Karen A. Cranston

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Education

2002-2007	PhD , Department of Medical Genetics, University of Alberta, Edmonton, AB, Canada. Thesis title: "Inferring, testing and summarizing a posterior distribution of phylogenies". Supervisor: Dr. Bruce Rannala.
1992-1996	B.Sc. (Honours, First Class), Genetics. University of Manitoba, Winnipeg, MB, Canada

Employment

2017-2018	Technical Advisor, Bioinformatics , Biological Informatics Centre of Excellence, Agriculture and Agri-food Canada, Ottawa, ON
2015-2017	Research Scientist, Department of Biology, Duke University, Durham, NC
2010-2015	Training Coordinator and Bioinformatics Project Manager , National Evolutionary Synthesis Center (NESCent), Durham, NC
2010-2015	Training Coordinator and Bioinformatics Project Manager , National Evolutionary Synthesis Center (NESCent), Durham, NC
2009-2010	Postdoctoral Research Associate , Biodiversity Synthesis Center / Encyclopedia of Life at the Field Museum of Natural History, Chicago, IL
1997-2002	Laboratory technician in cell culture, protein chemistry and analytical method validation, Cangene Corporation, Winnipeg, MB
1996-1997	Technical writer Integrated Engineering Software, Winnipeg, MB.

Peer-reviewed publications

McTavish, E. J., Drew, B. T., Redelings, B., & Cranston, K. A. (2017). How and Why to Build a Unified Tree of Life. BioEssays, 39(11). https://doi.org/10.1002/bies.201700114

Wilson, G., et al., (2017). Good Enough Practices in Scientific Computing. PLoS Comput Biol 13(6): e1005510 https://doi.org/10.1371/journal.pcbi.1005510

Rees, J., & Cranston, K. (2017). Automated assembly of a reference taxonomy for phylogenetic data synthesis. Biodiversity Data Journal, 5, e12581. https://doi.org/10.3897/BDJ.5.e12581

McTavish, E. J., Hinchliff, C. E., Allman, J. F., Brown, J. W., Cranston, K. A., Holder, M. T., ... & Smith, S. A. (2015). Phylesystem: a git-based data store for community curated phylogenetic estimates. Bioinformatics, btv276. http://dx.doi.org/10.1093/bioinformatics/btv276

Hinchliff, C. E., Smith, S. A., Allman, J. F., Burleigh, J. G., Chaudhary, R., Coghill, L. M., ... & Cranston, K. A. (2015). Synthesis of phylogeny and taxonomy into a comprehensive tree of life. Proceedings of the National Academy of Sciences, 112(41), 12764-12769. http://dx.doi.org/10.1073/pnas.1423041112. In the Altmetrics Top 100 for 2015 and in top 5% overall; 53 news reports; 832 Tweets, 14 blogs; https://www.altmetric.com/details/4529894

Ksepka, D.T., Parham, J.F., Allman, J.F., Benton, M.J., Carrano, M.T., Cranston, K.A., ... & Warnock, R.C.M. (2015) The Fossil Calibration Database, A New Resource for Divergence Dating. Systematic Biology. http://dx.doi.org/10.1093/sysbio/syv025

Teal, T. K., Cranston, K. A., Lapp, H., White, E., Wilson, G., Ram, K., Pawlik, A. (2015) Data Carpentry: Workshops to Increase Data Literacy for Researchers. *International Journal of Digital Curation*. 10(1):135-143. http://dx.doi.org/10.2218/ijdc.v10i1.351

Cranston, K. A., Harmon, L. J., O'Leary, M. A., Lisle, C. (2014) Best Practices for Data Sharing in Phylogenetic Research. *PLOS Currents Tree of Life*. Jun 19. Edition 1. http://dx.doi.org/10.1371/currents.tol.bf01eff4a6b60ca4825c69293dc59645

Stoltzfus, A., Lapp, H., Matasci, N., Deus, H., Sidlauskas, B., Zmasek, C. M., ... & Jordan, G. (2013). Phylotastic! Making tree-of-life knowledge accessible, reusable and convenient. *BMC bioinformatics*, 14(1), 158. http://dx.doi.org/10.1186/1471-2105-14-158

Rodrigo, A., Alberts, S., Cranston, K., Kingsolver, J., Lapp, H., McClain, C., ... & Wiegmann, B. (2013). Science incubators: synthesis centers and their role in the research ecosystem. PloS biology, 11(1), e1001468. http://dx.doi.org/10.1371/journal.pbio.1001468

