Trevor Bedford

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Faculty positions

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.020-	Affiliate Associate Professor, University of Washington Department of Epidemiology
.019–	Associate Professor, Fred Hutchinson Cancer Research Center Human Biology Division
.019–	Affiliate Associate Professor, University of Washington Department of Genome Sciences
.018–	Associate Professor, Fred Hutchinson Cancer Research Center Vaccine and Infectious Disease Division and Public Health Sciences Division
.016–2020	Affiliate Assistant Professor, University of Washington Department of Epidemiology
.013-2017	Assistant Professor, Fred Hutchinson Cancer Research Center Vaccine and Infectious Disease Division and Public Health Sciences Division
	Postgraduate training
.012-2013	Newton International Fellow, University of Edinburgh Institute of Evolutionary Biology
011	EMBO Fellow, University of Edinburgh Institute of Evolutionary Biology
.008–2011	HHMI Associate, University of Michigan Department of Ecology and Evolutionary Biology
	Education
.003-2008	PhD in Biology, Harvard University
999-2002	BA in Biological Sciences (with honors), University of Chicago
	Honors & awards
.021-2026	MacArthur Fellow, John D. and Catherine T. MacArthur Foundation.
.021-	HHMI Investigator, Howard Hughes Medical Institute.
.020	40 Under 40, Fortune Magazine.

2020	Webby Award, Special Achievement, International Academy of Digital Arts and Sciences.
2017	Open Science Prize Winner, NIH, Wellcome Trust and HHMI.
2016-2020	Pew Scholar in the Biomedical Sciences, Pew Charitable Trusts.
2016	Open Science Prize Finalist, NIH, Wellcome Trust and HHMI.
2012-2014	Marie Curie International Incoming Fellow, European Commission (Declined).
2012-2013	Newton International Fellow, The Royal Society.
2011-2013	Long Term Fellow, European Molecular Biology Organization.
2005-2008	NSF Graduate Research Fellowship, National Science Foundation.
2003-2005	NIH Genetics and Genomics Training Program, National Institutes of Health.
2001	HHMI Undergraduate Fellowship, Howard Hughes Medical Institute.
1999-2002	University Scholarship, University of Chicago.
	Research support
	Ongoing research support
2021-2028	75N93021C00015. National Institutes of Health. Center of Excellence for Influenza Research and Response (CEIRR). Co-investigator with PI Scott Hensley.
2020-2023	INV-018979. Bill and Melinda Gates Foundation. Nextstrain funding for Africa PGI. Principal investigator.
2018-2021	Seattle Flu Study. Gates Ventures. Principal investigator.
2017-2022	Wellcome Trust Collaborative Award 206298/Z/17/Z. Wellcome Trust. Putting genomic surveillance at the heart of viral epidemic response. Co-investigator with PI Andrew Rambaut.
2017-2022	Ro1 AI125392. National Institutes of Health. Mechanisms of influenza transmission bottlenecks: impact on viral evolution. Co-investigator with PI Thomas Friedrich.
2016-2021	Ro1 AI127893. National Institutes of Health. High-throughput experiments to guide influenza vaccine strain selection. Co-investigator with PI Jesse Bloom.
2016-2026	R ₃₅ GM ₁₁₉₇₇₄ . National Institutes of Health. Real-time tracking of virus evolution for vaccine strain selection and epidemiological investigation. Principal investigator.

COMPLETED RESEARCH SUPPORT

2015–2020 U19 AI117891-01. National Institutes of Health.

Dynamics and evolution of immune responses to influenza viruses.

Co-investigator with PIs Rafi Ahmed and Rustom Antia.

2014-2019 Ro1 GM113246-01. National Science Foundation and National Institutes of Health.

Leveraging deep sequencing data to understand antibody maturation.

Co-investigator with PI Erick Matsen.

2014-2019 U54 GM111274. National Institutes of Health.

Center for Statistics and Quantitative Infectious Diseases.

Co-investigator with PI M. Elizabeth Halloran, co-lead of project on "Understanding transmission with integrated genetic and epidemiologic inference".

Teaching & mentoring

Courses and workshops

- 2020- Instructor, Introduction to Computational Molecular Biology, UW Genome Sciences.
- Instructor, Tools for Computational Biology, UW MCB.
- Instructor, Pathogen Evolution, Selection and Immunity, Summer Institute in Statistics and Modeling in Infectious Diseases, University of Washington.
- 2014-2016 Lectures on data science and visualization for interns at the FHCRC.
- Presented phylodynamic inference practical at GenEpi workshop, London School of Hygiene and Tropical Medicine.
- Lecturer, Structured Population Models in Epidemiology, École Normale Supérieure.
- Completed an intensive 7-week "Postdoctoral Short-Course on College Teaching in Science and Engineering" at the University of Michigan.
- Teaching fellow, Population Genetics, Harvard University.
- Teaching fellow, Genetics and Genomics, Harvard University.

STUDENT AND POSTDOCTORAL MENTORING

- Miguel Paredes, MSTP student, Department of Epidemiology, University of Washington.
- 2020- Marlin Figgins, PhD student, Department of Applied Mathematics, University of Washington.
- 2020- Cassia Wagner, MSTP student, Department of Genome Sciences, University of Washington.
- Nicola Müller, postdoc, FHCRC.
- 2019- Maya Lewinsohn, MSTP student, Department of Genome Sciences, University of Washington.
- 2018– Katie Kistler, PhD student, Molecular and Cellular Biology Program, University of Washington.
- John Huddleston, PhD student, Molecular and Cellular Biology Program, University of Washington.

