Delasa Aghamirzaie

Contact information

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Education

- Ph.D. in Genetics, Bioinformatics, and Computational Biology (GBCB), May 2016, Virginia Polytechnic Institute and State University (Virginia Tech), Blacksburg, Virginia, USA.
 - o Major: Computer Science, Minor: Life Sciences
 - Dissertation: isoform-specific expression during Arabidopsis and soybean embryo development
- M.Sc. in Computer Engineering, Computer Architecture, October 2011, Department of Computer Engineering and Information Technology, Amirkabir University of Technology (Tehran Polytechnic), Tehran, Iran.
 - Thesis: Accelerating Protein Secondary Structure Prediction Using Artificial Neural Networks on GPU
- B.Sc. Computer Engineering, September 2009, Hardware, Department of Computer Engineering and Information Technology, Amirkabir University of Technology (Tehran Polytechnic), Tehran, Iran.
 - Thesis: Suppression of Variation Effects on FPGAs Using Multiple Configurations

Research Interests

- Large-scale data-driven problems in Computational Biology and Bioinformatics
- Developing Computational Tools and Statistical Models for Genomics and Transcriptomics Applications using *Machine Learning* and *Deep Learning*
- Integrative Analysis of Next Generation Sequencing Data
- Single Cell Genomics
- Big Data Analysis
- Functional Genomics
- Gene Regulatory Network Inference
- Long noncoding RNAs

Research Experience

- Post Doctoral Fellow, Trapnell lab, Department of Genome Sciences, University of Washington,
 July 2016- present
- Graduate Research Assistant, Group of Dr. Ruth Grene, Co-advised by Dr. Lenwood Heath (Department of Computer Science) Virginia Tech (2012- June 2016)
 - Analysis of RNA-Seq Data of developing soybean embryos, mutant and wild type Arabidopsis seeds, and seedlings
 - Developed methods for analysis of transcriptomics data
 - CoSpliceNet for co-splicing network analysis available on http://delasa.github.io/co-spliceNet/
 - Co-Expression Network Analysis
 - Bayesian Network Inference of gene expression data incorporating prior knowledge
 - Analysis of Alternative Splicing Events During Embryo Development
 - Functional Analysis of Long Noncoding RNA
 - Development of CodeWise: A Support Vector Machine (SVM) Classifier for assigning Transcripts Into Coding And Noncoding Classes based both on sequence and on structural features
 - "Expresso": Exploring Transcription Factors And Their Inferred Target Genes From Arabidopsis ChIP-Seq Data

- Developing a Webserver And DB For Integrating And Browsing Available ChIP-Seq Data For Arabidopsis
- Research Intern in Computational Biology Group, IBM T.J Watson Research Center, Yorktown Heights, NY, May-August 2015
 - o Watson Genomics Analytics (WGA) Project
 - Molecular profile analysis of cancer patients
 - o Application of Deep Learning techniques to gene expression data sets
- Remote Research Scholar in Persepolis Group, Under Supervision of Dr. Amir H. Assadi,
 Department of Mathematics, University of Wisconsin-Madison (2010-2011)
 - Gene Expression Data Analysis of Arabidopsis on GPU
 - Clustering Gene Expression Data
 - o Super Structure Artificial Neural Networks on GPU
- Research Assistant in FPGA Laboratory, Department of Computer Engineering, Amirkabir University of Technology (9/2009-10/2011)
 - GPU Programming Using Nvidia CUDA Programming Model
 - o Protein Secondary Structure Prediction
 - Multi class Pattern Classification
- Research Assistant in Design Automation laboratory, Department of Computer Engineering, Amirkabir University of Technology (2008-2009)
 - Variation Effects on FPGAs
 - Effects of Multiple Configurations in Variation Suppression

