# Narges Rezaie

# Bioinformatics, Data Analytics, Computational Biologist

■ (949)426-8112 nargesr@uci.edu Personal website narges-rezaie nargesr 

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# Summary

Innovative Postdoctoral Scholar specializing in bioinformatics and computational biology with a Ph.D. in Mathematical, Computational, and Systems Biology from UC Irvine. Over 6 years of experience developing bioinformatics tools and automated pipelines for biomedical data analysis. Proficient in multiple programming languages and machine learning.

# Research Experience

Postdoctoral fellow Mortazavi Lab, University of California, Irvine

CA, USA March 2024 - Present

• Developing/applying novel bioinformatic tools to analyze short and long read RNA-seq at bulk and single-cell resolution

**Graduate student** Mortazavi Lab, University of California, Irvine

**CA, USA** Dec 2019 – March 2024

- Developing and applying novel bioinformatic tools to analyse bulk and single-cell RNA-seq datasets.
- Model-AD: We are part of an NIA Consortium to build better late-onset Alzheimer's Disease models in mice. I am mainly involved in analyzing/validating these new models as well as implementing standardized data acquisition and data analysis protocols for these mouse models of Alzheimer's disease
- **IGVF**: We are leveraging mouse genetic diversity to study the impact of genotype on molecular function in various tissues using single-cell resolution.
- ENCODE4: We are part of the ENCODE consortium's effort to profile full-length transcriptomes in humans and mice.
- **Gene Regulatory Networks**: We are building models of gene regulatory networks using either bulk or single-cell RNA-seq, and ATAC-seq in humans, rodents, and other vertebrates to discover how the logic of development is encoded in the genome.

**Lab Rotation** Swarup Lab, University of California, Irvine

**CA. USA** Sep 2019 – Dec 2019

- Comparing various methods of doublet detection
- Analyzing single nucleus RNA sequencing (snRNA-seq) data to detect doublet.

**Research Assistant** Sharif University of Technology

**Tehran, Iran** *May* 2016 – Aug 2019

• Statistical Analysis of Clinical Data - Golestan Cohort: Designing a statistical model to predict risk of cardiovascular disease

#### Education

PhD in Mathematical, Computational, and Systems Biology University of California, Irvine

USA 2019-2024

- **Thesis**: Computational approach to characterize gene dynamics using bulk and single-nucleus RNA sequencing to study Alzheimer's disease. **Supervisor**: Dr. Ali Mortazavi
- **GPA**: 3.98/4.0

B.S. in Information Technology in Computer Engineering Sharif University of Technology

Iran 2014-2018

# Skills

- **Programming Languages:** Python (Matplotlib, seaborn, NumPy, pandas, SciPy, Anndata, scikit-learn, TensorFlow, and PyTorch), R, shinyapps, jupyter notebook, Bash, Matlab, C/C++, Java, SQL, HTML, CSS, Django, and PHP
- Softwares: Blast, Lima, Minimap2, Cellbender, Samtools, bedtools, Picard, RSEM, Seurat, Scanpy, and Kallisto
- Developed by me: PyWGCNA and Topyfic
- Pipeline managements: NextFlow and Snakemake
- Assays: Single-cell/nucleus RNA-seq, Multi-omics, Pertub-seq, Bulk long-read RNA-seq, Bulk short-read RNA-seq, and Hi-C
- Frameworks: Git, Django, Docker, and Flask

# **Publications**

- LF Garcia-Agudo, ..., **N Rezaie**, ..., V Swarup, GR MacGregor, KN Green.  $BIN1^{K358R}$  suppresses glial response to plaques in mouse model of Alzheimer's disease. Alzheimer's & Dementia (2024)
- N Rezaie, ..., B Wold, A Mortazavi. Identification of robust cellular programs using reproducible LDA that impact sex-specific disease progression in different genotypes of a mouse model of AD. bioRxiv (2024)
- F Reese, ..., N Rezaie, ..., BJ Wold, A Mortazavi. The ENCODE4 long-read RNA-seq collection reveals distinct classes of transcript structure diversity. bioRxiv (2023)
- KM Tran, ..., N Rezaie, ..., GR MacGregor, KN Green. A Trem2R47H mouse model without cryptic splicing drives age-and disease-dependent tissue damage and synaptic loss in response to plaques. Molecular neurodegeneration (2023)
- N Rezaie, F Reese, A Mortazavi. PyWGCNA: A Python package for weighted gene co-expression network analysis. Bioinformatics (2023)
- M Bayati, N Rezaie and M Hamidi, MS Tahaei, H Rabiee. A New R Package for Categorizing Coding and Non-Coding Genes. Preprints (2023)
- H Alinejad-Rokny, ..., N Rezaie, KT Tam, ARR Forrest. MaxHiC: A robust background correction model to identify biologically relevant chromatin interactions in Hi-C and capture Hi-C experiments. PLOS Computational Biology (2022)
- N Rezaie, ..., H Alinejad-Rokny. Somatic point mutations are enriched in non-coding RNAs with possible regulatory function in breast cancer. Communications Biology (2022)
- DI Javonillo, ..., **N Rezaie**, ..., KN Green, FM LaFerla. Systematic phenotyping and characterization of the 3xTg-AD mouse model of Alzheimer's Disease. Frontiers in neuroscience (2022)
- S Forner, ..., **N Rezaie**, ..., FM LaFerla, KN Green. Systematic phenotyping and characterization of the 5xFAD mouse model of Alzheimer's disease. Scientific data (2021)
- G Balderrama-Gutierrez, H Liang, **N Rezaie**, ..., F LaFerla, A Mortazavi. Single-cell and nucleus RNA-seq in a mouse model of AD reveal activation of distinct glial subpopulations in the presence of plaques and tangles. bioRxiv (2021)

## Selected Talks and Posters

- Identifying Distinct Cellular Programs from Single Cell Datasets Using Topyfic
   N Rezaie (2023 Alzheimer's and Brain Awareness Month, PARSE biosciences)
- Identifying regulatory topics in the IGVF mouse bridge samples using Topyfic.
   N Rezaie, ..., BJ Wold, A Mortazavi (IGVF Consortium Meeting 2023)
- Topyfic identified robust cellular programs using reproducible Latent Dirichlet allocation (rLDA).
   N Rezaie, E Rebboah, A Mortazavi (Network Biology 2023 (CHSL)) and (Probabilistic Modeling in Genomics 2023 (CHSL))
- Bulk and single-cell gene expression network analysis in mouse models of AD using PyWGCNA.
   N Rezaie, MODEL-AD, A Mortazavi (Alzheimer's & Dementia 2022 (AAIC))
- Phenotypic characterization of UCI MODEL-AD mice using RNA-seq.
   M Gantuz, HY Liang, N Rezaie, MODEL-AD, A Mortazavi (Alzheimer's & Dementia 2022 (AAIC))
- The MODEL-AD Explorer: An open-access resource for comparing phenotypic data from mouse models of Alzheimer's disease.
  - A Greenwood, ..., N Rezaie, ..., G Carter. (Alzheimer's & Dementia 2021 (AAIC))
- Analyzing gene expression during neurodegeneration in a 3xTgAD mouse model at transcriptome level.
   N Rezaie, ..., A Mortazavi. (Neuroscience 2021 (SFN))

## References