Jonathan Belyeu

Computational Genetics



Education

August PhD Human Genetics, University of Utah, Salt Lake City, UT.

2016-Present

August BS Bioinformatics, Minor: Computer Science, Brigham Young University,

2009-April Provo, UT.

2016

Experience

Research/Academic

May Graduate Research Assistant, Aaron Quinlan, Ph.D., University of Utah, Salt 2017-Present Lake City, UT, Eccles Institute of Human Genetics.

Computational research in genetics/genomics

- Performing statistical analysis of de novo structural variation in a large WGS family
- Developing/distributing command-line tools for solving genomic problems
- Performing data visualization in Python/R
- Using supercomputing resources for analysis of genomic structural variants
- Developed web and command-line tool (SV-plaudit) for structural variant curation

September Research Assistant, Perry Ridge, Ph.D., Brigham Young University, Provo, UT, 2014–April Department of Biology.

2016 Computational research in genetics/genomics

- Built phylogenetic trees from species sets
- o Performed evolution-based analyses of synonymous codon bias
- Graphically analyzed data with R
- Used supercomputer resources for large-scale analyses
- Developed command-line tool (SelecT) for statistical identification of selection

April Research Assistant, Department of Biology, Richard Gill, Ph.D., Brigham

2015-April Young University, Provo, UT.

Ecological data pipeline development

- Built data pipeline for automated or manual storage of ecological time-series data
- Managed Ubuntu server and MySQL database of time-series data
- o Performed time-series data analyses and quality control

Scientific Communication

Selected Publications

Jonathan R Belyeu, Thomas A. Sasani, Brent S. Pedersen, Aaron R Quinlan. "Unfazed: parent-of-origin detection for large and small de novo variants" 2021. 10.1101/2021.02.03.429658

Jonathan R Belyeu, Harrison Brand, Harold Wang, Xuefang, Zhao, Brent S. Pedersen, Julie Feusier, Meenal Gupta, Thomas J Nicholas, Lisa Baird, Bernie Devlin, Stephan J Sanders, Lynne B Jorde, Michael E Talkowski, Aaron R Quinlan. "*De novo* structural mutation rates and gamete-of-origin biases revealed through genome sequencing of 2,396 families" 2020. 10.1101/2020.10.06.329011

Jonathan R Belyeu, Murad Chowdhury, Joseph Brown, Brent S. Pedersen, Michael J. Cormier, Aaron R. Quinlan. "Samplot: A Platform for Structural Variant Visual Validation and Automated Filtering" 2020. 10.1101/2020.09.23.310110

Jonathan Belyeu, Thomas J Nicholas, Brent S Pedersen, Thomas A Sasani, James M Havrilla, Stephanie N Kravitz, Megan E Conway, Brian K Lohman, Aaron R Quinlan, Ryan M Layer. "SV-plaudit: A cloud-based framework for manually curating thousands of structural variants." 2018. 10.1101/265058

Talks

Jonathan Belyeu. Analysis of spontaneous human genomic structural variation in 2300 WGS families. Invited seminar, Brigham Young University Department of Biology, February 2020.

Jonathan Belyeu, Aaron Quinlan. Analysis of de novo structural variation rates in a large cohort. University of Utah Department of Human Genetics Retreat, November 2019.

Jonathan Belyeu, Ryan Layer, Julie Feusier, Lynn Jorde, Aaron Quinlan. Direct measurement of de novo structural variation through whole-genome sequencing of three-generation human pedigrees. Genome Informatics, September 2018.

Jonathan Belyeu, Ryan Layer. SV-plaudit: Rapid Visual Review of Structural Variants. Virtual invited seminar, Genome in a Bottle Consortium, July 2018.

Darian Ramage, Artem Golotin, **Jonathan Belyeu**. Streaming Correlation-based Seismic Event Detector. Lawrence Livermore National Laboratory, April 2016.

Computational/programming skills

Bioinformatics Skills

- Unix tools
- Algorithm development
- Computational tool development
- Parallel processing
- Supercomputing
- AWS
- Conda
- Nextflow
- Matplotlib
- o ggplot2

Programming Languages

- Python
- Bash/Shell
- o R
- o C++ (some)
- Java (some)

Other

- o Git
- PTEX