

Jianying Li
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SUMMARY

- **Four-year** college teaching experiences
- **Ten-year** bioinformatics working/research experiences with focus on genome informatics, microarray and next generation sequencing
- **Three-year** hands-on experiences in Next Generation Sequencing (NGS) analysis and development
- Strong in statistical modeling and statistical/computational programming in **SAS, JMP, R, Perl, Java, and C++**, **Linux/Unix shell scripting**
- Self-motivated individual with smooth communication skills

WORKING EXPERIENCE

Strayer University 10/2010 to present Adjunct Faculty

- Teaching math and science related courses

Kelly Government Solutions, Inc. 05/2012 to present Bioinformatics scientist

- Major duties
 - Providing biostatistics/bioinformatics consultation
 - Designing and leading internal training courses for bench scientists
 - Developing and conducting bioinformatics methodologies
 - Developing and conducting individual bioinformatics research

Duke University, Center for Human Genome Variation
04/2011 to 05/2012 Bioinformatician II

- Selected recent projects involved
 - Led and participated in upgrading in-house variant calling pipeline for **whole genome** and **exome** projects. In the project, we evaluated main third party software include BWA, Samtools, GATK, etc. and optimized the processes for the best performance. Eventually, deployed it for regular process.
 - Constructed an in-house MySQL database keeping track of the variant calling project progress, including three major components (GWAS experiment, Illumina sequencing and variant calling).
 - Developed a sequencing QC platform for real time comparison to traditional GWAS experiment, this involves direct process sequencing .bam file, curating Illumina “topstrand”

output from GWAS, consulting current dbSNP databases and performing two parts of concordance check.

UNC Chapel Hill, Lineberger Comprehensive Cancer Center

10/2008 to 03/2011 Senior Bioinformatician

■ Selected recent projects involved

- Led and participated in establishing in-house pipeline for RNA-seq project. I am a main developer for RNAseq QC modules and analytical workflow in SeqWare (http://sourceforge.net/apps/mediawiki/seqware/index.php?title=Main_Page). This project involves benchmarking and testing open source software package, implementing in-house statistical analysis, constructing customized database for data annotation, building LIMS and data storage database for *ad hoc* analyses. All statistical components are implemented in **R**, sequence processing and manipulation is implanted in **Perl**, and the final software/application is wrapped with **Java**
- Lead author on the Principal Variance Component Analysis (**PVCA**) procedure leveraging mixed linear model for genetic/genomic data analysis. The theory was published as a book chapter and the model was developed both in **SAS** (currently an analytical procedure “AP” in JMP Genomics software package from SAS Institute) and **R** (<http://www.niehs.nih.gov/research/resources/software/pvca/index.cfm>) for open source implementation.
- Participated in recently funded (by NIH) **The Cancer Genome Atlas (TCGA)** project, mainly relying on next generation sequence platform (plus genotyping and other high throughput method) to detect new mutation in cancer cell, find new transcripts, and study novel cancer mechanistic system. My main responsibility is statistical analysis and model on high-throughput NGS data.
- Collaborated and led the bioinformatics component on a project “to determine the DNA replication system for cancer mechanism using next generation whole genome sequencing”

Lockheed Martin (NIEHS contractor), 01/2003 to 10/2008 Bioinformatic Scientist

■ Bioinformatics application and statistical methodology development

- Developed in house scientific software in **Java**, for microarray quality control of in house RNA quality monitoring, hybridization quality assessment, data acquisition software performance evaluation
- Developed in house scientific software in **C++** implementing Fuzzy Logic algorithm for Rat acetaminophen dosing project
- Developed in house software in **R** providing analytical component in an in-house developed Oracle database Chemical Effect Biological System (CEBS)
- Implemented Mixed Linear Model in **SAS** for customized Micro Array data analysis, include data quality evaluation, data normalization, and statistical modeling

■ Statistical consultation and bioinformatics support

- Provided training of application statistics using **JMP**: acted as a leading role to coordinate with IT training department to design class, construct class materials and present in a live training class
- Performed regular microarray gene expression and proteomic data analysis with commercial software package, freeware as well as in house developed software in **Java, R, SAS** and **JMP**

