

RAZVAN IOAN PANEA

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Summary

- Initiative-taking with strong knowledge and experience in computer science and bioinformatics.
- Led the design, innovation and development of production-level genomics compute infrastructure that yearly processes and analyses petabytes-scale multi-omics data
- Developed from scratch a cloud-based workflow management system for bioinformatics in Python.
- Led a group of software engineers to develop a complete cloud-based infrastructure using Kubernetes and Django.
- Processed a terabytes-scale genomics dataset of Burkitt lymphoma and performed analysis using R.

Education

- 08/2014 – 05/2020 **Duke University | Durham, NC, USA**
Ph.D. in *Computational Biology and Bioinformatics*
Research: *Cloud computing and cancer genomics*
- 09/2011 – 06/2014 **Jacobs University Bremen | Bremen, Germany**
Bachelor of Science in *Applied and Computational Mathematics*
Specialization: *Bioinformatics and Computational Biology*

Research and Work Experience

- 08/2020 – Present **Regeneron Pharmaceuticals | Tarrytown, NY, USA**
Senior Genomic Data Engineer – Manager: William Salerno
► Led the design, innovation and development of the production genomics compute infrastructure
Technologies used: *Python, Bash, Git, AWS, DNANexus, Databricks, Jira*
- Optimized and innovated genomics workflows that process and analyze over 500,000 samples per year
 - Led the software engineering efforts by applying software development best practices, specifically version control using Git, software development life cycle tracking using Jira, and continuous integration, delivery and testing of methods through Amazon Web Services CodePipeline
 - Worked with bioinformaticians and data scientists to develop tools on data platforms such as DNANexus to facilitate at-scale, at-speed genomics pipelines for QC, variant calling and multi-omics analysis
 - Worked closely with other data engineers to integrate production with distributed compute environments
 - Interacted with software engineers and external technology partners to ensure 24/7 production uptime
- 08/2014 – 06/2020 **Duke University | Durham, NC, USA**
Ph.D. Candidate – Advisor: Sandeep Dave, MD
► Developed a cloud-based infrastructure for cancer genomics
Technologies used: *Python, Bash, Git, Docker, Kubernetes, Google Cloud, AWS, Apache libcloud*
- Developed CloudConductor, a cloud-based, modular, scalable, parallelizable and extensible workflow engine to generate and execute analysis pipelines. Available at github.com/labdave/CloudConductor
 - Led a team of 5 software engineers and bioinformaticians to extend CloudConductor to a complete cloud-based infrastructure for bioinformatics analysis using Kubernetes and Django
 - Improved CloudConductor to a platform-agnostic workflow engine using Apache libcloud
 - Integrated and evaluated the workflow engine on Google Cloud Platform and Amazon Web Services
 - Integrated 50+ bioinformatics tools using Docker and utilized them to analyze terabytes of genomics data
- Characterized the mutational patterns in Burkitt lymphoma subtypes
Technologies used: *R, Python, Bash*
- Sequenced the whole genome and transcriptome of 101 Burkitt Lymphoma cases of different subtypes
 - Identified genetic alterations in the sequencing data using the newly developed cloud-based infrastructure
 - Identified mutations in driver genes, such as *MYC*, *ID3* and *DDX3X*, the presence of the Epstein-Barr virus (EBV) in the samples, and translocations characteristic to Burkitt Lymphoma
 - Identified associations between lymphoma subtypes and EBV status
 - Work has been published in the *Blood* journal (*Panea et al., Blood 2019*)

06/2013 – **Max Planck Institute for Marine Microbiology | Bremen, Germany**
08/2013 *Software Developer* - Supervisor: Dr. Frank-Oliver Glöckner
► Improved features of the SILVA Project (www.arb-silva.de) software using Python and C++ binding
Technologies used: C++, Python

- Researched and tested SWIG, a software for Python and C++ binding
- Implemented SWIG in the project to bind and call methods implemented in Python on C++ objects

► Tested the performance and efficiency of a phylogenetic tree generating software
Technologies used: C++

- Benchmarked a new phylogenetic tree generator software
- Compared the performance and quality of the new tree generator software to the one currently used
- Deployed the new, more efficient software

03/2013 – **Jacobs University Bremen | Bremen, Germany**
06/2014 *Research Assistant* – Advisor: Dr. Georgi Muskhelishvili
► Determined the interaction between two neighboring genes in *E.coli*
Technologies used: Java

- Developed a software in Java to aggregate, analyze and report the expression levels of all neighboring genes at different time points in *E.coli* life cycle
- Identified a correlation between the differential expression between two neighboring genes and their transcriptional orientation to each other

Publications

- 09/2019 **Panea R***, Love C*, Shingleton J*, Reddy A* et al., "The whole genome landscape of Burkitt lymphoma subtypes", Blood, 2019
- 02/2020 Shingleton J., [and 16 authors, including **Panea R.**] (2020). "Non-Hodgkin Lymphomas: Malignancies Arising from Mature B Cells". In M. Kharas, R. L. Levine, A. M. Melnick (Ed.), "Leukemia and Lymphoma: Molecular & Therapeutic Insights". Cold Spring Harbor, NY: Cold Spring Harbor Laboratory Press, (In Press).
- 11/2019 Li X., [and 75 authors, including **Panea R.**], "Whole Exome and Transcriptome Sequencing in 1042 Cases Reveals Distinct Clinically Relevant Genetic Subgroups of Follicular Lymphoma" Blood (2019), (Manuscript in preparation).

Posters and Presentations

- 07/2018 **Google Next '18 | San Francisco, CA, USA**
Presentation and Panel Discussion
Title: Speeding up Research in Genomics
Link: <https://youtu.be/goBFt3B976A?t=11m31s>
- 06/2016 **Intelligent Systems for Molecular Biology | Orlando, FL, USA**
Poster Presentation
Title: Defining the Microbiome of Lymphomas
- 10/2017 **Duke Cancer Institute | Durham, NC, USA**
Poster Presentation
Title: A Cloud-Based Framework for Cancer Genomics

Skills

Programming: Python, R, Bash, C, Java, C++, SQL, Git
Libraries/Packages: Apache Libcloud, Plotly, Pandas, SQLAlchemy
Computing Platforms: Google Cloud, SLURM, Kubernetes, AWS
Operating Systems: Linux, Windows, OS X, Chrome OS
Softwares: PyCharm, Jupyter Notebook, RStudio, Adobe Illustrator

