## Siavash Mirarab

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Information UC San Diego e-mail: smirarab@ucsd.edu

9500 Gilman Drive, Mail code 0407 website: eceweb.ucsd.edu/~smirarab/ La Jolla, CA 92093-0407 google scholar: http://goo.gl/gEswQ5

Research Computational biology and Bioinformatics, Large-scale phylogenetics, Phylogenomics,

Interests Multiple sequence alignment, Metagenomics, Parallel computing, Big Data

ACADEMIC UC San Diego, San Diego, CA, United States

Positions Department of Electrical and Computer Engineering

Assistant Professor July 2015 – present

Education University of Texas at Austin, Austin, Texas, United States

Department of Computer Science

Ph.D January 2011 – July 2015

• Advisors: Prof. Tandy Warnow, Prof. Keshav Pingali

• Thesis: Novel scalable approaches for multiple sequence alignment and phylogenetic reconstruction

University of Waterloo, Waterloo, Ontario, Canada

Department of Electrical and Computer Engineering

Master of Science September 2006 – September 2008

• Advisors: Prof. Ladan Tahvildari

• Thesis: A Bayesian Framework for Software Regression Testing

University of Tehran, Tehran, Iran

Bachelor of Science

AWARDS

Department of Electrical and Computer Engineering

• Computer engineering, specialized in software engineering.

MAJOR Howard Hughes Medical Institute International Student Fellowship. 2012 - present

National Science and Engineering Research Council of Canada (NSERC), Canada Graduate Scholarship (CGS-D) converted to PGS-D outside Canada. **2011-2012** 

September 2001 – December 2005

University of Texas at Austin, MCD fellowship (I declined to take CGS instead)

Other University of Texas at Austin, Computer Science, Bert Kay Dissertation Award, 2015

University of Texas at Austin, College of Natural Sci., Dean's Excellence Award, 2011

University of Texas at Austin, Graduate Dean's Prestigious Fellowship Supplement

Award, 2011, 2012, 2013, 2014

#### **Publications**

- 1. James E Tarver, Mario Dos Reis, **Siavash Mirarab**, Raymond J Moran, Sean Parker, Joseph E O'Reilly, Benjamin L King, Mary J O'Connell, Robert J Asher, Tandy Warnow, Kevin J Peterson, Philip C J Donoghue, and Davide Pisani. The Interrelationships of Placental Mammals and the Limits of Phylogenetic Inference. *Genome biology and evolution*, page evv261, jan 2016.
- 2. Siavash Mirarab, Md. Shamsuzzoha Bayzid, Bastien Boussau, and Tandy Warnow. Response to Comment on "Statistical binning enables an accurate coalescent-based estimation of the avian tree". *Science*, 350(6257):171, 2015.
- 3. Joel Cracraft, Peter Houde, Simon Y W Ho, David P Mindell, Jon Fjeldså, Bent Lindow, Scott V Edwards, Carsten Rahbek, **Siavash Mirarab**, Tandy Warnow, M Thomas P Gilbert, Guojie Zhang, Edward L Braun, and Erich D Jarvis. Response to Comment on "Whole-genome analyses resolve early branches in the tree of life of modern birds". *Science*, 349(6255):1460, 2015.
- 4. Ruth Davidson, Pranjal Vachaspati, **Siavash Mirarab**, and Tandy Warnow. Phylogenomic species tree estimation in the presence of incomplete lineage sorting and horizontal gene transfer. *BMC Genomics*, 16(Suppl 10):S1, 2015.
- 5. Jed Chou, Ashu Gupta, Shashank Yaduvanshi, Ruth Davidson, Mike Nute, **Siavash Mirarab**, and Tandy Warnow. A comparative study of SVDquartets and other coalescent-based species tree estimation methods. *BMC Genomics*, 16(Suppl 10):S2, 2015.
- 6. **Siavash Mirarab** and Tandy Warnow. ASTRAL-II: coalescent-based species tree estimation with many hundreds of taxa and thousands of genes. *Bioinformatics*, 31(12):i44–i52, 2015.
- 7. Nam Nguyen, **Siavash Mirarab**, Keerthana Kumar, and Tandy Warnow. *Genome Biology*, 16(1):124, 2015.
- 8. Md Shamsuzzoha Bayzid, **Siavash Mirarab**, Bastien Boussau, and Tandy Warnow. Weighted Statistical Binning: Enabling Statistically Consistent Genome-Scale Phylogenetic Analyses. *PLoS ONE*, 10(6):e0129183, 2015.
- 9. (Erich D Jarvis, **Siavash Mirarab**)\*, 100 other authors, Edward L Braun, Tandy Warnow, Wang Jun, M Thomas P Gilbert, and Guojie Zhang. Wholegenome analyses resolve early branches in the tree of life of modern birds. *Science*, 346(6215):1320–1331, 2014.
- 10. **Siavash Mirarab**, Md. Shamsuzzoha Bayzid, Bastien Boussau, and Tandy Warnow. Statistical binning enables an accurate coalescent-based estimation of the avian tree. *Science*, 346(6215):1250463, 2014.

