Ashok Rajaraman

+1.217.417.9746 | ashok.rajaraman@gmail.com | ashokr@andrew.cmu.edu | Homepage | GitHub

EDUCATION

Doctorate in Mathematics

April 2015 | Burnaby, BC Supervisor: Cedric Chauve

Thesis topic: Algorithms and complexity results for variants of the consecutive ones property.

IIT ROORKEE

B. Tech in Metallurgical and Materials Engineering

June 2009 | Roorkee, Uttarakhand, India

PROJECTS

DeClone: 2015 FPSAC: 2013 ANGES: 2012

WORKSHOPS

PIMS IGTC Mathematical Biology Summit

Novamber 2013 | Banff, Alberta

ENUMEX

September 2012 | Bertinoro, Italy

IPAM Evolutionary Genomics November 2011 | Los Angeles, CA

AWARDS

Michael Stevenson Scholarship: 2014 SFU President's PhD Scholarship: 2014 PIMS IGTC Fellowship: 2012-2014 SFU Travel Awards: 2011,2012 SFU Graduate Fellowship: 2010,2012

SKILLS

Programming:

(Most used first)

Python C++ shell perl

Mathematical software:

MATLAB R

Typesetting/markup:

LATEX - HTML

REFERENCES

On request

EXPERIENCE

SIMON FRASER UNIVERSITY CARNEGIE MELLON UNIVERSITY | Postdoctoral Research Associate, Computational Biology Department, School of Computer Science January 2016 - | Pittsburgh, PA

UNIVERSITY OF ILLINOIS AT URBANA-CHAMPAIGN

Postdoctoral Research Associate, Department of Bioengineering, College of Engineering

May 2015 - December 2015 | Urbana, IL

Host: Jian Ma. Research on mathematical models in computational genomics.

RESEARCH

COMPUTATIONAL MODELS IN COMPARATIVE GENOMICS

- Integrated ancestral genome reconstruction at multiple resolutions.
- Hypergraph models for genome mapping problems.
- Dynamic programming algorithms in phylogenetics.
- Approximations for generalized vertex ordering problems.

PUBLICATIONS

- Accept Rajaraman, A. and Ma, J. Reconstructing ancestral gene orders with duplications guided by synteny level genome reconstruction. In RECOMB-CG 2016.
 - 2016 Rajaraman, A., Zanetti, J.P.P., Maňuch, J., and Chauve, C. Algorithms and complexity results for genome mapping problems. IEEE/ACM Transactions on Computational Biology and Bioinformatics, PP(99):1-1, 2016. ISSN 1545-5963.
 - 2015 Neafsey, D.E., Waterhouse, R.M. et al. Highly evolvable malaria vectors: The genomes of 16 anopheles mosquitoes. Science, 347(6217), 2015.
 - Rajaraman, A., Chauve, C., and Ponty, Y. Assessing the robustness of parsimonious predictions for gene neighbourhoods from reconciled phylogenies. In Bioinformatics Research and Applications - 11th International Symposium, ISBRA 2015, Norfolk, VA, USA, June 7-10, 2015 Proceedings, pages 260-271. 2015.
 - 2013 Chauve, C., Patterson, M., and Rajaraman, A. Hypergraph covering problems motivated by genome assembly questions. In Combinatorial Algorithms - 24th International Workshop, IWOCA 2013, Rouen, France, July 10-12, 2013, Revised Selected Papers, pages 428-432. 2013.

Rajaraman, A., Tannier, E., and Chauve, C. FPSAC: Fast Phylogenetic Scaffolding of Ancient Contigs. Bioinformatics, 29(23):2987–2994, 2013.

Rajaraman, A., Tannier, E., and Chauve, C. The genome of the medieval Black Death agent. In JOBIM 2013 (Extended abstract).

2012 Jones, B.R., Rajaraman, A., Tannier, E., and Chauve, C. ANGES: Reconstructing ANcestral GEnomeS maps. Bioinformatics, 28(18):2388–2390, 2012.