hamiltonian dynamics. Physical Review A, 72(1):013406, Jul 2005.

‡ denotes equal contributions; † denotes corresponding author

In progress

- 1. Castellano D, Uricchio LH, Munch K, Enard D. Viruses rule over adaptation in conserved human proteins. to be submitted: preprint at https://www.biorxiv.org/content/10.1101/555060v1, 2019.
- 2. COUPER L, FARNER J, CALDWELL J, CHILDS M, HARRIS M, KIRK D, NOVA N, SHOCKET M, SKINNER E, URICCHIO L, EXPOSITO-ALONSO M, MORDECAI E. How will mosquitoes adapt to climate change? Under review: preprint at https://www.authorea.com/users/330221/articles/494182-how-will-..., 2020.
- 3. Song C, Uricchio LH, Mordecai EA, Saavedra S. Understanding the emergence of contingent and deterministic exclusion in multispecies communities. Under review: preprint at https://www.biorxiv.org/content/10.1101/2020.09.23.310524v2.full.pdf, 2020.

Dissertation

1. URICCHIO LH. Models and forward simulations of selection, human demography, and complex traits. UNIVERSITY OF CALIFORNIA, SAN FRANCISCO, 2014.

Teaching	2018	Co-instructor, Evolutionary genetics (BIOL 118), San Jose State University*	
	2017	Co-instructor, Ecology (BIOL 160), San Jose State University*	
		st semester-long undergraduate courses for which I led 40-50% of instructional time.	
		Co-founder, Stanford postdoc pedagogy miniseries, Stanford University	
		Instructor, Undergraduate Biology Exploration, Stanford University	
	2016-2017 Guest lecturer (twice), Evolution, Stanford University 2014 Graduate teaching assistant, Computational Evolutionary Genomics, UCSF		
	2013	Graduate student instructor, Computational Biology, UC Berkeley	
	2009	Guest lecturer, Genes, Networks, and Cells, University of Chicago	
	2003-2004	Teaching assistant, tutor, & grader, Introduction to Physics, Classical & Computational	
		Mechanics, Contemporary Experimental Physics, Carleton College	
Mentoring	2016	Student research co-mentor to Alan Aw, Rosenberg Lab	
	2013	Student research mentor to Isela Hernandez, Biological Health Sciences Internship Program	
	2010	Student research mentor to Sam Neal, Summer Link High School Program	
	2006-2007	$\hbox{Co-supervisor/trainer of undergraduate lab members, UCLA}$	

Service	2013-	Invited reviewer for Genetics, Nature Genetics, BMC Evolutionary Biology, PLoS Genetics Nature Ecology & Evolution, Molecular Biology & Evolution, Genome Biology & Evolution Molecular Ecology, Heredity, Theoretical Population Biology, PLoS ONE, G3,IEEE/ACM Transactions on Computational Biology and Bioinformatics, Functional Ecology, Peer Communities in Evolutionary Biology, American Journal of Human Genetics, and the Journal of Allergy and Clinical Immunology,	
	2016	Committee member & session leader, Stanford Postdoc Pedagogy Journal Club	
	2015-2016	Committee member, Stanford CEHG diversity outreach committee	
Competitive Funding	2016-2018 2016-2017 2015-2016 2012-2014 2014	NIH IRACDA Fellow, Stanford & SJSU (\$53,600 per year) Stanford Teaching & Mentoring Academy Award (\$6,870) Stanford CEHG Postdoctoral Fellowship (\$50,000) UCSF Achievement Rewards for College Scientists Fellow (\$12,000 per year) UCSF Discovery Fellow (\$4,000 for travel & research over 2 years)	

Presentations

- 1. Back to the future: the population genetics of adaptation in of our rapidly changing world. CSU East Bay, Hayward, CA. Seminar, 2020.
- 2. Adaptation and constraint through the lens of genome-wide association studies. *University of Wisconsin, Madison, WI.* Seminar, 2020.
- 3. Adaptation and constraint through the lens of genome-wide association studies. Temple University, Philadelphia, PA. Seminar, 2020.
- 4. Adaptation and constraint through the lens of genome-wide association studies. *Tufts University, Somerville, MA*. **Seminar**, 2020.
- 5. Adaptation and constraint through the lens of genome-wide association studies. Syracuse University, Syracuse, NY. Seminar, 2020.
- 6. Adaptation and constraint through the lens of genome-wide association studies. Vanderbilt University, Nashville, TN. Seminar, 2020.
- 7. Evolutionary perspectives on human complex traits and missing heritability. CSUN, Northridge, CA. Seminar, 2019.
- 8. Can modern genomic sequence data fulfill their evolutionary potential? Indiana University, Bloomington, IN. Seminar, 2019.
- 9. Statistical inference of evolutionary processes from genomic data. Macalester College, St. Paul, MN. Seminar, 2019.
- 10. Can modern genomic sequence data fulfill their evolutionary potential? Cornell University, Ithaca, NY. Seminar, 2019.
- 11. Exploiting selection at linked stites to infer the rate and strength of adaptation. Evolution Meeting, Providence, RI. Talk, 2019.
- 12. Genome-scale inference of adaptive evolution: new approaches for answering old questions. *Boise State University, Boise, ID.* Seminar, 2019.
- 13. Genome-scale inference of adaptive evolution: new approaches for answering old questions. *Chapman University, Orange, CA.* Seminar, 2019.
- 14. Evolutionary processes shaping the human genome. Linfield College, McMinnville, OR. Seminar, 2018.
- 15. Scientific teaching workshops for stanford postdocs. Stanford Education Day, Stanford, CA. Talk, 2018.
- 16. Modulation of adaptation rate by background selection in the human genome. Bay Area Population Genomics Meeting, Santa Cruz, CA. Talk, 2018.
- 17. Modulation of adaptation rate by background selection in the human genome. American Society of Naturalists Meeting, Monterey, CA. Talk, 2018.
- 18. Designing and implementing scientific teaching workshops for postdocs. Stanford Teaching and Mentoring Academy seminar series, Stanford, CA. Talk., 2018.
- 19. An analytical upper bound on the number of loci required for all splits of a species tree to appear in a set of gene trees. RECOMB Comparative Genomics Meeting, Montreal, Canada. Talk, 2016.
- Detecting causal genetic variation in populations with complex evolutionary histories. University of Washington, Bothell, WA. Seminar, 2016.
- 21. Explosive growth and the genetic architecture of polygenic traits under selection. American Society of Naturalists Meeting,

Monterey, CA. Talk, 2016.

- 22. Selection and explosive growth may hamper the performance of rare variant association tests. Bay Area Population Genomics Meeting, Stanford, CA. Talk, 2015.
- 23. Recent demography and natural selection hamper the power of rare variant association tests. American Society of Human Genetics Meeting, San Diego, CA. Talk, 2014.
- 24. Model-based simulations of selection and demography with implications for heritable phenotypes and rare variant association tests. UC Berkeley Center for Theoretical Evolutionary Genomics, Berkeley, CA. Talk, 2014.
- 25. Simulations and inference of simultaneous positive and negative selection. Society of Molecular Biology and Evolution Meeting, San Juan, Puerto Rico. Poster, 2014.
- 26. Parameter rescaling for forward simulations of recurrent hitchhiking. Bay Area Population Genomics Meeting, San Francisco, CA. Poster, 2013.
- 27. Forward simulations of recurrent selection and demographics with rescaled parameters. American Society of Human Genetics Meeting, Boston, MA. Poster, 2013.
- 28. Accurate pedigree-based imputation. Department of Human Genetics, University of Chicago, Chicago, IL. Seminar, 2011.

References

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