- Louise Moncla, postdoc, FHCRC.
- James Hadfield, postdoc, FHCRC.
- 2016-2018 Gytis Dudas, postdoc, FHCRC.
- 2016-2017 Stephanie Stacy, Undergraduate, Summer Undergraduate Research Program, FHCRC.
- 2015–2020 Allison Black, PhD student, Department of Epidemiology, University of Washington.
- 2015–2019 Sidney Bell, PhD student, Molecular and Cellular Biology Program, University of Washington.
- 2015-2016 Charlton Callender, Undergraduate, Summer Undergraduate Research Program, FHCRC.
- 2014–2016 Alvason Li, postdoc, FHCRC.
- 2014–2015 Charles Cheung, postdoc, FHCRC.

Professional service

- 2020–2021 Member: Standing Committee on Emerging Infectious Diseases and 21st Century Health Threats,
 - National Academies of Sciences, Engineering, and Medicine.
- 2020–2021 Member: Advisory Board, The COVID Tracking Project.
- Advisor: WHO Global Influenza Surveillance and Response System.
- 2016–2017 Organizing committee: Epidemics⁶ conference.
- 2016–2017 Technical working group member: WHO Public Health Research Agenda for Influenza.
- 2015 Organizing committee: Epidemics 5 conference.
- Associate editor: Virus Evolution.

Ad-hoc referee: American Naturalist, Bioinformatics, BMC Evolutionary Biology, Cell, Complex Systems, eLife, Emerging Infectious Diseases, Genome Biology and Evolution, Genome Research, Journal of Biology, Journal of the Royal Society Interface, Molecular Biology and Evolution, Molecular Ecology, National Science Foundation, Nature, Nature Communications, Nature Genetics, Phil Trans R Soc B, PLoS Biology, PLoS One, PLoS Pathogens, Proc Natl Acad USA, Science, Virology, Virulence, Wellcome Trust.

Publications & talks

Journal articles (98 papers, 9930 citations, h-index of 42)

- Weil AA, Sohlberg SL, O'Hanlon JA, Casto AM, Emanuels AW, ..., Lockwood CM, Starita LM, **Bedford T**, Shendure JA, Chu HY (36 authors). SARS-CoV-2 epidemiology on a public university campus in Washington State. Open Forum Infect Dis: ofab464.
- Annavajhala MK, Mohri H, Wang P, Nair M, Zucker JE, ..., Tagliavia M, Huang Y, **Bedford T**, Ho DD, Uhlemann A-C (13 authors). Emergence and expansion of SARS-CoV-2 B.1.526 after identification in New York. Nature: s41586-021-03908-2.
- Braun K, Moreno G, Wagner C, Accola MA, Rehrauer WM, Baker D, Koelle K, O'Connor DH, Bed-

- ford T, Friedrich TC, Moncla LH. Acute SARS-CoV-2 infections harbor limited within-host diversity and transmit via tight transmission bottlenecks. PLoS Pathog 17: e1009849.
- Srivatsan S, Heidl S, Pfau B, Martin BK, Han PD, ..., **Bedford T**, ..., Konnick EQ, Debley JS, Shendure J, Lockwood CM, Starita LM (36 authors). SwabExpress: An end-to-end protocol for extraction-free COVID-19 testing. Clin Chem: hvab132.
- Corey L, Beyrer C, Cohen MS, Michael NL, **Bedford T**, Rolland M. SARS-CoV-2 variants in patients with immunosuppression. N Engl J Med 385: 562–566.
- Perchetti GA, Zhu H, Mills MG, Shrestha L, **Wagner** C, ..., Mathias P, **Bedford** T, Jerome KR, Greninger AL, Roychoudhury P (14 authors). Specific allelic discrimination of N501Y and other SARS-CoV-2 mutations by ddPCR detects B.1.1.7 lineage in Washington State. J Med Virol 93: 5931–5941.
- Konings F, Perkins MD, Kuhn JH, Pallen MJ, Alm EJ, ..., **Bedford T**, ..., Tong S, van der Werf S, von Gottberg A, Ziebuhr J, Van Kerkhove MD (47 authors). SARS-CoV-2 Variants of Interest and Concern naming scheme conducive for global discourse. Nat Microbiol 6: 821–823.
- Chung E, Chow EJ, Wilcox NC, Burstein R, Brandstetter E, ..., Hughes JP, Starita LM, **Bedford T**, Englund JA, Chu HY (23 authors). Symptoms and RNA levels in pediatric vs adult SARS-CoV-2 infections in the community setting. JAMA Pediatr. *In press*.
- Heimonen JT, McCulloch DJ, O'Hanlon J, Ashley Erin Kim, Emanuels A, ..., Boeckh M, Starita LM, Bedford T, Englund JA, Chu HY (19 authors). A remote household-based approach to influenza self-testing and antiviral treatment. Influenza Other Respir Viruses 15: 469–477.
- Müller NF, Wagner C, Frazar CD, Roychoudhury P, Lee J, ..., Jerome KR, Lindquist S, Greninger AL, Nickerson DA, Bedford T (43 authors). Viral genomes reveal patterns of the SARS-CoV-2 outbreak in Washington State. Sci Transl Med: eabfo202.
- Moncla LH, Black A, DeBolt C, Lang M, Graff NR, Pérez-Osorio AC, Müller NF, Haselow D, Lindquist S, Bedford T. Repeated introductions and intensive community transmission fueled a mumps virus outbreak in Washington State. eLife 10: e66448.
- Jackson ML, Hart GR, McCulloch DJ, Adler A, Brandstetter E, ..., Starita L, Englund JA, **Bedford T**, Chu H, Famulare M (17 authors). Effects of weather-related social distancing on city-scale transmission of respiratory viruses. BMC Infect Dis 21: 335.
- Barrat-Charlaix P, **Huddleston J**, **Bedford T**, Neher RA. Limited predictability of amino acid substitutions in seasonal influenza viruses. Mol Biol Evol 38: 2767–2777.
- Kinganda-Lusamaki E, **Black A**, Mukadi DB, **Hadfield H**, Mbala-Kingebeni P, ..., Peeters M, Wiley MR, Ahuka-Mundeke S, **Bedford T**, Muyembe Tamfum J-J (29 authors). Integration of genomic sequencing into the response to the Ebola virus outbreak in Nord Kivu, Democratic Republic of the Congo. Nat Med 27: 710–716.
- Mbala-Kingebeni P, Pratt P, Mutafali Ruffin M, Pauthner MG, Bile F, ..., **Bedford T**, Andersen KG, Wiley MR, Ahuka-Mundeke S, Muyembe Tamfum J-J (50 authors). Ebola virus transmission initiated by systemic Ebola virus disease relapse. N Engl J Med 384: 1240-1247.
- Kim AE, Brandstetter E, Wilcox N, Heimonen J, Graham C, ..., Rieder MJ, **Bedford T**, Boeckh M, Englund JA, Chu HY on behalf of the Seattle Flu Study Investigators (25 authors). Evaluating specimen quality and results from a community-wide, home-based respiratory surveillance study. J Clin

