Computational Biology Program Fred Hutchinson Cancer Research Center 1100 Fairview Ave. N., S2-105 Mail stop: S2-140 Seattle, WA 98109-1024 matsen@fredhutch.org http://matsen.fredhutch.org/

Positions	Investigator , Howard Hughes Medical Institute. October 2021 – Present
	Professor , Fred Hutchinson Cancer Research Center, Seattle. April 2020 – Present
	Associate Member , Fred Hutchinson Cancer Research Center, Seattle. April 2015 – March 2020
	Affiliate Associate Professor, Genome Sciences, University of Washington, Seattle. March 2017 – Present
	Affiliate Associate Professor, Statistics, University of Washington, Seattle. June 2016 – Present
	Assistant Member, Fred Hutchinson Cancer Research Center, Seattle. August 2010 – March 2015
	Miller Postdoctoral Scholar , UC Berkeley, Berkeley, California. July 2007 – July 2010
	Allan Wilson Postdoctoral Scholar , University of Canterbury, Christchurch, New Zealand. July 2006 – June 2007
Education	Harvard University , Cambridge, MA. Ph.D. in Mathematics, June 2006.
	Université de Paris 13 , Paris, France. D.E.A. in Mathematics, June 2001.
	Stanford University , Stanford, CA. B.S. in Mathematics with honors, June 2000.
Grants:	HHMI Investigator Award (2021/10/01 – 2028/09/31)
primary investigator	R01 AI162611 (2021/07/01 – 2026/06/31) Fast and flexible Bayesian phylogenetics via modern machine learning.
	R01 AI146028 (2019/06/01 – 2024/05/31) Blending deep learning with probabilistic mechanistic models to predict and understand the evolution and function of adaptive immune receptors.
	HHMI-Simons Faculty Scholar Award (11/01/2016 – 11/01/2021)
	NSF Award 1564137 (2016/07/01 – 2020/06/30) Enabling phylogenetic inference for modern data sets. Co-PI with Vladimir Minin.
	R01 GM1132461 (2014/08/01 – 2019/08/01) Leveraging deep sequencing data to understand antibody maturation.
	NSF Award 1223057 (2012/09/01 – 2017/08/31) New theorems and algorithms for comprehensive analysis of metagenomic data via statis- tical phylogenetics. Co-PI with Aaron Darling.

R01 HG005966 (2010/09/27 - 2013/06/30) Novel Computational Tools for Studying the Human Microbiome. Co-PI with David Fredricks. **ITHS pilot grant** (2014/02/01 – 2015/02/01) Inferring statistical models of antibody somatic hypermutation from high-throughput sequencing data **CFAR New Investigator Award** (2014/01/01 – 2016/01/01) Machine learning, informed by structural knowledge, to expand our knowledge of APOBEC3 specificity and improve hypermutation inference R01 AI138709 (2018/02/07 - 2023/02/07) Grant support: co-investigator Characterizing the Broad Antibody Response to HIV Superinfection PI: Julie Overbaugh. Our role: our group will design and perform all the B cell receptor sequence analysis for this project. R01 AI120961 (2016/01/01 - 2020/12/31) Defining the Infant Immune Response to HIV. PI: Julie Overbaugh.

Our role: our group will design and perform all the B cell receptor sequence analysis for this project.

Bill and Melinda Gates Foundation OPP1110049 (2014/08/31 – 2016/07/31) Biostatistical, Computational Biology, and Mathematical Modeling for the Assessment of Immune Correlates of Protection in the HVTN 701 and 702 Efficacy Trials in South Africa. PI: Peter Gilbert.

Our role: Improve methods for reconstruction of HIV molecular sequences which found infection using later time point samples.

R37 AI038518-16A2 (2010/03/01 – 2015/02/28) Early and Reinfection in High Risk Women. PI: Julie Overbaugh. Our role: our group has designed and performed all the bioinformatics for this project.

