Ryan P Smith

 $\begin{array}{l} linkedin.com/in/rpseq\\ github.com/rpseq \end{array}$

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Profile

Bioinformatics scientist turned data infrastructure engineer. Strong background with 7 years of bioinformatics computing work in academia and 5 years of professional data engineering at Amazon and Recursion.

TECHNICAL SKILLS

- Languages: Python, Bash, R (ggplot), awk, SQL, C++, golang, LATEX, Mathematica, Java, Scala
- Technologies: Linux/Unix, Kubernetes, Slurm, HPC, Helm, AWS (S3, EC2, Redshift, EKS, EMR), GCP(GKE), Grafana, Prometheus, PostgreSQL, Terraform, Looker, git, Docker, Airflow, CircleCI, GitLab CI, Databricks, Spark, HBase, Pinot, Druid, Avro/Parquet, Kafka, GATK, bedtools, bwa-mem

EXPERIENCE

Recursion

Salt Lake City, UT

 $Senior\ Software\ Engineer$

September 2020 - Present

- **Inception Labs**: Work with senior leadership to develop novel product ideas and take them through proof-of-concept. Rapidly design and build a large-scale Perturbseq sample processing pipeline using our on-prem HPC cluster.
- Transcriptomics Data Processing: Workflow orchestration. Design and build out automated RNAseq data processing systems in GCP. This includes a metadata tracking service and cloud computing pipelines to process millions of cell samples from an automated laboratory process.

eero, an Amazon company

San Francisco, CA

Software Development Engineer, Data

January 2019 - September 2020

- Data Infrastructure and Operations: Design, provision, maintain, and scale k8s clusters to power our data systems. Provisioned with Terraform and automated deployments. Spark for custom batch and streaming transformations. Metrics, monitoring and alerting using Prometheus and Grafana. Manage a diverse collection of k8s data services.
- Real-time Data Systems: Build and maintain data systems powering customer-facing features and internal tools. Visual dashboards for observing device bandwidth usage as well as security and content filtering events across a fleet of more than 6 million routers. Kafka for ingesting up to 60k records per second to a large HBase cluster, while simultaneously serving 60k customer queries per second.
- Data Analytics Platform: Design and provision AWS infrastructure for a data platform built around Pinot in k8s. Powers sub-second queries over very large datasets with continuous realtime ingestion of fleet data from Kafka, using Superset for data visualization and exploration.

Girihlet, Inc., Oakland Genomics Center

Oakland, CA

 $Bion formaticist + IT\ Engineer$

Aug 2018 - Oct 2018

- **Bioinformatics**: Provide a quality control dashboard for Illumina short read data. General Bash, Python, and Perl scripting to automate mitochondrial genome variant calling.
- IT: Upgrade and maintain the Oakland Genomics Center networks. Maintain two on-premises CentOS servers for data processing and web hosting (Apache, NGINX)

The McDonnell Genome Institute, Washington University

St Louis, MO

Graduate Research Scientist, Ira Hall Lab, Computational Genetics

July 2014 - May 2018

- o **Distributed Computing, Bioinformatics**: Process and analyze population-level whole-genome sequencing data to improve our understanding of structural variation in human genomes. Computation is done in a Linux environment with Bash pipelines, distributed across a high-performance computing cluster. Pipelines often use tools combining Bayesian statistics and machine learning approaches, packaged and deployed via Docker containers.
- Single-cell Sequencing: Develop novel computational and molecular biology methods for whole-genome sequencing of single mammalian neurons. Process up to 10 TB of raw Illumina sequencing data in 36 hours.
- **Teaching Assistant**: Advise students sequencing the genomes of unknown bacteriophages. Lab course using bioinformatics tools to identify genes, predict their function, and classify into phylogenetic groups.

University of Iowa

Iowa City, IA

Undergraduate Research Fellow, Adam Dupuy Lab, Cancer Genetics

Aug 2010 - May 2014

• Genome Editing, Bioinformatics: Design a viral gene delivery vector for simulating human cancers in mice alongside a DNA sequencing method to detect resulting transgene insertions. Process whole-genome sequencing data using Linux bioinformatics tools followed by ad-hoc statistical analyses visualization in Python and R.

EDUCATION

Washington University

MA, Molecular Genetics and Genomics

University of Iowa BS, Microbiology and Informatics

St Louis, MO Aug. 2014 - May 2018 Iowa City, IA Aug. 2009 - May 2014