- Microbiol 59: e02934-20.
- Kistler KE, Bedford T. Evidence for adaptive evolution in the receptor-binding domain of seasonal coronaviruses OC43 and 229E. eLife 10: e64509.
- Huddleston J, Hadfield J, Sibley TR, Lee J, Fay K, Ilcisin M, Harkins E, Bedford T, Neher RA, Hodcroft EB. Augur: a bioinformatics toolkit for phylogenetic analyses of human pathogens. J Open Source Softw 6: 2906.
- Jackson LK, **Potter B**, Schneider SE, Fitzgibbon M, Blair K, Farah H, Krishna U, **Bedford T**, Peek R, Salama NR. *Helicobacter pylori* diversification during chronic infection within a single host generates sub-populations with distinct phenotypes. PLoS Pathog 16: e1008686.
- Müller NF, Wüthrich D, Goldman N, Sailer N, Saalfrank C, ..., **Bedford T**, Battegay M, Schneider-Sliwa R, Egli A, Stadler T (25 authors). Characterising the epidemic spread of influenza A/H₃N₂ within a city through phylogenetics. PLoS Pathog 16: e1008984.
- Emanuels A, Heimonen J, O'Hanlon J, Kim AE, Wilcox N, ..., Starita LM, Nickerson DA, **Bedford** T, Englund JA, Chu HY, on behalf of Seattle Flu Study investigators (21 authors). Remote household observation for non-influenza respiratory viral illness. Clin Infect Dis: ciaa1719.
- 2020 Chu HY, Boeckh M, Englund JA, Famulare M, Lutz BR, ..., Newman KL, Sibley TR, Zigman Suchsland ML, Wolf C, Shendure J, Bedford T, on behalf of the Seattle Flu Study Research Group (30 authors). The Seattle Flu Study: a multi-arm community-based prospective study protocol for assessing influenza prevalence, transmission, and genomic epidemiology. BMJ Open 10: e037295.
- Rogers JH, Link AC, McCulloch D, Brandstetter E, Newman KL, ..., Richardson M, Nickerson DA, Starita LM, **Bedford T**, Chu HY, on behalf of Seattle Flu Study Investigators (21 authors). Characteristics of COVID-19 in homeless shelters: A community-based surveillance study. Ann Intern Med: 10.7326/M20-3799.
- Bedford T, Greninger AL, Roychoudhury P, Lea M Starita, Famulare M, ..., Armstrong GL, Baird GS, Chu HY, Shendure J, Jerome KR (56 authors). Cryptic transmission of SARS-CoV-2 in Washington State. Science 370: 571–575.
- Weil AA, Newman KL, Ong TD, Davidson GH, Logue J, ..., Zhong W, Starita LM, **Bedford T**, Roxby AC, Chu HY (17 authors). Cross-sectional prevalence of SARS-CoV-2 among skilled nursing facility employees and residents across facilities in Seattle. J Gen Intern Med 35: 3302–3307.
- Huddleston J, Barnes JR, Rowe T, Kondor R, Wentworth DE, ..., Barr I, Subbarao K, Barrat-Charlaix P, Neher RA, Bedford T (19 authors). Integrating genotypes and phenotypes improves long-term forecasts of seasonal influenza A/H₃N₂ evolution. eLife 9: e60067.
- Hilton SK, **Huddleston JL**, **Black A**, North K, Dingens AS, **Bedford T**, Bloom JD. dms-view: Interactive visualization tool for deep mutational scanning data. J Open Source Softw 5: 2353.
- Black A, MacCannell DR, Sibley TR, Bedford T. Ten recommendations for supporting open pathogen genomic analysis in public health. Nat Med 26: 832–841.
- Deng X, Gu W, Federman S, Du Plessis L, Pybus O, ..., **Bedford T**, ..., Chai S, Villarino E, Bonin B, Wadford D, Chiu CY (47 authors). A genomic survey of SARS-CoV-2 reveals multiple introductions into Northern California without a predominant lineage. Science 369: 582–587.
- 2020 CDC COVID-19 Response Team, Jorden MA, Rudman SL, Villarino E, Hoferka S, ..., Bedford T,