	R21 AI104449-01 (2013/02/15 – 2015/01/31)
	PrEP Exposure during HIV-1 Infection and the Risk of Low-Frequency ARV Resistance. PI: Dara Lehman.
	Our role: our group has designed and performed all the bioinformatics for this project.
	U54GM111274-01 (2014/09/12 – 2019/06/30) Center for Statistics and Quantitative Infectious Diseases. PI: Elizabeth Halloran.
	Our role: design and implement computational phylogenetic algorithms for pathogen se- quence data.
Awards	HHMI-Simons faculty scholar (November 2016 – October 2021) Research support for five years.
	Miller fellow (July 2007 – July 2010) Support for three years of postdoctoral research at UC Berkeley.
	Allan Wilson postdoctoral fellow (July 2006 – June 2007) Support for one year of postdoctoral research.
	NSF graduate research fellow (July 2003 – June 2006) Support for three years of study.

Harvard University certificate of distinction in teaching (Fall 2004) For teaching second semester calculus.

Jean D. Valpine fellowship (September 2001 – June 2003) Support for two years of study.
Fulbright fellowship (July 2000 – June 2001) To study mathematics in Paris.
phylogenetics, innate and adaptive immunity, infectious disease, human microbiome, Markov processes, coalescent theory, molecular evolution, macroevolutionary models, discrete ge- ometry, combinatorics, Bayesian statistics, bioinformatics.
Masters and PhD studentsAmrit Dhar2016-2019Data Scientist, GoogleJean Feng2016-2020Assistant professor (tenure track), UCSFBranden Olson2015-2020Data Scientist, WorkdayHassan Nasif2020-2022Data Scientist, MicrosoftAndy Magee2017-2021Senior Data Scientist, CDCWill DeWitt2017-2022Postdoc, UC BerkeleyThayer Fisher2015-2017Assistant professor (tenure track), University of DelawareDavid Shaw2015-2017Assistant professor (tenure track), University of DelawareJulia Fukuyama2017-2018Assistant professor (tenure track), University of IndianaChris Whidden2013-2018Assistant professor (tenure track), Dalhousie UniversityCheng Zhang2016-2019Assistant professor (tenure track), Peking UniversityMichael Karcher2019-2021Assistant professor (tenure track), Muhlenberg CollegeSeong-Hwan Jun2019-2021Research assistant professor, University of Rochester
 PhD committee member: Miguel Parades, since 2023. PhD committee member: Marlin Figgins, Alex Willcox and Tim Yu, since 2022. PhD committee member: Malika Hale, William Hannon, Zachary Montague, and Elena Violeta Romero, since 2021. PhD committee member: Frances Welsh and Zak Yaffe, since 2020. PhD committee member: Caelan Radford, since 2019. Conference Co-organizer: Advances in Immune Repertoire Measurement and Analysis, Chan-Zuckerberg Biohub, January 2019. PhD committee member: Allison Black, John Huddleston and Anna Minkina, since 2018. PhD committee member: Sarah Hilton and Sydney Bell, since 2016. Invited session organizer: Bernoulli Society World Congress in Probability and Statistics, July 2016. Mentor: Summer Undergraduate Research Program, Summer 2016. Mentor: Summer High School Internship Program, Summer 2014-6. Organizer: Spotlight session on next generation phylogenetic inference, Joint Evolution Meetings, June 2016. Leader: Tools and Resources Working Group, Adaptive Immune Receptor Repertoire Community. (http://airr-community.org) Founder and organizer: B-T.CR online immunogenetics discussion forum (http://b-t.cr/), since February 2016. Associate editor: PLOS Computational Biology, since December 2015.

Editor: Special issue of *Philosophical Transactions of the Royal Society B* on antibody dynamics, 2015.

Invited program committee member: International Conference on Intelligent Systems for Molecular Biology (ISMB) / European Conference on Computational Biology (ECCB), 2009 and 2015.

