Profile

I'm a scientist with training in computer science, statistics, and biology. I work with high dimensional data on clustering, classification and modeling problems for disease diagnostics. Computing to me is a hobby as well as a profession, and I constantly work on improving my skills through reading and contributing to open source projects. I seek and would thrive in an interdiciplinary, team oriented and technical environment. See github (https://github.com/joshuaar) or linkedin (https://linkedin.com/in/joshuaar) for more details.

Skills

Analytical Skills, Experience in supervised and unsupervised machine learning, genomics data analysis, high dimensional data, multivariate and nonparametric statistics, data visualization in R and Python.

Languages, Python, R, Scala, Java, C++, Javascript.

Software/Libraries, Linux, Apache Lucene, Vim, Git, Illustrator.

Experience

02.2014- USDA Agricultural Research Service, Bioinformatics Specialist, Maricopa, AZ. Present Run a small informatics core facility focusing primarily on RNA-Seq transcriptomics in insects.

- Built a web interface exposing tools for parallel processing of genomics data to research scientists.
- Designed an analysis pipeline available on Galaxy for de-novo transcriptome assembly, annotation, searching and visualization. Used by three participating research groups.
- Support and parallelization of FORTRAN code for radiative transfer atmospheric models.

2010-Present Biodesign Institute, Research Associate, Tempe, AZ.

Conduct research projects involving high density **peptide microarrays**. Design experiments, algorithms and software geared toward turning this technology into a universal disease diagnostic called Immunosignaturing (http://en.wikipedia.org/wiki/Immunosignature).

- Designed and tested a system for rapidly mapping B-Cell epitopes from human sera and monoclonal antibodies. Includes a novel statistical algorithm for epitope determination from noisy sequence data.
- Designed and built a rapid pattern search engine for biological sequences using Apache Lucene and Scala. It is over 100x faster than naive methods, and useful for identifying peptide motifs in large sequence databases. Includes trigram index, pattern compiler, web interface.
- Wrote non-parametric (bootstrapping) validation procedures for a an immunosignature based Dengue Fever diagnostic. Includes feature selection, SVM classification, cross validation, ROC analysis and visualization. I packaged this into a command line application which became standard protocol in the lab.
- Assorted infrastructure work SQL database design, data warehousing, organization and retrieval. PostgreSQL Database and web interface for storing, cleaning and retrieving high throughput peptide microarray data.

2009-2010 Entech, Quality Control Engineer, White Pigeon, MI.

Directed quality control for a recycling plant specializing in tire rubber for artificial turf, molding and extrusion.

- Built and supported a database for QC results with **nightly backups to offsite storage**.
- Wrote procedures creating formatted reports for management.
- Technologies used: PostgreSQL, Python, Excel
- Ensured data and product samples were available and accurate, which was important for occasional disputes over quality.
- 01.2009- University of Notre Dame, Research Visitor, Notre Dame, IN.
- 07.2009 Research internship at UND in the lab of Dr. Paul Helquist. Synthesized TSA analogs for treatment of Niemann-Pick disease, co-authoring two papers in peer reviewed journals.

Education

2010-Present Arizona State University, Ira. A. Fulton School of Engineering, Ph.D, Biological Design (conc. Bioinformatics), Tempe, AZ.

2005-2009 Goshen College, B.A, Chemistry, Mol. Biology, Goshen, IN.

Quantitative Coursework (Selected).

- Nonparametric Statistics
- Linear Regression
- Statistical Computing
- Data Structures and Algorithms
- Theoretical Computer Science
- Operating Systems
- Assembly Language Programming

Publications

Josh Richer, Stephen Johnston, and Phillip Stafford. Epitope identification from fixed-complexity random-sequence peptide microarrays. *Molecular and Cellular Proteomics*, Accepted with minor revisions.

Josh Richer, Xiao Wang, and Stephen Johnston. Immunosignatures for dengue diagnosis. Manuscript in Preparation.

Joe Hull, Josh Richer, and Jeffrey Fabrick. Sequencing and de novo assembly of the western tarnished plant bug (lygus hesperus) antennal transcriptome. Manuscript in Preparation.

Casey Cosner, Vijaya Bhaskara, Anamitra Chaterjee, John Markiewicz, Steven Corden, Joakim Lofstedt, Ankner Tobias, Joshua Richer, Tyler Hulett, Douglas Shauer, Olaf Wiest, and Paul Helquist. Evolution of concise and flexible synthetic strategies for trichostatic acid and the potent histone deacetylase inhibitor trichostatin a. *European Journal of Organic Chemistry*, 2013(1):162–172, 2013.

Anamitra Chatterjee, Joshua Richer, Tyler Hulett, Vijaya Bhaskara Reddy Iska, Olaf Wiest, and Paul Helquist. An efficient synthesis of (\pm) -trichostatic acid and analogues: A new route to (\pm) -trichostatin a. *Organic letters*, 12(4):832–834, 2010.

Contact

Phone **630-319-2073**.

Email joshamosricher@gmail.com.

Email (work) Josh.Richer@ars.usda.gov.