# JACOB KROL CURRICULUM VITAE

Computational Biologist/Professional RA

GitHub: jakekrolin LinkedIn: jacob-krol

P Denver, Colorado & Saline, Michigan; United States

## **Education**

Non-degree seeking - University of Colorado Anschutz Medical Campus- Aurora, CO

Fall 2023

• BIOS 7747: Machine Learning for Biomedical Applications, graduate course offered by the University of Colorado School of Public Health

B.Sc. in Computational Neuroscience - Michigan State University - East Lansing, MI

2020-2022

- · GPA: 3.89/4.0
- · Graduated 'With Honor'
- · Semester awards: Dean's List

Math and Science Transfer Program - Washtenaw Community College - Ann Arbor,

2018-2020

MI

- · GPA: 3.52/4.0
- · Semester awards: Honor Roll
- Transferred

## **Professional Experience**

**Information Sciences Professional (PRA)** - Dept. of Biomedical Informatics, Center for Health Artificial Intelligence, University of Colorado Anschutz School of Medicine - JRaviLab - Aurora, CO

2022-2023

- Redesigned the front and backend of MolEvolvR, a web app for protein characterization (http://jravilab.org/molevolvr; Krol, et al., 2023 \_bioRxiv\_; DOI:https://doi.org/10.1101/2022.02.18.461833)
- Developed a machine learning (ML) pipeline for classifying antimicrobial resistance (AMR) in bacterial strains (publication in-prep for early 2024 submission)
- Listed in 'Acknowledgements' section of publication \*provisionally accepted\* at \*mSystems DOI:https://doi.org/10.1101/2020.09.24.301986
- Presented research talks and gave programming workshops at 2 international conferences (Bioconductor 2023 and GLBIO 2023)
- Mentored 3 PhD students and 1 undergraduate to assist in omics data featurization, model development, and model outcome analysis
- Developing two R packages for AMR project (source code for Bioconductor and data for ExperimentHub)
- Used University of Colorado Anschutz School of Medicine (CU Anschutz)
   Alpine high performance computing cluster (HPC) to aggregate, transform, and train ML models on over 100GB of omics data
- Containerization of AMR data collection code and the R packages I developed, and worked with Faisal Alquaddoomi on containerizing the frontend, back-end, and slurm instance for the MolEvolvR web-application
- Resolved over 50 GitHub issues for multiple lab repositories and signficantly cleaned up consolidated projects with multiple repositories
- Mentored by members of CU Anschutz Department of Biomedical Informatics (DBMI) software engineering team
- Assisted in hiring process (interviewing and feedback) for various lab positions: post-doctoral and research assistants
- Implemented various ML approaches and evaluation techniques: logistic regression, gradient boosting machines, random forests, linear discriminant analysis, stratified-cross-validation, class-weighting, hyperparamter searching, and evaluating auROC, balanced accuracy, etc., on hold-out dataset
- Hosted department wide workshops on Bash, Git, R package development, and ssh-workflow basics
- Assisted in writing and generating figures for lab grant proposals and publications
- Performed system administrator duties (e.g., server onboarding, dependency/user/data/resource management) for our lab's webserver

## **Publications**

MolEvolvR: A web-app for characterizing proteins using molecular evolution and phylogeny. **Jacob D Krol**\*, Joseph T Burke\*, Samuel Z Chen\*, Lo M Sosinski\*, Faisal S Alquaddoomi, Evan P Brenner, Ethan P Wolfe, Vincent P Rubinetti, Shaddai Amolitos, Kellen M Reason, John B Johnston, Janani Ravi bioRxiv 2022.02.18.461833; doi: https://doi.org/10.1101/2022.02.18.461833 (\* co-primary author)

Accurately predicting AR in ESKAPE pathogens using ML. **Jacob D Krol**, Ethan P Wolfe, Evan P Brenner, Keenan Manpearl, Vignesh Sridhar, Joe Burke, Jill Bilodeoux, Janani Ravi (*In preparation*.).

#### **Presentations**

 $\begin{tabular}{ll} \textbf{Great Lakes Bioinformatics Conference} & - & MolEvolvR & a web-app & for protein characterization- McGill University, Montreal, CA \\ \end{tabular}$ 

2023

• Discussed the development and future directions of a web-app I develop: http://jravilab.org/molevolvr

**Great Lakes Bioinformatics Conference** - How and when to build a web-app or R package?- McGill University. Montreal, CA

2023

· Hosted a 4 hour in-person workshop on how to build an R package using

**R/Bioconductor** - MolevolvR a web-app for protein characterization - Boston University, Boston, MA

 Presented on MolEvolvR application methodology for the Cancer and Evolution talks section