<sup>\*</sup> co-first authorship

- 11. (Norman J. Wickett, Siavash Mirarab)\*, Nam Nguyen, Tandy Warnow, Eric Carpenter, Naim Matasci, Saravanaraj Ayyampalayam, Michael Barker, J. Gordon Burleigh, Matthew A. Gitzendanner, Brad R. Ruhfel, Eric Wafula, Joshua P. Der, Sean W. Graham, Sarah Mathews, Michael Melkonian, Douglas E. Soltish, Pamela S. Soltish, Nicholas W. Miles, Carl J. Rothfels, Lisa Pokorny, A. Jonathan Shaw, Lisa DeGironimo, Dennis W. Stevenson, Barbara Surek, Juan Carlos Villarreal, Béatrice Rourev, Hervé Philippe, Claude W. dePamphilis, Tao Chen, Michael K. Deyholos, Regina S. Baucom, Toni M. Kutchan, Megan M. Augustin, Jun Wang, Yong Zhang, Zhijian Tian, Zhixiang Yan, Xiaolei Wu, Xiao Sun, Gane Ka-Shu Wong, and James Leebens-Mack. Phylotranscriptomic analysis of the origin and early diversification of land plants. Proceedings of the National Academy of Sciences (PNAS), 111(45):E4859–4868, 2014.
- 12. **Siavash Mirarab**, Md. Shamsuzzoha Bayzid, and Tandy Warnow. Evaluating summary methods for multi-locus species tree estimation in the presence of incomplete lineage sorting. *Systematic Biology*, doi: 10.1093/sysbio/syu063, 2014.
- 13. **Siavash Mirarab**, Nam Nguyen, Sheng Guo, Li-San Wang, Junhyong Kim, and Tandy Warnow. PASTA: Ultra-Large Multiple Sequence Alignment for Nucleotide and Amino-Acid Sequences. *Journal of Computational Biology*, 22(05):377–386, 2015.
- 14. **Siavash Mirarab**, Nam Nguyen, and Tandy Warnow. PASTA: ultra-large multiple sequence alignment. In *Proceedings of the 18th International Conference on Research in Computational Molecular Biology (RECOMB)*, pages 177–191, 2014.
- 15. **Siavash Mirarab**, Rezwana Reaz, Md. Shamsuzzoha Bayzid, Théo Zimmermann, M Shel Swenson, and Tandy Warnow. ASTRAL: Genome-Scale Coalescent-Based Species Tree. *Bioinformatics ECCB Special Issue*, 30(17):i541–i548, 2014.
- 16. Nam Nguyen, **Siavash Mirarab**, Bo Liu, Mihai Pop, and Tandy Warnow. TIPP: Taxonomic identification and phylogenetic profiling. *Bioinformatics*, doi: 10.1093/bioinformatics/btu721, 2014.
- 17. Théo Zimmermann, **Siavash Mirarab**, and Tandy Warnow. BBCA: improving the scalability of \*BEAST using random binning. *BMC Genomics*, 15(Suppl 6):S11, 2014.
- 18. Arlin Stoltzfus, Hilmar Lapp, Naim Matasci, Helena Deus, Brian Sidlauskas, Christian M Zmasek, Gaurav Vaidya, Enrico Pontelli, Karen Cranston, Rutger Vos, Campbell O Webb, Luke J Harmon, Megan Pirrung, Brian O'Meara, Matthew W Pennell, **Siavash Mirarab**, Michael S Rosenberg, James P Balhoff, Holly M Bik, Tracy A Heath, Peter E Midford, Joseph W Brown, Emily Jane