Awards and Honors

- 2016: Translational Plant Sciences Travel Award
- 2016: Cold Spring Harbor Scholarship Award for Plant Genomics Meeting
- 2015-2016: Genetics, Bioinformatics, and Computational Biology Academic Fellowship
- 2015: Graduate Student Assembly Travel Award
- 2015: Molecular Plant Science Student Grant Proposal Winner
- 2014: Molecular Plant Science Travel Award Fellowship for 2nd Plant Genomics Conference
- 2013: NIH Fellowship for National Short Course in System Biology, University of California, Irvine
- 2011: Best Master Thesis Award in The Amirkabir University of Technology
- 2011: Ranked 3rd among M.Sc. Students in Computer Engineering
- 2009: Accepted as an Honored/Talented Student to Computer Engineering M.Sc program
- 2005-2009: Ranked 2nd among B.Sc. Students in Computer Hardware Engineering.
- 2007: Cisco Certified Network Associates (CCNA) Degree

Selected Course Projects at Virginia Tech

- Deep Learning and Perceptron, Fall 2015
- Computational Biochemistry in Bioinformatics (Spring 2014)
- Bayesian Statistics, Spring 2013
- Introduction to Machine Learning and Perception (Spring 2013)

Modeling and Identifying Regulatory Modules in Soybean Time Series Gene Expression Data using Bayesian Networks

Technical skills

Next Generation Sequencing:

RNA-seq analysis: Tuxedo Suite Protocol

ChIP-Seq analysis

DEXseq, Limma R package (Linear Models for Microarray Data)

Gene Set Enrichment Analysis_

Ontologizer for Gene Ontology Enrichment Analysis

Machine Learning:

Naïve Bayes Classifiers, Logistic Regression, Support Vector Machines, Artificial Neural Networks, Bayesian Networks, Time-Course Clustering, SplineCluster, k-means

· Deep Learning:

Restriced Boltzman Machines, Stacked Autoencoderes, recurrent neural networks

Tools: PyLearn2, Torch

Programming languages

Python, C++: Proficient

MATLAB, R, Assembly, Pascal, Visual Basic.Net, Java: Intermediate

• Database Programming, and Web Development

MySQL, SQL Server, HTML, PHP: Intermediate

Asp.net: Familiar

Statistics:

Classical: Hypothesis testing, Classical Regression analysis, ANOVA, One Sample and Two

sample t-tests

Bayesian: Bayesian Regression analysis, MCMC

GPU programming

CUDA: Familiar

Version Control:

Github

Invited Talks

- Delasa Aghamirzaie, "Fundamentals of Computational Algorithms for Data Analysis of RNA-Seq Data", Paradigms of Bioinformatics. October 2015
- **Delasa Aghamirzaie,** Erhan Bilal, Takahiko Koyama, Fang Wang, Filippo Utro, Kahn Rhrissorrakrai, Raquel Norel, Laxmi Parida, and Ajay Royyuru, "Interpreting Cancer Patient Genomics Data to Assist Clinicians", IBM T.J Watson Research Center, Intern Seminar Series, August 2015
- Delasa Aghamirzaie, "Fundamentals of Next Generation Sequencing", Plant Stress Physiology Course, April 2015.
- Delasa Aghamirzaie, "CodeWise: A Support Vector Machine Classifiers for Accurate detection of Noncoding RNAs", Graduate Student Assembly Symposium, March 2015
- Delasa Aghamirzaie, "Comprehensive Functional Characterization Reveals Novel Relationships Among Differentially Expressed Transcripts in Developing Soybean Embryos", MPS/TPS Mini-symposium, Feb, 2015
- Delasa Aghamirzaie, "An Accurate Support Vector Machine Classifier For Assessing Coding Potential
 Of Transcripts Using Several Sequential And Structural Features", Biological Data Science Meeting,
 Cold Spring Harbor Laboratories, New York, November 2014
- Ruth Grene, **Delasa Aghamirzaie**, "Toward a Functional Classification of Splice Variants From Soybean Seed Embryos", 2nd Plant Genomics Congress, St. Louis, USA, September 2014