- ..., Famulare M, Nickerson DA, Rieder MJ, Shendure J, Starita LM (27 authors). Evidence for limited early spread of COVID-19 within the United States, January–February 2020. Morb Mortal Wkly Rep 69: 680–684.
- Chu HY, Englund JA, Starita LM, Famulare M, Brandstetter E, ..., Jackson M, Hughes JP, Boeckh M, Shendure J, **Bedford T** (30 authors). Early detection of Covid-19 through a citywide pandemic surveillance platform. N Engl J Med 383: 185–187.
- Caudill VR, Qin S, Winstead R, Kaur J, Tisthammer K, ..., **Bedford T**, ..., Tran K, Tran L, Winters EJ, Wong A, Pennings PS (56 authors). CpG-creating mutations are costly in many human viruses. Evol Ecol 34: 339–359.
- Castro LA, **Bedford T**, Meyers LA. Early prediction of antigenic transitions for influenza A/H₃N₂. PLoS Comput Biol 16: e100768₃.
- Moncla LH, Bedford T, Dussart P, Horm SV, Rith S, Buchy P, Karlsson EA, Li L, Liu Y, Zhu H, Guan Y, Friedrich TC, Horwood PF. Quantifying within-host evolution of H5N1 influenza in humans and poultry in Cambodia. PLoS Pathog 16: e1008191.
- Dudas G, Bedford T. The ability of single genes vs full genomes to resolve time and space in outbreak analysis. BMC Evol Biol 19: 232.
- Potter BI, Kondor R, Hadfield J, Huddleston J, Barnes J, Rowe T, Guo L, Xu X, Neher RA, Bedford T, Wentworth DE. Evolution and rapid spread of a reassortant A(H₃N₂) virus that predominated the 2017-2018 influenza season. Virus Evol 5: vezo46.
- Black A, Moncla LH, Laiton-Donato K, Potter B, Pardo L, Tovar C, Rojas DP, Longini IM, Halloran ME, Peláez-Carvajal D, Ramírez JD, Mercado-Reyes M, Bedford T. Genomic epidemiology supports multiple introductions and cryptic transmission of Zika virus in Colombia. BMC Infect Dis 19: 963.
- Hadfield J, Brito AF, Swetnam DM, Vogels CBF, Tokarz RE, Andersen KG, Smith RC, Bedford T, Grubaugh ND. Twenty years of West Nile virus spread and evolution in the Americas visualized by Nextstrain. PLoS Pathog 15: e1008042.
- Bell SM, Katzelnick L, Bedford T. Dengue genetic divergence generates within-serotype antigenic variation, but serotypes dominate evolutionary dynamic. eLife 8: e42496.
- Lee JM, Eguia R, Zost SJ, Choudhary S, Wilson PC, **Bedford T**, Stevens-Ayers T, Boeckh M, Hurt A, Lakdawala SS, Hensley SE, Bloom JD. Mapping person-to-person variation in viral mutations that escape polyclonal serum targeting influenza hemagglutinin. eLife 8: e49324.
- Dale GA, Wilkins DJ, Bohannon CD, Dilernia D, Hunter E, **Bedford T**, Antia R, Sanz I, Jacob J. Clustered mutations at the murine and human IgH locus exhibit significant linkage consistent with templated mutagenesis. J Immunol: ji1801615.
- Soh YQS, Moncla LH, Eguia R, Bedford T, Bloom JD. Comprehensive mapping of avian influenza PB2 protein as a function of virus adaptation to humans. eLife 8: e45079.
- Neher R, **Bedford T**. Real-time analysis and visualization of pathogen sequence data. J Clin Microbiol 56: e00480-18.
- Wen FT, **Bell SM**, **Bedford T**, Cobey S. Estimating vaccine-driven selection in seasonal influenza. Viruses 10: 509.