- McTavish, Jeet Sukumaran, Mark Westneat, Michael E Alfaro, Aaron Steele, and Greg Jordan. Phylotastic! Making tree-of-life knowledge accessible, reusable and convenient. *BMC bioinformatics*, 14(1):158, 2013.
- 19. Md. Shamsuzzoha Bayzid, **Siavash Mirarab**, and Tandy Warnow. Inferring optimal species trees under gene duplication and loss. In *Proceedings of the Pacific Symposium on Biocomputing (PSB)*, pages 250–61, 2013.
- 20. Nam Nguyen, **Siavash Mirarab**, and Tandy Warnow. MRL and SuperFine+MRL: new supertree methods. *Algorithms for Molecular Biology*, 7(1):3, 2012.
- 21. **Siavash Mirarab**, Nam Nguyen, and Tandy Warnow. SEPP: SATé-Enabled Phylogenetic Placement. In *Proceedings of the Pacific Symposium on Biocomputing* (*PSB*), pages 247–58, 2012.
- 22. **Siavash Mirarab**, Soroush Akhlaghi, and Ladan Tahvildari. Size-constrained regression test case selection using multicriteria optimization. *IEEE Transactions on Software Engineering*, 38(4):936–956, 2012.
- 23. **Siavash Mirarab** and Tandy Warnow. FastSP: linear time calculation of alignment accuracy. *Bioinformatics*, 27:3250–8, 2011.
- 24. Hyunsook Do, **Siavash Mirarab**, Ladan Tahvildari, and Gregg Rothermel. The effects of time constraints on test case prioritization: A series of controlled experiments. *IEEE Transactions on Software Engineering*, 36(5):593–617, 2010.
- 25. **Siavash Mirarab** and Ladan Tahvildari. An empirical study on bayesian network-based approach for test case prioritization. In *Proceedings of the 1st International Conference on Software Testing, Verification, and Validation (ICST)*, pages 278–287. IEEE, 2008.
- 26. **Siavash Mirarab**, Afshar Ganjali, Ladan Tahvildari, Shimin Li, Weining Liu, and Mike Morrissey. A requirement-based software testing framework: An industrial practice. In *Proceedings of the 24th International Conference on Software Maintenance (ICSM)*, pages 452–455. IEEE, 2008.
- 27. Hyunsook Do, **Siavash Mirarab**, Ladan Tahvildari, and Gregg Rothermel. An empirical study of the effect of time constraints on the cost-benefits of regression testing. In *Proceedings of the 16th ACM SIGSOFT International Symposium on Foundations of Software Engineering (FSE)*, pages 71–82. ACM, 2008.
- 28. Mehdi Amoui, Mazeiar Salehie, **Siavash Mirarab**, and Ladan Tahvildari. Adaptive action selection in autonomic software using reinforcement learning. In *Proceedings of the Fourth International Conference on Autonomic and Autonomous Systems (ICAS)*, pages 175–181. IEEE, 2008.

- 29. Siavash Mirarab and Ladan Tahvildari. A prioritization approach for software test cases based on bayesian networks. In Fundamental Approaches to Software Engineering (FASE), pages 276–290. Springer, 2007.
- 30. Siavash Mirarab, Alaa Hassouna, and Ladan Tahvildari. Using bayesian belief networks to predict change propagation in software systems. In Proceedings of the 15th International Conference on Program Comprehension (ICPC), pages 177– 188. IEEE, 2007.
- 31. Mehdi Amoui, Siavash Mirarab, Sepand Ansari, and Caro Lucas. A genetic algorithm approach to design evolution using design pattern transformation. Journal of Information Technology and Intelligent Computing, 1(2):235–244, 2006.

### Software (Publicly AVAILABLE)

ASTRAL (https://github.com/smirarab/ASTRAL)

Coalescent-based species tree estimation from gene trees

PASTA (https://github.com/smirarab/pasta)

Multiple sequence alignment of ultra-large datasets (up to a million sequences)

SEPP, TPP (https://github.com/smirarab/sepp)

Phylogenetic placement and taxonomic identification of metagenomic reads

UPP (https://github.com/smirarab/sepp/blob/master/README.UPP.md) Multiple sequences alignment using families of HMMs for ultra-large datasets

FastSP (https://github.com/smirarab/FastSP)

Fast comparison of very large multiple sequence alignments

See https://github.com/smirarab/?tab=repositories for more.

# EXPERIENCE

Professional International Business Machines (IBM), Vancouver, British Columbia, Canada

IT Specialist

September 2008 – December 2010

Worked on a software platform for wireless pace-maker devices

### Teaching EXPERIENCE

### Teaching Assistance:

University of Waterloo, Software Testing and Quality Assurance

Winter 2007

University of Tehran, Scientific & Technical Communication

Fall 2005

### Workshops, Tutorials, and Short Courses:

Ann Arbor, Michigan, Phylogenomics Symposium and Software School

2015

UT Austin, Summer school on Phylogenomics and Metagenomics **Summer 2014** Evolution meetings (NC), Software School on ASTRAL and PASTA Summer 2014 UT Austin, Workshop on New Methods for Phylogeny and Alignment Estimation,

Tutorials on SEPP and SATe

Winter 2013

HIGHLIGHTS Talks

RECOMB, Warsaw, Poland, "Statistical binning enables an accurate ..."

2015

IPAM, UCLA, "Ultra-large multiple sequence alignments". **2015** INFORMS, Philadelphia, "Reconstruction of species histories using genomic data" **2015** 

PROFESSIONAL Program Committee: ISMB/ECCB 2015
SERVICES Paper Review: TCBB, Bioinformatics, ISMB, RECOMB-CG, Theoretical Population

Genetics, TSE, ICPC 2007-2015