Goff, S. A., Vaughn, M., McKay, S., Lyons, E., Stapleton, A. E., Gessler, D., ... & Kleibenstein, D. J. (2011). The iPlant collaborative: cyberinfrastructure for plant biology. *Frontiers in plant science*, 2. http://dx.doi.org/10.3389/fpls.2011.0003

Evans, M. E., Hearn, D. J., Theiss, K. E., Cranston, K., Holsinger, K. E., & Donoghue, M. J. (2011). Extreme environments select for reproductive assurance: evidence from evening primroses (Oenothera). *New Phytologist*, 191(2), 555-563. http://dx.doi.org/10.1111/j.1469-8137.2011.03697.x

Ammiraju, J. S., Fan, C., Yu, Y., Song, X., Cranston, K. A., Pontaroli, A. C., ... & Wing, R. A. (2010). Spatiotemporal patterns of genome evolution in allotetraploid species of the genus Oryza. *The Plant Journal*, 63(3), 430-442. http://dx.doi.org/10.1111/j.1365-313X.2010.04251.x

Cranston, K. A. (2010) Quantifying gene tree incongruence across varying phylogenetic depths. In L. Lacey Knowles, Laura S. Kubatko (Eds.) *Estimating Species Trees: Practical and Theoretical Aspects*, Wiley-Blackwell.

Cranston, K. A., Hurwitz, B., Sanderson, M. J., Ware, D., Wing, R. A., & Stein, L. (2010). Phylogenomic analysis of BAC-end sequence libraries in Oryza (Poaceae). Systematic botany, 35(3), 512-523. http://dx.doi.org/10.1600/036364410792495872

Cranston, K. A., Hurwitz, B., Ware, D., Stein, L., & Wing, R. A. (2009). Species trees from highly incongruent gene trees in rice. *Systematic Biology* 58(5), 489-500. http://dx.doi.org/10.1093/sysbio/syp054

Sanderson, M. J., Boss, D., Chen, D., Cranston, K. A., & Wehe, A. (2008). The PhyLoTA Browser: processing GenBank for molecular phylogenetics research. *Systematic Biology* 57(3), pp. 335-346. http://dx.doi.org/10.1080/10635150802158688

Cranston, K. A., & Rannala, B. (2007). Summarizing a posterior distribution of trees using agreement subtrees. *Systematic Biology* 56(4), pp. 578-590.

http://dx.doi.org/10.1080/10635150701485091

Cranston, K., & Rannala, B. (2005). Closing the gap between rocks and clocks. Heredity, 94(5), 461-462. http://dx.doi.org/10.1038/sj.hdy.6800644

Commentaries, preprints, other non-peer reviewed

(published preprints not shown)

Katz, D.S., Choi, S.T., Wilkins-Diehr, N., Hong, N.C., Venters, C.C., ... & Littauer, R. Report on the Second Workshop on Sustainable Software for Science: Practice and Experiences (WSSSPE2). arXiv preprint arXiv:1507.01715

Katz, D. S., Allen, G., Hong, N. C., Cranston, K., Parashar, M., Proctor, D., ... & Wilkins-Diehr, N. (2014). Second Workshop on on Sustainable Software for Science: Practice and Experiences (WSSSPE2): Submission, Peer-Review and Sorting Process, and Results. arXiv preprint arXiv:1411.3464.

Vision, T., & Cranston, K. A. (2014) Open data for evolutionary synthesis: an introduction to the NESCent collection. Nature Scientific Data. doi:10.1038/sdata.2014.30

Cranston, K. A., Blackburn, D., Brown, J. W., Dececchi, A., Gardner, N., Greshake, B., ... & Wolfe, J. (2014): Simple rules for sharing phylogenetic data. figshare. doi:10.6084/m9.figshare.997763

Rees, J. A., Cranston, K. A., Lapp, H., & Vision, T. (2013): Response to GBIF request for consultation on data licenses. figshare. doi:10.6084/m9.figshare.799766

Cranston, K. A., Vision, T., O'Meara, B., & Lapp, H. (2013): A grassroots approach to software sustainability. figshare. doi:10.6084/m9.figshare.790739

