Alison F. Feder

Contact

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Appointments

2021-Assistant Professor, Department of Genome Sciences, University of Washington 2018-2021 Miller Fellow, Department of Integrative Biology, University of California, Berkeley

Hosts: Oskar Hallatschek & Monty Slatkin

Education

2013-2018 PhD, Biology, Stanford University, Stanford, CA

Advisor: Dmitri Petrov

MSc (by Research), Statistics, University of Oxford, Oxford, UK 2012-2013

Advisor: Gil McVean

2008-2012 BA, Mathematics, summa cum laude, University of Pennsylvania, Philadelphia, PA

Advisor: Joshua Plotkin

Research Funding

2018-2021	Miller Fellowship [Website]
2017-2018	Stanford Center for Computational, Evolutionary & Human Genomics Fellowship [Website]
2016-2017	Gerald J. Lieberman Fellowship [Website]
	Awarded yearly to twelve Stanford graduate students whose teaching, research and

university service demonstrate potential for academic leadership.

2014 Center for Computational, Evolutionary and Human Genomics Trainee Grant

2012-2017 National Science Foundation Graduate Research Fellowship [Website]

2012-2013 Thouron Award [Website]

Awards

2018	Milner Prize in Evolutionary Biology
2018	Samuel Karlin Prize in Mathematical Biology
	Awarded to one graduate student in the Stanford Department of Biology per year
	whose dissertation reaches the highest standard of mathematical biology
2018	James F. Crow Early Career Researcher Finalist (Genetics Society of America)
2017	Gil Omenn Prize for the best article in evolutionary medicine published in the previous year
	Awarded for 'More effective drugs lead to harder selective sweeps in the evolution of
	drug resistance in HIV-1'

2015 Excellence in Teaching Award (Department of Biology, Stanford)

2012 Penn Genome Frontiers Institute Excellence in Genomics Undergraduate Award

2012 Phi Beta Kappa (University of Pennsylvania)

Peer-Reviewed Publications (* denotes equal contributions)

- A. F. Feder, K. Harper, C. J. Brumme, P. S. Pennings (2021). Understanding patterns of HIV multi-drug resistance through models of temporal and spatial drug heterogeneity. eLife, 10:e69032. [Link]
- 10. **A. F. Feder**, P. S. Pennings, D. A. Petrov (2021). The clarifying role of time series data in the population genetics of HIV. *PLOS Genetics* 17(1): e1009050. [Link]
- 9. **A. F. Feder**, P. S. Pennings, J. Hermisson*, D. A. Petrov* (2019). Evolutionary dynamics in structured populations under strong population genetic forces. (G3: GENES, GENOMES, GENETICS) 9(10):3395-3407. [Link]