Publications

- **D. Aghamirzaie**, S. Li, E. Collakova, R. Grene, "CoSpliceNet: A framework for co-splicing network inference from transcriptomics data", submitted.
- Y. Ni, **D. Aghamirzaie**, E. Collakova, S. Li, R. Grene, L.S. Heath, "A Machine Learning Approach to Predict Gene Regulatory Networks in Seed Development in Arabidopsis", submitted.
- A. Schneider, D. Aghamirzaie, H. Elmarakeby, A. Poudel, A.J. Koo, L.S. Heath, R. Grene, E. Collakova, "Potential Targets Of VAL1 Repression In Developing Arabidopsis Thaliana Embryos", The Plant Journal, January 2016.
- **D. Aghamirzaie**, D. Batra, L.S.Heath, A. Schneider, R. Grene, E. Collakova, "Transcriptome-Wide Functional Characterization Reveals Novel Relationships Among Differentially Expressed Transcripts In Developing Soybean Embryos", BMC Genomics, November 2015.
- **D. Aghamirzaie,** M. Nabiyouni, Y. Fang, C. Klumas, L.S.Heath, R. Grene, E. Collakova, "Changes in RNA Splicing in Developing Soybean (Glycine max) Embryos", Biology 2013.
- E. Collakova, D. Aghamirzaie, Y. Fang, C. Klumas, F. Tabataba, A. Kakumanu, E. Myers, L. Heath, R. Grene, "Metabolic and Transcriptional Reprogramming in Developing Soybean (Glycine max) Embryos", Metabolites, May 2013.
- M. Nabiyouni, D. Aghamirzaie, "A Highly Parallel Multi-Class Pattern Classification on GPU," 12th IEEE/ACM International Symposium on Cluster, Cloud and Grid Computing (ccgrid), 2012, Best Paper Candidate.
- **D. Aghamirzaie**, A. Razavi, M. Saheb Zamani, M. Nabiyouni, "Reduction of Process Variation Effect on FPGAs Using Multiple Configurations", IEEE/IFIP International Conference on Very Large-Scale Integration (VLSI-SOC'10), September 2010.
- Manuscripts in preparation
 - **D. Aghamirzaie**, Sh. Wu, K. Raja Vulmugran, D. Altarawy, L.S. Heath, R. Grene, "Expresso: Exploring Arabidopsis' Transcription Factors and Target Genes from ChIP-Seq Data", Under Preparation.

Poster Presentations at international conferences

- **D. Aghamirzaie**, Sh. Wu, K. Raja Vulmugran, R. Grene", Expresso: Exploring Arabidopsis' Transcription Factors and Target Genes from ChIP-Seq Data", Biological Data Science, Cold Spring Harbor Laboratory, USA, November 2014.
- **D. Aghamirzaie**, Sh. Wu, K. Raja Vulmugran, R. Grene", Expresso: Exploring Arabidopsis' Transcription Factors and Target Genes from ChIP-Seq Data", 2nd Plant Genomics Congress, St. Louis, USA, Septermber 2014.
- **D. Aghamirzaie**, L. Heath, R. Grene, E. Collakova, "Changes in RNA Splicing in Soybean Seed Embryos", 8th q-bio Conference, Santa Fe, NM, August 2014.
- D. Aghamirzaie, D. Batra, E. Collakova, L. Heath, R. Grene, "Modeling and Identifying Regulatory Modules in (Glycine max) Soybean Time Series Gene Expression Data using Bayesian Networks". International Conference on Computational Cell Biology (ICCCB 2013), Blacksburg, Virginia, August 2013.

References:

- Dr. Ruth Grene (Professor of Plant Physiology), grene@vt.edu, 540-231-6761
- Dr. Eva Collakova (Associate Professor of Plant Physiology), collakov@vt.edu, 540-231-7147
- Dr. Lenwood Heath (Professor of Computer Science), heath@vt.edu, 540-231-4352
- Dr. Song Li (Assistant Professor of Plant Genomics and Bioinformatics), songli@vt.edu, 540-231-2756
- Dr. Erhan Bilal, Research Staff Member, IBM T.J Watson Research Center, ebilal@us.ibm.com