Adrin Jalali

Curriculum Vitae

Berlin, Germany ⊠ adrin.jalali@gmail.com '` github/adrinjalali

Experience

2018-present scikit-learn, Core Developer.

2020-present fairlearn, Core Developer.

2020-present **Machine Learning Engineer**, *HuggingFace*, Remote.

- Maintain scikit-learn and fairlearn
- Work on HuggingFace Hub integration of third part libraries and lead scikit-learn integration.

2020-2021 **Senior Applied Scientist**, *Zalando SE*, Berlin, Germany.

- Center of Excellence for Algorithmic Privacy and Fairness
 - Assess and deploy model cards across the organization
 - Help teams understand and mitigate ethical related risks
 - Audit and review other teams' work to assess potential risks and harms in their models and processes

2019-2020 Open Source Developer, Anaconda Inc., Berlin, Germany.

- o scikit-learn
- o fairlearn: a fairness in ML lib originally by Microsoft

2016-2019 **Senior Machine Learning Consultant**, Ancud IT, Berlin, Germany.

- Machine Learning (scikit-learn, tensorflow, GPy, etc.)
- NLU/NLP (intent and entity recognition, conversation context, etc.)
- NoSQL (Cassandra, Elasticsearch, Solr)
- MLOps (PipelinelO: kubernetes, weave-scope, AWS, MS-Azure, docker, etc.)

2012-2016 PhD Candidate, Max Planck Institute for Informatics, Saarbrücken, Germany.

- Interpretable Sparse methods for cancer associations using an ensemble of sparse support vector machines (SVMs)
- 2011-2012 PhD Candidate, British Columbia Cancer Research Center, Vancouver, Canada.
 - Automated flow cytometry data analysis and visualization using dynamic programming, clustering, graph algorithms, and SVMs
- 2009-2011 **Team Lead**, *Tosan Intelligent Data Miners*, Tehran, Iran.
 - o Offline fraud detection and real-time fraud prevention on debit transactions

2006-2007 Al and Statistics Programmer, Fidofa Software Group, Tehran, Iran.

Automated trading in Future stock market

Education

2012-2021 **PhD (cont.) in Computer Science/Bioinformatics**, Max Planck Institute for Informatics, Computational Biology and Applied Algorithmics Department, Saarbrücken, Germany,

Dissertation Topic: Interpretable Methods in Cancer Diagnosis .

2011-2012 **PhD in Bioinformatics**, *University of British Columbia, British Columbia Cancer Research Center, Terry Fox Laboratory*, Vancouver, Canada, *Topic:* Automated analysis of flow-cytometry data.

2006-2009 **MSc in Computer Science**, *University of Tehran, School of Mathematics, Statistics, and Computer Science*, Tehran, Iran, *Thesis Topic:* Finding DNA Motifs Using Bidirectional Recurrent Neural Networks.

2002-2006 **BSc in Computer Science**, *Tehran Polytechnic*, *Department of Mathematics*, and *Computer Science*, Tehran, Iran.

Service

2017-present PyData Berlin coorganizer, NumFocus, Berlin, Germany.

- 2015 Chemistry, Physics, and Technology Section Representative, *PhDnet steering group*, Max Planck Society, Germany.
- 2012 **Bioinformatics Representative**, *College for Interdisciplinary Studies Graduate Student Society*, University of British Columbia, Vancouver, Canada.
- 2012 "Bioinformatics" and "British Columbia Cancer Agency Graduate Student and Post Doctoral Fellow Society" representative, *Graduate Student Society*, University of British Columbia, Vancouver, Canada.

Languages

Persian Native

English Fluent

German Fluent

Publications

- 1. Handl L, Jalali A, Scherer M, Pfeifer N., Partially blind domain adaptation for age prediction from DNA methylation data, arXiv preprint arXiv:1612.06650, 2016.
- 2. Jalali A., and Pfeifer N., *Interpretable per Case Weighted Ensemble Method for Cancer Associations*, BMC Genomics, volume 17, no. 1, 2016.
- 3. Courtot M., Meskas J., Diehl A. D., Droumeva R., Gottardo R., Jalali A., Taghiyar M.J., *flowCL:* ontology-based cell population labelling in flow cytometry, Bioinformatics 31, no. 8 (2015): 1337-1339.
- 4. Jalali A., and Pfeifer N., *Interpretable per Case Weighted Ensemble Method for Cancer Associations*, Algorithms in Bioinformatics, pp. 352-353. Springer Berlin Heidelberg, 2014.
- 5. O'Neill K.*, Jalali A.*, Aghaeepour N.*, Hoos H.H., and Brinkman R.R., Enhanced flow-Type/RchyOptimyx: A Bioconductor pipeline for discovery in high-dimensional cytometry data, Bioinformatics (2014), doi: 10.1093/bioinformatics/btt770.
- 6. Jalali A.*, Aghaeepour N.*, O'Neill K., Chattopadhyay P.K., Roederer M., Hoos H.H., Brinkman

- R.R., *RchyOptimyx: cellular hierarchy optimization for flow cytometry*, Cytometry Part A 81, no. 12 (2012): 1022-1030.
- 7. Aghaeepour N., Chattopadhyay P.K., Ganesan A., O'Neill K., Zare H., Jalali A., Hoos H.H., Roederer M., and Brinkman R.R., *Early Immunologic Correlates of HIV Protection can be Identified from Computational Analysis of Complex Multivariate T-cell Flow Cytometry Assays*, Bioinformatics, 2012: 28(7):1009–1016.

^{*} co-authors contributed equally