Selected for 2019 G3 Spotlight issue

- 8. R. S. Mehta, **A. F. Feder**, S. M. Boca, N. A. Rosenberg (2019). The relationship between haplotype-based F_{ST} and haplotype length. *Genetics* 213(1):281-295. [Link]
- 7. K. Theys*, A. F. Feder*, M. Gelbart*, M. Hartl, A. Stern, and P. S. Pennings (2018). Withinpatient HIV mutation frequencies reveal fitness costs of CpG dinucleotides, drastic amino acid changes and $G \to A$ mutations. *PLoS Genetics* 14(6): e1007420. [Link]
- 6. A. F. Feder, C. Kline, P. Polacino, M. Cottrell, A. D. Kashuba, B. F. Keele, S.-L. Hu, D. A. Petrov, P. S. Pennings*, and Z. Ambrose* (2017). A spatio-temporal assessment of simian/human immunodeficiency virus (SHIV) evolution reveals a highly dynamic process within the host. *PLoS Pathogens*, 13(5): e1006358. [Link]
- 5. B. A. Wilson*, N. R. Garud*, A. F. Feder*, Z. J. Assaf*, and P. S. Pennings (2016). The population genetics of drug resistance evolution in natural populations of viral, bacterial and eukaryotic pathogens. *Molecular Ecology*, 25(1):42–66. [Link]
- 4. **A. F. Feder**, S.-Y. Rhee, S. P. Holmes, R. W. Shafer, D. A. Petrov*, and P. S. Pennings* (2016). More effective drugs lead to harder selective sweeps in the evolution of drug resistance in HIV-1. *eLife*, 5:e10670. [Link]
- 3. A. F. Feder*, S. Kryazhimskiy*, and J. B. Plotkin (2014). Identifying signatures of selection in genetic time series. *Genetics*, 196(2):509–522. [Link]
- 2. **A. F. Feder**, D. A. Petrov, and A. O. Bergland (2012). LDx: estimation of linkage disequilibrium from high-throughput pooled resequencing data. *PLoS One*, 7(11):e48588. [Link]
- K. E. Lohmueller, A. Albrechtsen, Y. Li, S. Y. Kim, T. Korneliussen, N. Vinckenbosch, G. Tian, E. Huerta-Sanchez, A. F. Feder, N. Grarup, T. Jørgensen, T. Jiang, D. R. Witte, A. Sandbæk, I. Hellmann, T. Lauritzen, T. Hansen, O. Pedersen, J. Wang, R. Nielsen (2011). Natural selection affects multiple aspects of genetic variation at putatively neutral sites across the human genome. PLoS Genetics, 7(10):e1002326. [Link]

Invited Presentations v virtually 2021^{v} Temporal Genomics Working Group 2021^{v} Miller Institute for Basic Research in Science, UC Berkeley, Berkeley, USA 2021^{v} Quantitative Evolution, Phylogeny and Ecology: from models to data and back, IHP Workshop, Paris, France 2021^{v} Institute of Ecology & Evolution, University of Oregeon, Eugene, USA 2020^{v} Ecology & Evolution Seminar, University of California, Davis, USA 2020 Department of Genome Sciences, University of Washington, Seattle, USA 2019 Department of Ecology & Evolutionary Biology, University of Chicago, Chicago, USA 2019 Department of Computational Biology, Cornell University, Ithaca, USA 2019 Science & Mathematics Seminar, University of Puget Sound, Tacoma, USA European Society of Evolutionary Biology, Turku, Finland 2019 Society of Molecular Biology & Evolution, Manchester, UK 2019 Trainee Invited Speaker Series, Arjun Raj Lab at Penn, Philadelphia, USA 2019 Science & Technology Seminar, Joint Genome Institute, Walnut Creek, USA 2019 Departmental seminar, University of San Francisco, San Francisco, USA 2019 2018 Palo Alto Research Center, Palo Alto, USA 2018 Milner Prize Lecture, University of Bath, Bath, UK Systems Biology Seminar, Cancer Research UK Cambridge Institute, UK 2018 2018 Ad hoc seminar, University of California, Davis, USA Institute for Disease Modeling Annual Symposium, Seattle, USA 2018 2017 Center for Theoretical Evolutionary Genomics, University of California, Berkeley, USA Institute for Disease Modeling, Bellevue, USA 2017 2017 Center for Inference and Dynamics of Infectious Disease, Fred Hutchinson Cancer Research Institute, Seattle, USA 2017 Omenn Prize talk at the International Society of Evolution, Medicine and Public Health, Groningen, Netherlands 2017 Program for Evolutionary Dynamics, Harvard University, Cambridge, USA 2016 "Darwin's Weekly" Seminar, University of Chicago, Chicago, USA Contributed/selected presentations * talk † poster [*] Society for Molecular Biology & Evolution, Yokohama, Japan 2018 2018 [*] James F. Crow Award finalist session at PEQG, Madison, USA 2018 [*] HIV Dynamics & Evolution, Leavenworth, USA 2017 [†] Gordon Research Conference: Microbial Population Biology, Andover, USA 2017 [*] Gordon Research Seminar: Microbial Population Biology, Andover, USA 2017 [*] Society for Molecular Biology & Evolution Annual Meeting, Austin, USA [*] International Society of Evolution, Medicine and Public Health, Raleigh, USA 2016 2016 [*] International HIV Drug Resistance Workshop, Boston, USA