Member: Statistical Genetics faculty group at the University of Washington, since October 2014.

Founder: *fredhutch.io* initiative to improve access to and education on scientific computing at the Fred Hutch (http://fredhutch.io/), since July 2014.

Founder and organizer: *Phylobabble* online phylogenetics discussion forum (http://phylobabble.org/), March 2014 – 2022.

Editorial board: Systematic Biology, since February 2013.

Host for professor on sabbatical: Vladimir Minin, academic year 2013-4.

Founder and organizer: *Phyloseminar* online phylogenetics seminar (http://phyloseminar.org/), since November 2009.

Contributor: *Biopython project* (http://github.com/biopython/), since December 2010.

Conference co-organizer: *Molecular Evolution and Phylogenetics*, Statistical and Applied Mathematical Sciences Institute (SAMSI), Spring 2009.

Paper and grant reviewer (selected list):

Publications Gabriel E Boyle, Katherine Sitko, Jared G Galloway, Hugh K Haddox, Aisha Haley Bianchi, Ajeya Dixon, Raine E S Thomson, Riddhiman K Garge, Allan E Rettie, Alan Rubin, Renee C Geck, Elizabeth M J Gillam, William S DeWitt, Frederick A <u>Matsen</u>, and Douglas M Fowler. Deep mutational scanning of CYP2C19 reveals a substrate specificity-abundance tradeoff. October 2023.

Hugh K. Haddox, Jared G. Galloway, Bernadeta Dadonaite, Jesse D. Bloom, Frederick A. <u>Matsen</u> IV, and William S. DeWitt. Jointly modeling deep mutational scans identifies shifted mutational effects among sars-cov-2 spike homologs. August 2023.

Zak A Yaffe, Shilei Ding, Kevin Sung, Vrasha Chohan, Lorie Marchitto, Laura Doepker, Duncan Ralph, Ruth Nduati, Frederick A <u>Matsen</u>, Andrés Finzi, and Julie Overbaugh. Reconstruction of a polyclonal ADCC antibody repertoire from an HIV-1 Non-Transmitting mother. *iScience*, page 106762, April 2023.

Michael Karcher, Cheng Zhang, and Frederick A Matsen, IV. Variational Bayesian supertrees. *arXiv*, April 2021.

Christiaan Swanepoel, Mathieu Fourment, Xiang Ji, Hassan Nasif, Marc A Suchard, Frederick A <u>Matsen</u>, IV, and Alexei Drummond. TreeFlow: probabilistic programming and automatic differentiation for phylogenetics. *arXiv*, November 2022.

Karthik Gangavarapu, Xiang Ji, Guy Baele, Mathieu Fourment, Philippe Lemey, Frederick A <u>Matsen</u> IV, and Marc A Suchard. Many-core algorithms for high-dimensional gradients on phylogenetic trees. *Bioinformatics*, 40(2), February 2024.

Andrew Magee, Michael Karcher, Frederick A <u>Matsen</u> IV, and Volodymyr M Minin. How trustworthy is your tree? bayesian phylogenetic effective sample size through the lens of monte carlo error. *Bayesian Analysis*, 19(2):565–593, June 2024.

Magdalena L Russell, Carolyn S Fish, Sara Drescher, Noah A J Cassidy, Pritha Chanana, Sarah Benki-Nugent, Jennifer Slyker, Dorothy Mbori-Ngacha, Rose Bosire, Barbra Richardson, Dalton Wamalwa, Elizabeth Maleche-Obimbo, Julie Overbaugh, Grace John-Stewart, Frederick A <u>Matsen</u>, 4th, and Dara A Lehman. Using viral sequence diversity to estimate time of HIV infection in infants. *PLoS Pathog.*, 19(12):e1011861, December 2023.