## **Posters**

**CU DBMI Retreat** - Robust machine learning-based classification of antimicrobial resistance in high-impact pathogens- CU Anschutz, Aurora, CO

2023

2023

Presented on highly performant AMR classification ML models I developed

## **Funding**

- Submitted Fall 2023 NSF GRFP (in-review)
- Contributed to minor revisions NIH NIAID U01 grant submission (awarded; PI: Janani Ravi)
- Contributed revision and figures to NIH NIAID R01 grant submission (in-review; PI: Janani Ravi)
  Submitting Winter 2023 Department of Energy Computational Science Graduate Fellowship (inpreparation)

**Student Research Assistant II -** Computational Mathematics Science and Engineering program, Michigan State University - Krishnan Lab & Malmstrom Lab - East Lansing, MI

- Learned, presented on, and implemented statistical methods on viral protein datasets: fisher test, logistic regression, & principal component analysis
- Trained machine learning classifiers to predict plant virus' host types; also, trained models to predict plant virus taxonomy.
- Featurization of protein sequences and data wrangling with Pandas, Biopython, NumPy, and R (Tidyverse + Bioconductor) packages for biological feature extraction
- Further preparing data with one-hot-encoding and z-score normalization
- Analyzing and visualizing model performance with Matplotlib/Seaborn & Scikit-learn performance metrics

## **Professional Summary**

- Develop bioinformatics tools specializing in applying ML to large omics datasets using R, Python, and Bash/shell
- Use large, public databases (e.g., BV-BRC, NCBI, InterPro) with applied machine learning to study the relationship between genotypes and phenotypes
- Mentored 3 PhD students and 1 undergraduate student
- Developed a protein analysis web-app MolEvolvR
- Presented research talks and workshops at international conferences, hosted department-wide workshops, and presented research posters
- Exceptional IT knowledge: experienced with high performance computing (HPC), version control (GitHub), dependency management, containerization, package development, web-application hosting, using web APIs for data, etc.
- Basic familiarity of other common languages like Javascript, C, Java, and Perl
- Apply statistical methods to analyze omics data: supervised classification, fisher's test, under/oversampling, cross-validation, rank-based hypothesis testing, etc
- Assist in various sub-tasks throughout the lab such as one-on-one mentorship with undergraduates and graduate students

## **Peer-Mentees**

#### **PhD Students**

- Keenan Manpearl; 2023 (Computational Biosciences program; CU Anschutz)
- Jill Bilodeaux; 2023 (Computational Biosciences program; CU Anschutz)
- · Charmie Vang; 2023 (Biophysical Sciences program; CU Anschutz)

#### **Undergraduates**

• Ethan Wolfe; 2023 (Biochemistry & Molecular Biology (BS) with CMSE and additional minors)

### **Project summaries**

## Machine learning-based classification of antimicrobial resistance:

- · Project lead, first author
- Developed a computational pipeline to gather bacterial genomes and AMR data from public databases, featurize the genomes, develop ML models to classify AMR, and analyze top predictors to discover novel AMR genes
- Mentored 3 PhD students and 1 undergraduate to assist in omics data featurization, model development, and model outcome analysis
- Supervised learning with large (over 10k bacterial isolates) sample sizes and tackling class imbalance with weighted loss functions and undersampling
- · Model evaluation using confusion matrix performance metrics such as balanced accuracy, auROC, etc.
- Implemented Fisher's Exact test for nearly a million genes to determine significant presence/absence in the binary classes (resistant/susceptible phenotypes) yielding ranked gene lists for AMR contribution
- Developed the omics data featurization, ML, and non-ML pipelines as Bioconductor R package with planned submission in early 2024

- All source code for data wrangling, machine learning, and presentation is already installable as an R package on https://github.com/jravilab/amR (private until submission)
- · Planned submission for datasets and results to Bioconductor's ExperimentHub in early 2024
- Future directions:
  - Build a sequence database of top resistance genes by clustering across species to address the uninterprebility of gene cluster assignment across species using species-wise pangenomics tools
  - Design a web-application to showcase results of ML models in classifying AMR for various drugs/species
  - Design a web-application which allows submission of bacterial genomes for AMR classification

## References

Janani Ravi - University of Colorado Anschutz School of Medicine- Aurora, CO

2022-Present

- · janani.ravi@cuanschutz.edu
- Assistant Professor, Principal investigator of JRaviLab

Faisal Alquaddoomi - University of Colorado Anschutz Anschutz School of Medicine-

2022-Present

Aurora, CO

- · faisal.alquaddoomi@cuanschutz.edu
- IT Principal Professional

Arjun Krishnan - Michigan State University- East Lansing, MI

2022

- · arjun.krishnan@cuanschutz.edu
- Associate Professor, Principal invesigator of Krishnan Lab