Computational biology researcher and leader. After pivoting my career from software engineering leadership to reunite with my passion for biological research in 2017, I came to the Pulmonary division at Northwestern. Here, I set up computational biology operations from sample tracking to interpretable multimodal analyses to support translational research in lung diseases, building the foundation for the division NIH funding of \$5M+ and impactful publications. My interest is to use vast amount of published and private multimodal data via novel machine learning models to improve our understanding of diseases and factors that determine patient individual outcomes.

Expertise

Research leadership

Cross-functional collaboration, project management, team management, mentorship, scientific communication

Machine learning

Deep learning frameworks (torch, jax), large language models, multimodal fusion, embedding representations, bayesian modeling

Healthcare and biological data

Electronic health records (EHR), clinical notes analysis, single-cell RNA-seq, TCR-seq, spatial transcriptomics, imaging data, statistics

Software & tools

Python, R, SQL, Linux, computational clusters, interactive visualizations, pipelines

Education

Ph.D. in Computational Biology, Northwestern University, Chicago, USA Driskill Graduate Program in Life Sciences

2022-2025 (expected in December)

M.S. in Bioinformatics, Newcastle University, Newcastle upon Tyne, UK With distinction

2017-2018

Undergraduate coursework in Biology, Genetics, Moscow State University, Moscow, Russia

2003-2006

Work Experience

Ph.D. researcher, Division of Pulmonary and Critical Care Medicine Feinberg School of Medicine, Northwestern University, Chicago, USA 2022-present

- Led 4 large scientific collaborative projects with 500+ patients and multimodal data to publication
- Acquired external funding for my training (competitive AHA predoctoral fellowship, \$67,000)
- Authored and co-authored 18, including 4 first/co-first, publications or preprints
- Led analysis of scRNAseq patient samples in the context of their clinical course with novel machine learning approach (clustering of patient-day representations and late fusion)
- Identified cell population as a biomarker and potential therapeutic target of ILD in SSc (bioRxiv 2025)
- Consulted 7 Northwestern grad students, postdocs and faculty on deep learning, data science and data visualization, including setting up and training in paw tracking on videos for mouse experiments

Research data analyst, bioinformatics, Division of Pulmonary and Critical Care Medicine,

2019-2022

Feinberg School of Medicine, Northwestern University, Chicago, USA

- Created data processing pipelines, data exploration and management infrastructure for the division
- Delivered analytical insights from scRNAseq and other data to principal investigators for 7 publications
- Formulated activated T cell-macrophage circuit in Nature 2021, which supported successful clinical trials of Auxora in COVID-19 (NCT04345614)
- Supported grant writing for U19, R01 and other NIH grants for the division, resulting in \$5M+ funding
- Hire'd and trained incoming data analysts to grow the team and replace myself

Head of maintenance tools development group, Yandex, Moscow, Russia

2014-2017

- Managed a team of 6 engineers: hiring, mentoring, resolving conflicts, improving performance
- Synthesized internal customers' needs into technical roadmaps for supporting web-services
- Owned various web-services to improve employees' workflows

2007-2014

Full-stack software engineer, Yandex, Moscow, Russia
Automated deploy workflows of system administrators for better consistency and transparency Deployed and maintained various web-services to improve employees' workflows

Key publications

(* denotes first/co-first contribution)

* Profibrotic monocyte-derived alveolar macrophages as a biomarker and therapeutic target in systemic 2025 sclerosis-associated interstitial lung disease. **bioRxiv**. 10.1101/2025.08.07.669006 © 2025

Defining and benchmarking open problems in single-cell analysis. *Nature Biotechnilogy*. 10.1038/s41587-025-02694-w \circlearrowleft

2024

* A distinctive evolution of alveolar T cell responses is associated with clinical outcomes in unvaccinated patients with SARS-CoV-2 pneumonia. *Nature Immunology*. 10.1038/s41590-024-01914-w ©

2023

* Machine learning links unresolving secondary pneumonia to mortality in patients with severe pneumonia, including COVID-19. The Journal of Clinical Investigation (JCI). 10.1172/JCI170682 ♂ An integrated cell atlas of the lung in health and disease. **Nature Medicine**.

2023

UCSC Cell Browser: visualize your single-cell data. *Bioinformatics*. 10.1093/bioinformatics/btab503 &

2021

Circuits between infected macrophages and T cells in SARS-CoV-2 pneumonia. *Nature*. 10.1038/s41586-020-03148-w &

2021