Jay Lubow, Lisa M Levoir, Duncan K Ralph, Laura Belmont, Maya Contreras, Catiana H Cartwright-Acar, Caroline Kikawa, Shruthi Kannan, Edgar Davidson, Veronica Duran, David E Rebellon-Sanchez, Ana M Sanz, Fernando Rosso, Benjamin J Doranz, Shirit Einav, Frederick A <u>Matsen</u> IV, and Leslie Goo. Single B cell transcriptomics identifies multiple isotypes of broadly neutralizing antibodies against flaviviruses. *PLoS Pathog.*, 19(10):e1011722, October 2023.

Seong-Hwan Jun, Hassan Nasif, Chris Jennings-Shaffer, David H Rich, Anna Kooperberg, Mathieu Fourment, Cheng Zhang, Marc A Suchard, and Frederick A <u>Matsen</u> IV. A topologymarginal composite likelihood via a generalized phylogenetic pruning algorithm. *Algorithms Mol. Biol.*, 18(1):10, July 2023.

Will Dumm, Mary Barker, William Howard-Snyder, William S DeWitt III, and Frederick A <u>Matsen</u> IV. Representing and extending ensembles of parsimonious evolutionary histories with a directed acyclic graph. *J. Math. Biol.*, 87(5):75, October 2023.

Jared G Galloway, Kevin Sung, Samuel S Minot, Meghan E Garrett, Caitlin I Stoddard, Alexandra C Willcox, Zak A Yaffe, Ryan Yucha, Julie Overbaugh, and Frederick A <u>Matsen</u> IV. phippery: a software suite for phip-seq data analysis. *Bioinformatics*, 39(10):btad583, September 2023.

Caitlin I Stoddard, Kevin Sung, Zak A Yaffe, Haidyn Weight, Guillaume Beaudoin-Bussières, Jared Galloway, Soren Gantt, Judith Adhiambo, Emily R Begnel, Ednah Ojee, Jennifer Slyker, Dalton Wamalwa, John Kinuthia, Andrés Finzi, Frederick A <u>Matsen</u> IV, Dara A Lehman, and Julie Overbaugh. Elevated binding and functional antibody responses to SARS-CoV-2 in infants versus mothers. *Nat. Commun.*, 14(1):4864, August 2023.

Mathieu Fourment, Christiaan J Swanepoel, Jared G Galloway, Xiang Ji, Karthik Gangavarapu, Marc A Suchard, and Frederick A <u>Matsen</u>, Iv. Automatic differentiation is no panacea for phylogenetic gradient computation. *Genome Biol. Evol.*, 15(6), June 2023.

Magdalena L Russell, Noah Simon, Philip Bradley, and Frederick A <u>Matsen</u>, IV. Statistical inference reveals the role of length, GC content, and local sequence in V(D)J nucleotide trimming. *Elife*, 12:e85145, May 2023.

Timothy C Yu, Zorian T Thornton, William W Hannon, William S DeWitt, Caelan E Radford, Frederick A <u>Matsen</u> IV, and Jesse D Bloom. A biophysical model of viral escape from polyclonal antibodies. *Virus Evol*, 8(2):veac110, December 2022.

Branden J Olson, Stefan A Schattgen, Paul G Thomas, Philip Bradley, and Frederick A <u>Matsen</u>, IV. Comparing T cell receptor repertoires using optimal transport. *PLoS Comput. Biol.*, 18(12):e1010681, December 2022.

Frederick A <u>Matsen</u> IV and Peter L Ralph. Enabling inference for Context-Dependent models of mutation by bounding the propagation of dependency. *J. Comput. Biol.*, 29(8):802–824, August 2022.

Caitlin I Stoddard, Kevin Sung, Ednah Ojee, Judith Adhiambo, Emily R Begnel, Jennifer Slyker, Soren Gantt, Frederick A <u>Matsen</u>, John Kinuthia, Dalton Wamalwa, Julie Overbaugh, and Dara A Lehman. Distinct antibody responses to endemic coronaviruses preand Post-SARS-CoV-2 infection in kenyan infants and mothers. *Viruses*, 14(7):1517, July 2022.