Contributed/Selected Presentations (continued) * talk † poster 2016 [† †] Conference on Retroviruses and Opportunistic Infections (CROI), Boston, USA 2015 [†] Bio-X Interdisciplinary Initiatives Symposium, Stanford, USA 2015 [*] Society for Molecular Biology & Evolution Annual Meeting, Vienna, Austria 2015 [†] "Forecasting Evolution?" SFB 680 Conference, Lisbon, Portugul 2015 [*] Biomedical Computation at Stanford (BCATS), Stanford, USA 2011 [*] NIMBioS Undergraduate Research Conference at the Interface of Biology and Mathe-

matics, Knoxville, USA
2011 [††] Society for Molecular Biology & Evolution Annual Meeting, Kyoto, Japan

Research Supervision

2021-	Elena Romero, Genome Sciences PhD student, U. Washington
2020-	Will Hannon, Molecular & Cellular Biology PhD student, Fred Hutch (J. Bloom lab)
2020-	Maya Lewinsohn, MSTP student (Genome Sciences), U. Washington (T. Bedford lab)
2020	Helen Sakharova, Comp. Biology PhD rotation student, UC Berkeley (O. Hallatschek lab)
2016	Michael Herschl, undergraduate student, Stanford University (D. Petrov lab)

Teaching

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Fall 2015 Co-teacher for BioCore Exploration (3 hour course), 'Are we still evolving?' with L. Uricchio

Spring 2015 TA for Bio 143, Evolution

Spring 2014 TA for Bio 43, Evolution, Ecology & Plant Biology

High School:

2016 Guest lecturer, Evolutionary genomics theory, application and you!

Stanford Pre-Collegiate Institute

2014-2016 Stanford Splash! Teacher

Taught 6 one-session mini-courses to high school students (two each on mathematical/logical

thinking, population genetics and statistics/probability).

Public Outreach

2019	Invited speaker at Nerd Nite East Bay, a general audience seminar series
2017	Finalist in Evolution Film Festival for "Intra-patient Simian-HIV drug resistance evolution:
	does blood tell the whole story?"
2016	Finalist in Evolution Film Festival for "Better drugs lead to harder sweeps in HIV-1"

Competitive travel support

2018	Young Investigator Travel Award from SMBE (Yokohama, Japan)
2016	International Society for Evolutionary Medicine and Public Health Travel Award (Durham,
	USA)
2016	CROI Young Investigator Scholarship (Boston, USA)
2015	Wellcome Trust Travel Award (for "Forecasting Evolution?" meeting, Lisbon, Portugal)
2013	Cargese Summer School in Quantitative Genetics Grant (Cargese, France)
2011	NiMBioS Undergraduate Conference Grant (Knoxville, USA)
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Academic, Community & University Service

2020-2021	Miller Institute DEI Working Group
2019-2021	Miller Symposium Planning Committee
2018	Co-organizer of SMBE 2018 symposium on 'Intra-host evolutionary dynamics' with K. Xue
2017	Co-organizer of 'Petrino' joint lab retreat between D. Petrov and R. Andino (UCSF) labs
2016 - 2017	Department of Biology TA Mentorship Program mentor and program organizer
2014-2017	Stanford Bioscience Students Association new student Mentor
2014-2015	Mentored student writing NSF Graduate Research Fellowship application

Referee for American Society of Naturalists, Evolution, Genetics, Journal of Theoretical Biology, Molecular Biology and Evolution, Nature Ecology & Evolution, PCI Evolutionary Biology, PLOS Computational Biology, PLOS Genetics, PNAS, Virus Evolution