# Curriculum Vitae – Mr. Arash Shahsavari

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### **Education**

2016 - 2019 M.S. in Complex Adaptive Systems - Chalmers University of Technology

Gothenburg, Thesis: An Evaluation of Multi-Step Analyses of Single-Cell RNA Sequencing Data.

Sweden Advisor: Rebecka Jörnsten.

Selected Coursework: Neural Networks, Stochastic Optimization, Information Theory, Dynamical Systems.

Fall 2017 Exchange Studies - Dongguk University

Seoul, Selected Coursework: Multiple View Geometry, Deep Learning.

South Korea

2013 - 2016 Electrical Engineering - Lund University

Lund, Completed 3 years of a 5-year joint B.S./M.S. program in electrical engineering, equivalent to a B.S. degree. Sweden Selected Coursework: Multivariable Calculus, Control Theory, Mathematical Statistics, Numerical Analysis.

### **Experience**

### 2019- Bioinformatician at Cambridge Stem Cell Institute, University of Cambridge

Cambridge, UK

- Long-term data science support for multiple biomedical projects (50+ datasets for 10+ different projects).
- Machine learning and statistics on terabyte-scale datasets
- Involved in entire recruitment pipeline, from designing recruitment test, to selecting candidates for interviews, interviewing and final recruitment decisions
- Co-authoring peer-reviewed publications

### 2018-2019 Volunteer at Kodcentrum

Gothenburg, Sweden

- Teaching coding to children from underprivileged areas
- · Planning and structuring classes, hands-on support to children and other volunteers
- Long-term development of the learning platforms and curriculum

# **Selected Research Projects**

#### 2020-2021 ClustAssess: Tools for scRNA-seq Clustering Assessment - Cambridge

Cambridge, UK

- Development of R package for data-driven assessment of clustering
- Profiled code to identify computational bottlenecks and reimplemented performance-critical parts in
- High-resolution quantification of clustering stability
- Investigation of how cell types inferred from data depend on clustering results

# 2020-2021

# Mapping the Biogenesis of Forward Programmed Megakaryocytes - Cambridge

Cambridge, UK

- Characterization of temporal changes in time-course scRNA-seq data
   Integration with in vivo dataset to quantify accuracy of in vitre system
- Integration with in-vivo dataset to quantify accuracy of in-vitro system
- Identification of novel surface markers across multiple flow cytometry datasets

### 2020-2021

### $Consequences\ of\ Splitting\ 10X\ scRNA-seq\ Samples\ Across\ Sequencing\ Lanes\ -\ Cambridge$

Cambridge, UK

- Simulation study assessing the downstream consequences of sample splitting, compared to an unsplit sample, throughout the scRNA-seq analysis pipeline
- · Quantification of impact on clustering and cluster marker genes

#### Spring 2019 Master's Thesis - Dept. of Mathematical Sciences, Chalmers

Gothenburg, Sweden

- Benchmarking of scRNA-seq clustering methods across feature selections and datasets
- Assessing deconvolution methods' robustness to noise in cell type signatures
- Successful defense of thesis leading to M.S. degree

Fall 2018

### HIV Transcription Latency - Dept. of Microtechnology and Nanoscience, Chalmers

Gothenburg, Sweden

- Investigation of feedback and noise in mathematical model of HIV transcription
- Stochastic simulations of biochemical reactions with Gillespie algorithm
- Quantification of uncertainty of molecular concentrations over time

#### Spring 2016 Feedback Controller for Ballbot - Dept. of Automatic Control, Lund University

Lund, Sweden

- · Development of linear state feedback controller for ballbot
- Evaluating feedback controller and optimizing parameters in simulations with sensor noise
- Deployment on Arduino microcontroller including interfacing with Inertial Measurement Unit

# Mentoring

2021

Andi Munteanu, UAIC Computer Science M.S. student and Cambridge Stem Cell Institute bioinformatics intern

### **Skills and Development**

- **Programming Environments:** Python (Optimization example),
  R (Clustering evaluation package, Random Forest feature selection example),
  Julia (Outlier detection example, Optimization example), MATLAB, C++, Bash, 上下X, Arduino.
- Computing Tools: SLURM, git, Jupyter Notebook (Example), R Markdown (Example)
- Software packages: ClustAssess developed from scratch, currently package maintainer Github, Example, Documentation
- **Professional Courses:** Python for Bioimage Analysis week-long course by the Royal Microscopical Society.
- **Self-Studied Books and Courses:** Intermediate Linear Algebra, Computational Linear Algebra, Real Analysis, Combinatorics & Graph Theory, Reinforcement Learning.
- Webpage: sharash.github.io

### **Journal Publications**

- [1] **Arash Shahsavari** and I. Mohorianu. "ClustAssess: tools for assessing the robustness of single cell clustering". In preparation (2021).
- [2] J. Chen\*, V. Sathiaseelan\*, A. Moore, S. Tan, C. S. R. Chilamakuri, **Arash Shahsavari**, C. Jakwerth, S. B. Hake, A. Warren, I. Mohorianu, C. D'Santos, and I. Ringshausen. "ZAP-70 constitutively regulates gene expression and protein synthesis in chronic lymphocytic leukemia". *Blood* (2021), Commentary.
- [3] M. Lawrence\*, **Arash Shahsavari**\*, S. Bornelöv\*, T. Moreau, K. Kania, M. Paramor, R. McDonald, J. Baye, M. Perrin, M. Steindel, P. Jimenez, C. Penfold, I. Mohorianu, and C. Ghevaert. "Mapping the biogenesis of forward programmed megakaryocytes from induced pluripotent stem cells". Under review at Science Advances (2021).
- [4] C. M. Morell, S. G. Tilson, R. A. Tomaz, **Arash Shahsavari**, G. Canu, B. T. Wesley, M. Perrin, I. Geti, S. Mukhopadhyay, F. Mazzacuva, P. Gissen, J. Garcia-Bernardo, C. A. Rimland, F. Sampaziotis, I. Mohorianu, and L. Vallier. "Novel 3D Approach to Model Non-Alcoholic Fatty Liver Disease using human Pluripotent Stem Cells". In preparation (2021).
- [5] E. Williams\*, R. Chazarra-Gil\*, **Arash Shahsavari**\*, and I. Mohorianu. "The sum of two halves may be different from the whole. Effects of splitting samples across sequencing lanes". Submitted to Cell Reports (2021).

<sup>\*</sup> indicates joint first authorship