Alexandra C Willcox, Kevin Sung, Meghan E Garrett, Jared G Galloway, Jesse H Erasmus, Jennifer K Logue, David W Hawman, Helen Y Chu, Kim J Hasenkrug, Deborah H Fuller, Frederick A <u>Matsen</u>, Iv, and Julie Overbaugh. Detailed analysis of antibody responses to SARS-CoV-2 vaccination and infection in macaques. *PLoS Pathog.*, 18(4):e1010155, April 2022.

Duncan K Ralph and Frederick A <u>Matsen</u>, 4th. Inference of B cell clonal families using heavy/light chain pairing information. *PLoS Comput. Biol.*, 18(11):e1010723, November 2022.

Magdalena L Russell, Aisha Souquette, David M Levine, Stefan A Schattgen, E Kaitlynn Allen, Guillermina Kuan, Noah Simon, Angel Balmaseda, Aubree Gordon, Paul G Thomas, Frederick A <u>Matsen</u>, IV, and Philip Bradley. Combining genotypes and T cell receptor distributions to infer genetic loci determining V(D)J recombination probabilities. *Elife*, 11:e73475, March 2022.

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Hannah L Itell, Haidyn Weight, Carolyn S Fish, Jennifer K Logue, Nicholas Franko, Caitlin R Wolf, Denise J McCulloch, Jared Galloway, Frederick A <u>Matsen</u>, 4th, Helen Y Chu, and Julie Overbaugh. SARS-CoV-2 antibody binding and neutralization in dried blood spot eluates and paired plasma. *Microbiol Spectr*, page e0129821, October 2021.

Mackenzie M Shipley, Vidya Mangala Prasad, Laura E Doepker, Adam S Dingens, Duncan K Ralph, Elias Harkins, Amrit Dhar, Dana Arenz, Vrasha Chohan, Haidyn Weight, Kishor Mandaliya, Jesse D Bloom, Frederick <u>Matsen</u>, IV, Kelly K Lee, and Julie M Overbaugh. Functional development of a v3/glycan-specific broadly neutralizing antibody isolated from a case of HIV superinfection. *Elife*, 10:e68110, July 2021.

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Meghan E Garrett, Jared Galloway, Helen Y Chu, Hannah L Itell, Caitlin I Stoddard, Caitlin R Wolf, Jennifer K Logue, Dylan McDonald, Haidyn Weight, Frederick A <u>Matsen</u>, and Julie Overbaugh. High-resolution profiling of pathways of escape for SARS-CoV-2 spike-binding antibodies. *Cell*, May 2021.

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Jean Feng, William S DeWitt, III, Aaron McKenna, Noah Simon, Amy D Willis, and Frederick A <u>Matsen</u>, IV. Estimation of cell lineage trees by maximum-likelihood phylogenetics. *Annals of Applied Statistics*, 15(1):343–362, March 2021.

Mark D Pankau, Daniel B Reeves, Elias Harkins, Keshet Ronen, Walter Jaoko, Kishor Mandaliya, Susan M Graham, R Scott McClelland, Frederick A <u>Matsen</u>, IV, Joshua T Schiffer, Julie Overbaugh, and Dara A Lehman. Dynamics of HIV DNA reservoir seeding in a cohort of superinfected kenyan women. *PLoS Pathog.*, 16(2):e1008286, February 2020.

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Amrit Dhar, Duncan K Ralph, Vladimir N Minin, and Frederick A <u>Matsen</u> IV. A bayesian phylogenetic hidden markov model for B cell receptor sequence analysis. *PLoS Comput. Biol.*, 16(8):e1008030, August 2020.

Cheng Zhang, Vu Dinh, and Frederick A <u>Matsen</u>. Nonbifurcating phylogenetic tree inference via the adaptive LASSO. *J. Am. Stat. Assoc.*, pages 1–16, June 2020.