■ Selected research projects involved

- Developed statistic prediction models in the FDA MAQC consortium for standardized procedure for prediction phenotypic outcome using the transcriptome biomarkers, and currently involving in cross-platform and cross-tissue prediction research
- Participated in scientific director's CpG project, implementing genome searching algorithms to search for domains in human and mouse genome
- Participated in the collaborated research for statistic modeling in Toxicogenomics Research Consortium cross platform and cross laboratory gene expression

University of Hawaii/North Carolina State University, 1998-2002 Research Assistant

- Research on genes (in *Listeria monocytogenes*) responsible for surface antigen determinants involving all molecular technique, i.e. DNA/RNA manipulation, microarray experiment, and function study
- Research involving search for major genes contributing to favorable commercial properties in loblolly pine

EDUCATION

- Master of Bioinformatics, North Carolina State University, 2002
- Master of Science in Food Microbiology, North Carolina State University, 2001
- B.S. in Microbiology, Nankai University Tianjin, 1990

HARDWARE AND SOFTWARE SKILLS

- Operating systems: **Windows, Linux, and Unix**
- Programming languages: **Java, C++, SQL, Perl, SAS, R, JMP**
- Database system: MySQL, PostgreSQL
- Genetic/genomic analysis package: JMP Genomics Suite, Rosetta Resolver, GeneSpring, Bioconductor package, IPA, GoMiner
- NextGenSeq data analysis package: BWA, Samtools, Picard, GATK, SeqWare, Galaxy, FASTX-toolkit etc.

SELECTED RECENT PUBLICATIONS

Myrtle Davisa, Elaine Knighta, **Jianying Li**, Kellye K. Danielsc, James E. Doroshowa and Pierre R. Bushel Toxicogenomic Profiling of Bone Marrow from Rats treated with Topotecan in Combination with Oxaliplatin: A Proof of Concept Study to Identify a Strategy to Assess Risk of

Enhanced Toxicity for Combination Therapies (2013) Submitting to Molecular Cancer Therapeutics

Wang, C, ..**Li J**, ...Tong, W. RNA-Seq and Microarray Gene Expression Vie for Superiority of Biologic Elucidation Using A Comprehensive Study Design (2013) In prep.

Laura R. Barkleya, Komaraiah Palleb, Michael Durandob, Tovah A. Dayc, Aditi Gurkard, Naoko Kakushoe, **Jianying Li**, Hisao Masaie, and Cyrus Vazirib c-Jun N-terminal kinase-mediated Rad18 phosphorylation facilitates Polη recruitment to stalled replication forks **Mol. Biol. Cell** (2012) vol. 23 no. 10, 1943 - 1954

Chen H, **Li J**, Li H, Hu Y, Tevebaugh W, et al. (2012) Transcript Profiling Identifies Dynamic Gene Expression Patterns and an Important Role for Nrf2/Keap1 Pathway in the Developing Mouse Esophagus. PLoS ONE 7(5): e36504. doi:10.1371/journal.pone.0036504

J Huang, W Shi, J Zhang, J W Chou, R S Paules, K Gerrish, **J Li**, et al on the MAQC consortium, (2010) Genomic indicators in the blood predict a drug-induced liver injury response **The Pharmacogenomics Journal** (2010) 10, 267–277

Cabanski CR, Qi Y, Yin X, Bair E, Hayward MC, **Li J**, et al. (2010) SWISS MADE: Standardized WithIn class Sum of Squares to evaluate Methodologies And Dataset Elements **PloS ONE** 5(3): e9905

Jianying Li, Chu-TZ, Wolfinger, R, Bushel, P.R. (2009) Batch Effects and Noise in Microarray Experiments: Sources and Solutions. John Wiley & Sons, Ltd.

Song, M.O., **J. Li**, and J.H. Freedman (2009) Physiological and Toxicological Transcriptome Exposure Levels Changes in HepG2 Cells Exposed to Copper” **Physiol. Genomics**. 38: 386-401.

P. R. Bushel, A. N. Heinloth, **J. Li**, et al (2007) “Blood Gene Expression Signatures Predict Exposure Levels” **PNAS**. v.104, no. 46: 18211-18216