- Lee JM, **Huddleston J**, Doud MB, Hooper KA, Wu NC, **Bedford T**, Bloom JD. Deep mutational scanning of hemagglutinin helps predict evolutionary fates of human H₃N₂ influenza variants. Proc Natl Acad Sci USA 115: E8276–E8285.
- Hadfield J, Megill C, Bell SM, Huddleston J, Potter B, Callender C, Sagulenko P, Bedford T, Neher RA. Nextstrain: real-time tracking of pathogen evolution. Bioinformatics 34: 4121–4123.
- Xue KS, **Moncla LH**, **Bedford T**, Bloom JD. Within-host evolution of human influenza virus. Trends Microbiol 26: 781–793.
- Rouet F, Nouhin J, Zheng D, Roche B, **Black A**, ..., Barin F, Plantier JC, **Bedford T**, Ramos A, Saphonn V (26 authors). Massive iatrogenic outbreak of human immunodeficiency virus type 1 in rural Cambodia, 2014-2015. Clin Infect Dis 66: 1733-1741.
- Morris DH, Gostic KM, Pompei S, **Bedford T**, Łuksza M, Neher RA, Grenfell BT, Lässig M, Mc-Cauley JW. Predictive modeling of influenza shows the promise of applied evolutionary biology. Trends Microbiol 26: 102–118.
- Dudas G, Max Carvalho L, Rambaut A, Bedford T. MERS-CoV spillover at the camel-human interface. eLife 7: e31257.
- Cybis GB, Sinsheimer JS, **Bedford T**, Rambaut A, Lemey P, Suchard MA. Bayesian nonparametric clustering in phylogenetics: modeling antigenic evolution in influenza. Stat Med 37: 195–206.
- Langat P, Raghwani J, **Dudas G**, Bowden T, Edwards S, ..., **Bedford T**, ..., Russell C, Pybus O, Kellam P, McCauley J, Watson SJ (14 authors). Genome-wide evolutionary dynamics of influenza B viruses on a global scale. PLoS Pathog 13: e1006749.
- Bell SM, Bedford T. Modern-day SIV viral diversity generated by extensive recombination and crossspecies transmission. PLoS Pathog 13: e1006466.
- Grubaugh ND, Ladner JT, Kraemer MUG, **Dudas G**, Tan AL, ..., **Bedford T**, Pybus OG, Isern S, Palacios G, Andersen KG (67 authors). Genomic epidemiology reveals multiple introductions of Zika virus into the United States. Nature 546: 401–405.
- Faria NR, Quick J, Morales I, Thézé J, Jesus JG, ..., **Bedford T**, Teixeira MNR, Sabino EC, Alcantara LCJ, Loman N, Pybus OG (71 authors). Establishment and cryptic transmission of Zika virus in Brazil and the Americas. Nature 546: 406–410.
- Quick J, Grubaugh ND, Pullan ST, Claro IM, Smith AD, ..., **Bedford T**, ..., Simpson JT, Pybus OG, Andersen KG, Loman NJ (28 authors). Multiplex PCR method for MinION and Illumina sequencing of Zika and other virus genomes directly from clinical samples. Nat Protoc 12: 1261–1276.
- Dudas G, Max Carvalho L, Bedford T, Tatem AJ, Baele G, ..., Ströher U, Wury I, Suchard MA, Lemey P, Rambaut A (93 authors). Virus genomes reveal the factors that spread and sustained the West African Ebola epidemic. Nature 544: 309–315.
- Black A, Breyta R, Bedford T, Kurath G. Geography and host species shape the evolutionary dynamics of U genogroup infectious hematopoietic necrosis virus. Virus Evol 2: vewo34.
- Lipsitch M, Barclay W, Raman R, Russell CJ, Belser JA, ..., **Bedford T**, ..., Herfst S, Murcia PR, Roche B, Wilke CO, Russell CA (19 authors). Viral factors in influenza pandemic risk assessment. eLife 5: e18491.

- Bedford T, Malik HS. Did a single amino acid change make Ebola virus more virulent? Cell 167: 892–894.
- Wen F, **Bedford T**, Cobey S. Explaining the geographic origins of seasonal influenza A (H₃N₂). Proc R Soc B 283: 20161312.
- Harvey WT, Benton DJ, Gregory V, Hall JPJ, Daniels RS, **Bedford T**, Haydon DT, Hay AJ, McCauley JW, Reeve R. Identification of low- and high-impact hemagglutinin amino acid substitutions that drive antigenic drift of influenza A(H1N1) viruses. PLoS Pathog 12: e1005526.
- Neher RA, **Bedford T**, Daniels RS, Russell CA, Shraiman BI. Prediction, dynamics, and visualization of antigenic phenotypes of seasonal influenza viruses. Proc Natl Acad Sci USA 113: E1701–E1709.
- Karcher MD, Palacios JA, **Bedford T**, Suchard MA, Minin VN. Quantifying and mitigating the effect of preferential sampling on phylodynamic inference. PLoS Comp Biol 12: e1004789.
- Lorenzo-Redondo R, Fryer HR, **Bedford T**, Kim EY, Archer J, ..., Malim MH, Rambaut A, Haase AT, McLean AR, Wolinsky SM (16 authors). Persistent HIV-1 replication maintains the tissue reservoir during therapy. Nature 530: 51–56.
- Neafsey DE, Juraska M, **Bedford T**, Benkeser D, Valim C, ..., Yu Q, Lapierre D, Birren BW, Gilbert PB, Wirth DF (63 authors). Genetic diversity and protective efficacy of the RTS,S/ASo1 malaria vaccine. New Engl J Med 373: 2025–2037.
- Machkovech HM, **Bedford T**, Suchard MA, Bloom JD. Positive selection in CD8+ T-cell epitopes of influenza nucleoprotein revealed by a comparative analysis of human and swine viral lineages. J Virol 89: 11275-11283.
- Neher RA, **Bedford T**. nextflu: real-time tracking of seasonal influenza virus evolution in humans. Bioinformatics 31: 3546–3548.
- Meyer AG, Spielman SJ, **Bedford T**, Wilke CO. Time dependence of evolutionary metrics during the 2009 pandemic influenza virus outbreak. Virus Evol 1: vevoo6.
- Cybis G, Sinsheimer J, **Bedford T**, Mather AE, Lemey P, Suchard MA. Assessing phenotypic correlation through the multivariate phylogenetic latent liability model. Ann Appl Stat 9: 969–991.
- McCoy CO, **Bedford T**, Minin VN, Bradley P, Robins H, Matsen FA. Quantifying evolutionary constraints on B cell affinity maturation. Phil Trans R Soc B 370: 20140244.
- Bedford T, Riley S, Barr IG, Broor S, Chadha M, ..., Tashiro M, Wang D, Xu X, Lemey P, Russell CA (26 authors). Global circulation patterns of seasonal influenza viruses vary with antigenic drift. Nature 523: 217–220.
- Park DJ, Dudas G, Wohl S, Goba A, Whitmer SLM, ..., **Bedford T**, Gnirke A, Ströher U, Rambaut A, Garry RF, Sabeti PC (86 authors). Ebola virus epidemiology, transmission, and evolution during seven months in Sierra Leone. Cell 161: 1–11.
- Frost SDW, Pybus OG, Gog JR, Viboud C, Bonhoeffer S, **Bedford T**. Eight challenges in phylodynamic inference. Epidemics 10: 88–92.
- Vrancken B, Lemey P, Rambaut A, **Bedford T**, Longdon B, Gunthard H, Suchard MA. Simultaneously estimating evolutionary history and repeated traits phylogenetic signal: applications to viral and host