Julia Fukuyama, Branden J Olson, and Frederick A <u>Matsen</u>, 4th. Lack of evidence for a substantial rate of templated mutagenesis in B cell diversification. *J. Immunol.*, July 2020.

Chris Whidden, Brian C Claywell, Thayer Fisher, Andrew F Magee, Mathieu Fourment, and Frederick A <u>Matsen</u>, IV. Systematic exploration of the high likelihood set of phylogenetic tree topologies. *Syst. Biol.*, 69(2):280–293, March 2020.

Mathieu Fourment, Andrew F Magee, Chris Whidden, Arman Bilge, Frederick A <u>Matsen</u>, IV, and Vladimir N Minin. 19 dubious ways to compute the marginal likelihood of a phylogenetic tree topology. *Syst. Biol.*, 69(2):209–220, March 2020.

Laura E Doepker, Cassandra A Simonich, Duncan Ralph, Mackenzie M Shipley, Meghan Garrett, Theodore Gobillot, Vladimir Vigdorovich, D Noah Sather, Ruth Nduati, Frederick A <u>Matsen</u>, 4th, and Julie M Overbaugh. Diversity and function of maternal HIV-1-specific antibodies at the time of vertical transmission. *J. Virol.*, February 2020.

William Lees, Christian E Busse, Martin Corcoran, Mats Ohlin, Cathrine Scheepers, Frederick A <u>Matsen</u>, Gur Yaari, Corey T Watson, AIRR Community, Andrew Collins, and Adrian J Shepherd. OGRDB: a reference database of inferred immune receptor genes. *Nucleic Acids Res.*, 48(D1):D964–D970, January 2020.

Jean Feng, David A Shaw, Vladimir N Minin, Noah Simon, and Frederick A <u>Matsen</u>, IV. Survival analysis of DNA mutation motifs with penalized proportional hazards. *Ann. Appl. Stat.*, 13(2):1268–1294, June 2019.

Kristian Davidsen, Branden J Olson, William S DeWitt, III, Jean Feng, Elias Harkins, Philip Bradley, and Frederick A <u>Matsen</u>, IV. Deep generative models for T cell receptor protein sequences. *Elife*, 8:e46935, September 2019.

Cassandra A Simonich, Laura Doepker, Duncan Ralph, James A Williams, Amrit Dhar, Zak Yaffe, Lauren Gentles, Christopher T Small, Brian Oliver, Vladimir Vigdorovich, Vidya Mangala Prasad, Ruth Nduati, D Noah Sather, Kelly K Lee, Frederick A <u>Matsen</u> IV, and Julie Overbaugh. Kappa chain maturation helps drive rapid development of an infant HIV-1 broadly neutralizing antibody lineage. *Nat. Commun.*, 10(1):2190, May 2019.

David A Shaw, Vu C Dinh, and Frederick A <u>Matsen</u> IV. Joint maximum likelihood of phylogeny and ancestral states is not consistent. *Mol. Biol. Evol.*, May 2019.

Lam Si Tung Ho, Vu Dinh, Frederick A <u>Matsen</u>, 4th, and Marc A Suchard. On the convergence of the maximum likelihood estimator for the transition rate under a 2-state symmetric model. *J. Math. Biol.*, 80(4):1119–1138, March 2020.

Cheng Zhang and Frederick A <u>Matsen</u>, IV. Variational bayesian phylogenetic inference. In *International Conference on Learning Representations (ICLR)*, 2019.

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Amrit Dhar, Kristian Davidsen, Frederick A <u>Matsen</u>, 4th, and Vladimir N Minin. Predicting B cell receptor substitution profiles using public repertoire data. *PLoS Comput. Biol.*, 14(10):e1006388, October 2018.

