Alexander Larsen

PHONE (920) 246-8971

E-MAIL Alarsen525@gmail.com

GITHUB github.com/Zinthrow

LINKEDIN www.linkedin.com/in/a-w-larsen

SKILLS

Managerial & Communication Skills

- Managerial experience of a Bioinformatics team: employee reviews, project tracking.
- Clinician consultations, patient customer service, public speaking
- SOP & publication writing, creation of publication visuals.

Software Engineering & Development

- Principles: Code review, git versioning, project management tools (Trello, Jira).
- Pipeline development & module creation: Python, Perl, C++, AWS, SQL, command-line/bash.
- Web Development: HTML, CSS, Javascript.
- Workflow management: Docker, Nextflow.

Bioinformatics & Biophysical Modeling

- NGS protocols: optimization, data analysis, library preparation (Illumina NextSeq).
- Tools & software: Bowtie2, BWA, BLAST command line, GATK, Samtools, Annovar, Picard, usearch,
 CD-HIT, FastQC, ClinVar, Kraken2, CLARK, MetaPhlan2, Pathoscope2, NCBI Utilities, R programming.
- Algorithms: Vienna, Nupack, MSA, Hidden Markov Models, Smith-Waterman/Needlman-Wunsch, UPGMA, Nearest Neighbor agglomerative, DeBruijn.

Statistical & Data Analysis

- Machine Learning: Kmeans++, Linear Regression / Logistic regression (L1/L2), Classification, Random forests, Boosting, Bagging, Ensembling, SVM, KNN, A*.
- Statistical methods: Bayesian analysis, Regression, PCoA/PCA, alpha/beta diversity analysis, DeSEQ2.
- Python Libraries: numpy, pandas, matplotlib, seaborn, biopython, primer3-py, sklearn, scipy, sqlalchemy.

Laboratory Techniques & Protocols

 DNA/RNA extraction, Western Blotting, FISH, SDS-Page Electrophoresis, Mass Spectrometry, Neuronal electrophysiology, PCR.

High-Performance Computing (HPC)

• Defining environments, nodes, **linux**, operating systems, user groups, job scheduling (SLURM), and resource management.

EXPERIENCE

Bioinformatics Scientist I

Sherlock Biosciences Inc.

Nov 2020 - Current Boston, MA

Projects

- Primary developer on company bioinformatics pipelines optimizing RNA structure, CRISPR predictive performance, LAMP constraints, SDA ROI identification across two different chemistries.
- Achieved 99%+ reduction in genomic feature space for all company amplification technologies across Sense SDA, Cap SDA, LAMR and **CRISPR** spaces reducing clinical diagnostic failure post-deployment.
- Identified novel thermostable CRISPR-Cas proteins for publication, patent, and LAMP + Cas diagnostic.
- Created software for high specificity,/inclusivity/LOD and thermodynamically stable guide targets tool.

Clinical Bioinformatician I, II

June 2018 - Nov 2020

Sterling, VA

Aperiomics Inc.
Projects

- Designed automated COVID-19 report end to end software. Analysis, formatting, clinical sign-off,
- Developed metagenomic analysis library that automatically queries patient data for report statistics formatting. Accomplished using python, bash, sql, css/html, git, Django, Docker, AWS CLI.
- Presented on bioinformatics clinical value to investors, posters, and at other local events.

Associate Research Assistant

Apr 2013 - May 2016 University of Wisconsin Madison Madison, WI

 Characterizing circadian rhythm genes and their temporal effect on long term memory formation in drosophila. Particular effort towards the quality control of data and samples. - Professor Jerry Yin / Robin Fropf

• Characterizing zebrafish embryonic development, crayfish nerve reading frequencies, and rat hippocampal electrical activity. - Professor Tony Stretton

EDUCATION / CERTIFICATIONS

Bioinformatics M.S. Mar 2019 - Nov 2021

University of Maryland, Adelphi

Adelphi, MD

Alignment, Statistical Analysis, SQL Databases and Protein Structure. Biotechnology and Biomanufacturing.

Bioinformatics June 2017 - Apr 2018

Audited – UW Madison / MIT Open Courseware

De Pere. WI

- Graduate studies following UW Madison outlines for a master's in bioinformatics.
- Bioinformatic algorithms I/II, general algorithms / data structures, artificial intelligence.

Neurobiology B.S.

Sept 2012 - May 2016

University of Wisconsin Madison

Madison, WI

Graduated from a selective honors biology program which facilitated group directed research, scientific writing, and analysis. – UW Madison Biocore program.

edX Courseware Certificates

May 2018 - Current

University of Maryland / UC Berkley

- BIF001x: DNA Sequences
- BIF002x: Proteins: Alignment, Analysis and Structure
- BIF003x: Statistical Analysis in Bioinformatics
- Bioinformatics MicroMasters Certificate
- Foundations of Data Science: Computational Thinking with Python

PUBLICATIONS

Published

- Baugher, D., Larsen, A., Chen, Y., & others. (2021). Female Urinary Microbiome Analysis and Artificial Intelligence Enhances the Infectious Diagnostic Yield in Precision Medicine. Microbiol Infect Dis, 5(4), 1–8.
- Larsen, A., Icenhour, C., Crandall, K., Valencia, C. A., & Chen, Y. (2020). Increased Virulent Microorganism Associations in the Urinary Microbiome of Interstitial Cystitis Patients. George Washington University.
- Larsen, A. W., Chen, Y., Crandall, K. A., Icenhour, C. R., & Valencia, A. C. (2022). Characterization of the Interstitial Cystitis/Bladder Pain Syndrome Microbiome in Clinically Diagnosed Patients. Clinical Immunology & Research, 6(1), 1-9.
- Pena JM, Manning BJ, Li X, Fiore ES, Carlson L, Shytle K, Nguyen PP, Azmi I, Larsen A, Wilson MK, Singh S, DeMeo MC, Ramesh P, Boisvert H, Blake WJ. Real-Time, Multiplexed SHERLOCK for in Vitro Diagnostics. J Mol Diagn. 2023 Jul;25(7):428-437. doi: 10.1016/j.jmoldx.2023.03.009.
- Phillips, E; Silverman, A; Liu, M; Brown, C; Carlson, P; Coticchia, C; Shytle, K; Larsen, A; Goyal, N; Cai, V; Huang, J; Hickey, J; Ryan, E; Acheampong, J; Ramesh, P; Collins, J; Blake, W. Detection of viral RNAs at ambient temperature via reporter proteins produced through the target-splinted ligation of DNA probes. Nat. Biomed. Eng (2023). https://doi.org/10.1038/s41551-023-01028-v.

Various, US