- phenotypic evolution. Methods Ecol Evol 6: 67-82.
- Dudas G, **Bedford T**, Lycett S, Rambaut A. Reassortment between influenza B lineages and the emergence of a co-adapted PB1-PB2-HA gene complex. Mol Biol Evol 32: 162–172.
- Zinder D, **Bedford T**, Baskerville EB, Woods RJ, Roy M, Pascual M. Seasonality in the migration and establishment of H₃N₂ influenza lineages with epidemic growth and decline. BMC Evol Biol 14: 272.
- Faria NR, Rambaut A, Suchard MA, Baele G, **Bedford T**, Ward MJ, Tatem AJ, Sousa JD, Arinamin-pathy N, Pépin J, Posada D, Peeters M, Pybus OG, Lemey P. The early spread and epidemic ignition of HIV-1 in human populations. Science 346: 56–61.
- Roche B, Drake JM, Brown J, Stallknecht DE, **Bedford T**, Rohani P. Adaptive evolution and environmental durability jointly structure phylodynamic patterns in avian influenza viruses. PLoS Biol 12: e1001931.
- Lemey P, Rambaut A, **Bedford T**, Faria N, Bielejec F, Baele G, Russell CA, Smith DJ, Pybus OG, Brockmann D, Suchard MA. Unifying viral genetics and human transportation data to predict the global transmission dynamics of human influenza H₃N₂. PLoS Pathog 10: e1003932.
- Bedford T, Suchard MA, Lemey P, Dudas G, Gregory V, Hay AJ, McCauley JW, Russell CA, Smith DJ, Rambaut A. Integrating influenza antigenic dynamics with molecular evolution. eLife 3: e01914.
- Landis MJ, **Bedford T**. Phylowood: interactive web-based animations of biogeographic and phylogeographic histories. Bioinformatics 30: 123–124.
- Volz EM, Koelle K, Bedford T. Viral phylodynamics. PLoS Comput Biol 9: e1002947.
- Zinder D, **Bedford T**, Gupta S, Pascual M. The roles of competition and mutation in shaping antigenic and genetic diversity in influenza. PLoS Pathog 9: e1003104.
- Bedford T, Rambaut A, Pascual M. Canalization of the evolutionary trajectory of the human influenza virus. BMC Biol 10: 38.
- Baele G, Lemey P, **Bedford T**, Rambaut A, Suchard MA, Alekseyenko AV. Improving the accuracy of demographic and molecular clock model comparison while accommodating phylogenetic uncertainty. Mol Biol Evol 29: 2157–2167.
- Baskerville EB, Dobson AP, **Bedford T**, Allesina S, Anderson TM, Pascual M. Spatial guilds in the Serengeti food web revealed by a Bayesian group model. PLoS Comput Biol 7: e1002321.
- Bedford T, Cobey S, Pascual M. Strength and tempo of selection revealed in viral gene genealogies.