Kristian Davidsen and Frederick A <u>Matsen</u>. Benchmarking tree and ancestral sequence inference for B cell receptor sequences. *Front. Immunol.*, 9:2451, 2018.

William S DeWitt, Anajane Smith, Gary Schoch, John A Hansen, Frederick A <u>Matsen</u>, and Philip Bradley. Human T cell receptor occurrence patterns encode immune history, genetic background, and receptor specificity. *Elife*, 7:e38358, August 2018.

Vu Dinh, Lam Si Tung Ho, Marc A Suchard, and Frederick A <u>Matsen</u> IV. Consistency and convergence rate of phylogenetic inference via regularization. *Ann. Stat.*, 46(4):1481–1512, August 2018.

Branden J Olson and Frederick A <u>Matsen</u> IV. The Bayesian optimist's guide to adaptive immune receptor repertoire analysis. *Immunol. Rev.*, 284(1):148–166, June 2018.

William S DeWitt, Luka Mesin, Gabriel D Victora, Vladimir N Minin, and Frederick A <u>Matsen</u> IV. Using genotype abundance to improve phylogenetic inference. *Mol. Biol. Evol.*, February 2018.

F A <u>Matsen</u> IV, S C Billey, A Kas, and M Konvalinka. Tanglegrams: A reduction tool for mathematical phylogenetics. *IEEE/ACM Trans. Comput. Biol. Bioinform.*, 15(1):343–349, January 2018.

C Whidden and F <u>Matsen</u> IV. Efficiently inferring pairwise subtree prune-and-regraft adjacencies between phylogenetic trees. In 2018 Proceedings of the Fifteenth Workshop on Analytic Algorithmics and Combinatorics (ANALCO), Proceedings, pages 77–91. Society for Industrial and Applied Mathematics, January 2018.

Brian C Claywell, Vu Dinh, Mathieu Fourment, Connor O McCoy, and Frederick A <u>Matsen</u> IV. A surrogate function for one-dimensional phylogenetic likelihoods. *Mol. Biol. Evol.*, 35(1):242–246, January 2018.

Vu Dinh and Frederick A <u>Matsen</u> IV. The shape of the one-dimensional phylogenetic likelihood function. *Ann. Appl. Probab.*, 27(3):1646–1677, June 2017.

Vu Dinh, Arman Bilge, Cheng Zhang, and Frederick A <u>Matsen</u> IV. Probabilistic path Hamiltonian Monte Carlo. In *International Conference on Machine Learning*, pages 1009–1018. PMLR, 17 July 2017.

Vu Dinh, Aaron E Darling, and Frederick A <u>Matsen</u> IV. Online Bayesian phylogenetic inference: theoretical foundations via Sequential Monte Carlo. *Syst. Biol.*, December 2017.

Mathieu Fourment, Brian C Claywell, Vu Dinh, Connor McCoy, Frederick A <u>Matsen</u> IV, and Aaron E Darling. Effective online Bayesian phylogenetics via Sequential Monte Carlo with guided proposals. *Syst. Biol.*, November 2017.

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Felix Breden, Eline T Luning Prak, Bjoern Peters, Florian Rubelt, Chaim A Schramm, Christian E Busse, Jason A Vander Heiden, Scott Christley, Syed Ahmad Chan Bukhari, Adrian Thorogood, Frederick A <u>Matsen</u> IV, Yariv Wine, Uri Laserson, David Klatzmann, Daniel C Douek, Marie-Paule Lefranc, Andrew M Collins, Tania Bubela, Steven H Kleinstein, Corey T Watson, Lindsay G Cowell, Jamie K Scott, and Thomas B Kepler. Reproducibility and reuse of adaptive immune receptor repertoire data. *Front. Immunol.*, 8:1418, 2017.

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Chris Whidden and Frederick A <u>Matsen</u> IV. Quantifying MCMC exploration of phylogenetic tree space. *Systematic Biology*, 64(3):472–491, May 2015.

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