I excel in applying machine learning tools (newly published or custom built) to high-resolution and multimodal data, such as single-cell RNA-seq, spatial transcriptomics or EHR data, to deliver high-quality analytics, interpretability and interactive visualizations. My interest is to use vast amount of published and private multimodal data via novel machine learning models to improve our understanding of common diseases and factors that determine patient individual outcomes.

Education

Ph.D. in Computational Biology, Northwestern University, Chicago, USA

2022–2025 (anticipated in June)
Driskill Graduate Program in Life Sciences

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

Analyzed scPNAsses and EHP data is inthe to understand national resource programming.

 Analyzed scRNAseq and EHR data jointly to understand patient response to severe pneumonia, including COVID-19

Acquired external funding for the project (AHA predoctoral fellowship)

Consulted Northwestern students and faculty on deep learning, data science and data visualization

Presented and published our research in 3 peer-reviewed papers

Post-baccalaureate research fellow, bioinformatics, Division of Pulmonary and Critical Care Medicine, 2019–2022

Feinberg School of Medicine, Northwestern University, Chicago, USA
Analyzed scRNAseq and other high-throughput data from human samples and mouse experiments

Cleaned and analyzed clinical EHR data from patient cohorts

Delivered analytical insights to principal investigators in presentations and data exploration tools

Developed and maintained data processing pipelines, data exchange and management infrastructure

Head of maintenance tools development group, Yandex, Moscow, Russia 2014–2017

Managed a team of 6 engineers: hiring, mentoring, resolving conflicts, improving performance

• Developed, designed and supported tools for system administrators and other employees

Full-stack software engineer, Yandex, Moscow, Russia 2007–2014

Identified employees' workflows bottlenecks for automation

 Developed, designed and supported web-services and console tools for system administrators and other employees

Software engineer, Art. Lebedev Studio, Moscow, Russia

Developed and supported web-sites and a content management system

Computational skills

Programming languages:

Python, R, Java, C++, Ruby, Perl. Linux. Latex. SQL. HTML, JavaScript. Github: https://github.com/mxposed ♂ Techniques:

Machine learning, deep learning (torch, jax), statistics, bayesian modeling, scRNAseq (scanpy, Seurat), bioinformatics tools, interactive visualizations (d3.js, plotly), pipelines (nextflow, snakemake).

Key publications

(* denotes equal contribution)

Markov NS, [...], Budinger GRS, Singer BD, Morales-Nebreda L, NU SCRIPT Study Investigators. 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 ♂

Gao CA*, **Markov NS***, Stoeger T*, [...], Misharin AV, Singer BD, NU SCRIPT Study Investigators.

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 &

Sikkema L, Ramírez-Suástegui C, Strobl DC, Gillett TE, Zappia L, Madissoon E, **Markov NS**, [...], Theis FJ. 2023 An integrated cell atlas of the lung in health and disease. *Nature Medicine*. 10.1038/s41591-023-02327-2 &

Speir ML, Bhaduri A, **Markov NS**, [...], Pollen AA, Raney BJ, Seninge L, Kent WJ, Haeussler M. 2021 UCSC Cell Browser: visualize your single-cell data. *Bioinformatics*. 10.1093/bioinformatics/btab503 ©

Grant RA*, Morales-Nebreda L*, **Markov NS***, [...], Budinger GRS, Misharin AV, Singer BD, Wunderink RG.
Circuits between infected macrophages and T cells in SARS-CoV-2 pneumonia. *Nature*.
10.1038/s41586-020-03148-w ©