 BMC Evol Biol 11: 220.
- Rogers RL, **Bedford T**, Lyons AM, Hartl DL. Adaptive impact of the chimeric gene *Quetzalcoatl* in *Drosophila melanogaster*. Proc Natl Acad Sci USA 107: 10943–10948.
- Bedford T, Cobey S, Beerli P, Pascual M. Global migration dynamics underlie evolution and persistence of human influenza A (H₃N₂). PLoS Pathog 6: e1000918. (Faculty of 1000 "Must Read")
- Bedford T, Hartl DL. Optimization of gene expression by natural selection. Proc Natl Acad Sci USA 106: 1133-1138.
- 2009 Rogers RL, **Bedford T**, Hartl DL. Formation and longevity of chimeric and duplicate genes in *Drosophila*

- melanogaster. Genetics 181: 313-322.
- Bedford T, Hartl DL. Overdispersion of the molecular clock: temporal variation of gene-specific substitution rates in *Drosophila*. Mol Biol Evol 25: 1631–1638.
- Bedford T, Wapinski I, Hartl DL. Overdispersion of the molecular clock varies between yeast, *Drosophila* and mammals. Genetics 179: 977–984.
- Volkman SK, Lozovsky E, Barry AE, **Bedford T**, Bethke L, Myrick A, Day KP, Hartl DL, Wirth DF, Sawyer SA. Genomic heterogeneity in the density of noncoding single-nucleotide and microsatellite polymorphisms in *Plasmodium falciparum*. Gene 387: 1–6.
- Castillo-Davis CI, **Bedford TB**, Hartl DL. Accelerated rates of intron gain/loss and protein evolution in duplicate genes in human and mouse malaria parasites. Mol Biol Evol 21: 1422–1427.
- Nielsen KM, Kasper J, Choi M, **Bedford T**, Kristiansen K, Wirth DF, Volkman SK, Lozovsky ER, Hartl DL. Gene conversion as a source of nucleotide diversity in *Plasmodium falciparum*. Mol Biol Evol 20: 726–734.
- Feder ME, **Bedford T**, Albright DR, Michalak P. Evolvability of Hsp70 expression under artificial selection for inducible thermotolerance in independent populations of *Drosophila melanogaster*. Phys Biochem Zool 75: 325–334.

Invited presentations

- VIDD Seminar, Fred Hutchinson Cancer Research Center, Seattle, WA.
- Plenary talk, AGBT General Meeting.
- Mathematics and Statistics of Genomic Epidemiology Workshop, Casa Matemática Oaxaca and Banff International Research Station for Mathematical Innovation and Discovery.
- VIDD Seminar, Fred Hutchinson Cancer Research Center, Seattle, WA.
- AMD Virtual, Centers for Disease Control and Prevention, Atlanta, GA.
- 2020 COVID-19 Medical Grand Rounds, UCSF Department of Medicine, San Francisco, CA.
- 2020 COVID-19 Dynamics and Evolution.
- Keynote talk, AACR Virtual Meeting: COVID-19 and Cancer.
- Biology of Genomes, Cold Spring Harbor, NY.
- AAAS Annual Meeting, Seattle, WA.
- MCMB Seminar Series, SF State University, San Francisco, CA.
- 2020 CCB Seminar Series, UC Berkeley, Berkeley, CA.
- BioHub Seminar Series, BioHub, San Francisco, CA.
- EPPIcenter Seminar Series, UCSF, San Francisco, CA.
- Options X for the Control of Influenza, Singapore.
- Distinguished Scientist Seminar Series, Rocky Mountain Laboratories, Hamilton, MT.

Northwest Data Science Summit, University of Washington, Seattle, WA. 2019 Population Biology, Ecology and Evolution Seminar, Emory University, Atlanta, GA. 2019 SMBE Satellite Workshop on Genome Evolution in Pathogen Transmission and Disease, Kyoto, Japan. 2018 Grand Challenges Annual Meeting, Berlin, Germany. 2018 Plenary talk, International Conference on Emerging Infectious Diseases, Atlanta, GA. 2018 Emerging Infections and Pandemic Risk, Institut Pasteur, Paris, France. 2018 AMD Seminar, Centers for Disease Control and Prevention, Atlanta, GA. 2.018 Plenary talk, HIV Dynamics and Evolution, Leavenworth, WA. 2018 6th Annual IDM Symposium, Institute for Disease Modeling, Bellevue, WA. 2018 Bioinformatics and Genomics Seminar, University of North Carolina, Charlotte, NC. 2018 Annual International Symposium, Institute for Systems Biology, Seattle, WA. 2018 NIHE/OUCRU Workshop on Influenza Epidemiology and Evolution in Vietnam, Hanoi, Vietnam. 2018 Department of Hygiene Seminar, Hokkaido University, Sapporo, Japan. 2018 CREST International Symposium on Big Data Applications, Tokyo, Japan. 2018 BioHub Seminar, Chan Zuckerberg BioHub, San Francisco, CA. 2017 VIDD Seminar, Fred Hutchinson Cancer Research Center, Seattle, WA. 2017 Viral Pathogenesis Symposium, Fred Hutchinson Cancer Research Center, Seattle, WA. 2017 Global Infectious Disease Seminar, Center for Infectious Disease Reseach, Seattle, WA. 2017 Immunology and Evolution of Influenza Symposium, Emory University, Atlanta, GA. 2017 Applied Bioinformatics and Public Health Microbiology, Wellcome Genome Campus, Hinxton, UK. 2017 Biosystems Science and Engineering Seminar, ETH Zurich, Basel, Switzerland. 2017 Gordon Research Conference on Chemical and Biological Terrorism Defense, Ventura, CA. 2017 Pew Annual Meeting, Santa Barbara, CA. 2017 BD2K Open Data Science Symposium, Bethesda, MD. 2016 Plenary talk, Options IX for the Control of Influenza, Chicago, IL. 2016 WHO Workshop on Predictive Models for Improving Influenza Vaccine Virus Selection, Princeton 2016 University, Princeton, NJ. Microbiology Seminar, Korea University College of Medicine, Seoul, Republic of Korea. 2016 Federation Meeting of Korean Basic Medical Scientists, Incheon, Republic of Korea. 2016

4th Annual IDM Symposium, Institute for Disease Modeling, Bellevue, WA.