Invited talks

- Synthesizing phylogeny and taxonomy into a comprehensive tree of life, Biodiversity Informatics (TDWG) (2014)
- Data and services for the tree of life, Commonwealth Scientific and Industrial Research Organisation (CSIRO) (2014)
- Technical and social challenges in synthesizing the tree of life, Agriculture Canada (2014)
- Enabling science with the tree of life, Carleton University (2014)
- Technical and social challenges in synthesizing the tree of life, http://phyloseminar.org (2014)
- Open Tree of Life: Synthesizing phylogenetic data into a tree of all life, University of North Carolina Charlotte (2013)
- Phylogenetics, phylogenomics and phyloinformatics: Finding evolutionary signals in large data sets', University of Toronto (2010)
- Automating phylogenetic tree inference from public sequence databases, University of Missouri, 2009
- Automating phylogenetic tree inference from public sequence databases, Botany meetings, 2009
- Species trees and gene trees from high-throughput sequence data in rice, University of Michigan, 2009
- A species tree from 2000 gene trees in rice (Oryza), Evolution, 2008
- The PhyLoTA browser: Processing GenBank for molecular phylogenetic research, George Washington University, 2008
- Species trees from gene trees in rice: When 2.5 million characters are not enough..., University of California Davis, 2008

Grants and Awards

- NSF Special Creativity Award: Collaborative Research: Automated and community-driven synthesis of the tree of life. NSF AVAToL (2014) \$1.4 million (\$1.1 million to Duke, PI Cranston, 2015-2017)
- NSF AVAToL: Collaborative Research: Automated and community-driven synthesis of the tree of life. \$5.76 million (\$927,000 to Duke, PI Cranston, 2012-2015)

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- The iPlant Collaborative; a cyberinfrastructure-centered community for a new plant biology
 subcontract to Field Museum of Natural History, (\$62000, PI Cranston, 2009-2010) and to
 Duke University (\$14500, PI Vision, 2011-2011)
- Dissertation Fellowship, University of Alberta (2006)
- Province of Alberta Graduate Student Fellowship (2006)
- Mary Louise Imrie Graduate Student Award, University of Alberta (2006)
- 75th Anniversary Graduate Student Award, Faculty of Medicine and Dentistry, University of Alberta (2004)
- Medical Sciences Graduate Research Assistantship, Faculty of Medicine and Dentistry, University of Alberta (2006 & 2004)

Professional activities

- Board of Directors: Open Bioinformatics Foundation (https://open-bio.org/), 2015-present
- Co-organizer: Reproducible science hackathon (https://github.com/Reproducible-Science-Curriculum/Reproducible-Science-Hackathon-Dec-08-2014), 2014
- Data Carpentry founding board member (http://datacarpentry.org), 2014-present
- Organization administrator for NESCent Google Summer of Code participation, 2011-2013
- Program committee, Workshop on Sustainable Software for Science: Practice and Experiences (WSSSPE), http://wssspe.researchcomputing.org.uk, 2014 and 2016
- Panelist, First Workshop on Sustainable Software for Science: Practice and Experiences (WSSSPE), http://wssspe.researchcomputing.org.uk, 2013
- Society of Systematic Biology council, 2012-2014
- Systematic Biology Editorial Board, 2011-present
- Leadership team, Informatics for Evolutionary Biology (iEvoBio, http://ievobio.org) conference, 2013-present
- Leadership team, Hackathons, Interoperability and Phylogenetics (HIP) NESCent working group, 2011-present
- Phyloinformatics Research Foundation board member, 2010-present
- Team lead, iPlant Tree Of Life working group on Tree Visualization, 2009-2012
- Co-organizer: Phyloinformatics VoCamp for development of an ontology for evolutionary biology, November 2009, Montpellier, France
- Invited participant: National Evolutionary Synthesis Center (NESCent) Hackathon on Evolutionary Database Interoperability, March 2009, Durham, NC, USA

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- Invited participant: iPlant Collaborative (http://iplantcollaborative.org) Grand Challenge Workshop on Assembling the Tree of Life to Enable the Plant Sciences, November 2009, Biosphere 2, Oracle, AZ
- Journal referee for: Systematic Biology, Trends Ecology and Evolution, PeerJ, Nucleic Acids Research, Journal of Open Research Software, PLOS Computational Biology, Molecular Biology and Evolution. See Publons reviewer profile for recent activity.
- National Science Foundation panelist, 2011 2014

Teaching

- Coordinator: NESCent Academy (http://academy.nescent.org): a series of hands-on summer workshops in Evolutionary Biology and Informatics, 2010-2014.
- Certified Instructor: Software and Data Carpentry (http://software-carpentry.org & http://datacarpentry.org), 2013-present
- Instructor: Computational biology workshops, Wuhan Institute of Virology and Fudan University, China, 2013
- Instructor: Phylogenetics workshops, Kenya Medical Research Institute, 2012
- Instructor: Phylogenetics workshop, Research Experience for Undergraduates (REU) Program, Field Museum of Natural History, 2010
- Instructor: Workshop in applied phylogenetics, Bodega Bay Marine Laboratory, 2006