MIDAS Network Meeting, Washington, DC.

2016

2016

Ecology and Evolution Seminar, University of Chicago, Chicago, IL. Structure and Computation Affinity Group Seminar, Scripps Research Institute, San Diego, CA. 2016 Infectious Disease Epidemiology Seminar, Harvard School of Public Health, Boston, MA. 2016 VIDD Seminar, Fred Hutchinson Cancer Research Center, Seattle, WA. 2016 Influenza Division, VSDB Meeting, Centers for Disease Control and Prevention, Atlanta, GA. 2016 4th Informal Consultation on Influenza Vaccine Virus Selection, World Health Organization, Hong 2015 Kong, SAR China. Global Infectious Disease Seminar, AIDS Vaccine Research Laboratory, University of Wisconsin, Madi-2015 son, WI. Evolution Seminar, JF Crow Institute, University of Wisconsin, Madison, WI. 2015 Keynote talk, BCCDC Research Week Symposium, Vancouver, Canada. 2015 Forecasting Evolution, Calouste Gulbenkian Foundation, Lisbon, Portugal. 2015 Keynote talk, Ecology and Evolution of Infectious Diseases, Athens, GA. 2015 RAPIDD Workshop on Modeling and Predicting Influenza Phenotypes, Cambridge University, Cam-2015 bridge, UK. Laboratory Medicine Grand Rounds, University of Washington School of Medicine, Seattle, WA. 2014 Evolution of Drug Resistance Workshop, Kavli Institute for Theoretical Physics, University of Califor-2014 nia, Santa Barbara, CA. VIDD Seminar, Fred Hutchinson Cancer Research Center, Seattle, WA. 2014 Infectious Disease Dynamics Workshop, Isaac Newton Institute for Mathematical Sciences, Cambridge 2013 University, Cambridge, UK. Spring Symposium, Centre for Immunity, Infection and Evolution, University of Edinburgh, Edin-2013 burgh, UK. Institute of Biologie Seminar, École Normale Supérieure, Paris, France. 2012 Infectious Disease Epidemiology Seminar, School of Public Health, Imperial College, London, UK. 2012 Ecology and Evolution Seminar, Centre d'Ecologie Fonctionnelle et Evolutive, Université Montpellier, 2012 Montpellier, France. Institute Seminar, Institute of Medical Virology and Epidemiology of Viral Diseases, Universitäts Klinikum 2012 Tübingen, Tübingen, Germany. Virology Seminar, MRC National Institute for Medical Research, London, UK. 2012 Next-Generation Molecular and Evolutionary Epidemiology of Infectious Disease, The Royal Society, 2012 Kavli International Centre, Buckinghamshire, UK. VIDD Seminar, Fred Hutchinson Cancer Research Center, Seattle, WA. 2012 Scottish Phylogeny Discussion Group, University of Glasgow, Glasgow, UK. 2012

2016

2012	Viral Evolution: Linking Genetics to Epidemics, University of Cologne, Cologne, Germany.
2011	Thematic Seminar, Institute of Biodiversity, Animal Health and Comparative Medicine, University of Glasgow, Glasgow, UK.
2011	RAPIDD Phylodynamics Workshop, NESCent, Durham, NC.
201 I	Infectious Disease Epidemiology Seminar, Harvard School of Public Health, Boston, MA.
2010	Departmental Seminar, Department of Ecology and Evolutionary Biology, University of Michigan, Ann Arbor, MI.
2010	Phyloseminar.org, webcast.
2010	Departmental Seminar, Department of Ecology and Evolutionary Biology, University of Michigan, Ann Arbor, MI.
2009	Departmental Seminar, School of Computational Science, Florida State University, Tallahassee, FL.
2006	NKS Wolfram Science Conference, Washington, DC.
	Conference abstracts
2019	Epidemics ⁷ , Charleston, SC.
2017	Epidemics ⁶ , Sitges, Spain.
2017	Ecological Society of America Annual Meeting, Portland, OR.
2015	Epidemics ⁵ , Clearwater Beach, FL.
2013	Mathematical and Computational Evolutionary Biology, Montpellier, France.
2013	HIV Dynamics and Evolution, Utrecht, Netherlands.
2012	Society for Molecular Biology and Evolution Annual Meeting, Dublin, Ireland.
2012	Population Genetics Group, Nottingham, UK.
2011	Epidemics ³ , Boston, MA. (Award for Best Oral Presentation by a Young Investigator.)
2010	Ecology and Evolution of Infectious Diseases, Cornell University, Ithaca, NY.
2008	Society for Molecular Biology and Evolution Annual Meeting, Barcelona, Spain. Poster presentation.
2007	Society for Molecular Biology and Evolution Annual Meeting, Halifax, Canada. Poster presentation. (Graduate student poster award)
	Outreach
2020	COVID-19 pandemic scientific communication and outreach through Twitter to > 300k followers.
2015	Science outreach video "Tracking down the sneeze that started seasonal flu" published on Vox.com.