

Ioannis Mouratidis

Austin, Texas

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Computational biology researcher with 32 publications (10 first or senior author) in large-scale data analysis and ML applications in genomics. Co-founded startup in cancer diagnostics, authored grants securing \$4M+ in competitive funding and currently lead a 5-member team focused on evals and safety of AI models in biology.

Experience

University of Texas at Austin

Austin, TX

Senior Research Scientist Associate (Research Engineer/Scientist Associate V)

08/2025 – Present

- Directed 10 research initiatives (4 independently, 6 collaboratively) resulting in 2 peer-reviewed publications with an additional 2 under review and 6 in preparation.
- Design research projects and lead a 5-member team with a focus in developing novel computational methods and testing the capabilities and safety profiles of foundation models in biology.
- Spearheaded novel compression tool for multiple genomic file formats, achieving 10-20% reduced file sizes and 50-70% faster compression times compared to state-of-the-art tools costing up to \$20K/year.
- Trained genomic foundation models to evaluate their susceptibility to adversarial data poisoning attacks.
- Led benchmarking of generative genomic models against real genomes, identifying systematic discrepancies in k-mer distributions, non-B DNA motifs, and transcription factor binding sites.

The Pennsylvania State University

Hershey, PA

Research Associate & Graduate Research Assistant

08/2023 – 08/2025

- Engineered novel algorithms in Python and Bash for the analysis of short DNA and protein sequences; 12 peer-reviewed publications (7 as first/corresponding author).
- Built scalable High-Performance Computing (HPC) pipeline in Slurm enabling the identification of pathogen biomarkers across 45,000 genome assemblies; led scaling to 500,000 assemblies.
- Developed the *kmerDB* database, consolidating information on short DNA and protein sequences across over 60K genomes and proteomes; co-developed 4 additional public scientific databases.
- Created open-source *Zseeker* Python tool; co-developed 5 additional open-source bioinformatics tools.
- Delivered oral presentations at 4+ national and international conferences, ranging from 150-300 attendees.
- Supervised 8+ researchers and coordinated collaborative projects across multiple research groups; mentored 5 junior researchers with no prior bioinformatics experience to their first first-author publications.
- Designed research projects and drafted grant applications resulting in \$3M+ in competitive funding.

Neomer Diagnostics

Brussels, Belgium

Co-founder and Chief Technical Officer

01/2022 – 05/2023

- Co-founded diagnostics startup translating patented nullomer research into clinical cancer detection platform.
- Developed machine learning (ML) pipeline for the detection of cancer from liquid biopsies, achieving AUC ranging from 0.89 to 0.94 in lung and ovarian cancers.
- Contributed to securing \$850K through technology development and investor demos.

Independent contractor

Brussels, Belgium

Software Engineering Consultant

02/2020 – 08/2023

- Optimized cancer genomics ML pipeline using Julia, achieving 50-fold speed increase, saving >3 months of computational time and \$50K+ in costs.
- Created distributed computing pipelines for analysis of 100+ TB of genomic data.
- Deployed and managed genomic analysis pipeline to Google Cloud Platform, enabling processing of 14 TB of cell-free DNA sequencing data.
- Engineered compression procedure achieving 100-fold reduction, enabling development of *kmerDB* database.
- Consulted on study design and grant writing, securing \$400K+ in awarded funding.

Education

The Pennsylvania State University

Hershey, PA

Ph.D. in Bioinformatics and Genomics (Cumulative GPA: 4.0)

08/2023 – 01/2026

Doctoral thesis: Computational and machine learning methods for k-mer sequence analysis in large-scale omics data

KU Leuven

Leuven, Belgium

M.Sc. in Artificial Intelligence (Honours: Cum Laude)

09/2020 – 07/2023

Master's thesis: Frequentmers – a k-mer based approach for the detection of liver cirrhosis gut microbiome mNGS

Aristotle University of Thessaloniki

Thessaloniki, Greece

B.Sc. in Mathematics (Honours: Very Good)

10/2013 – 03/2018

Skills

Technical skills:

- Programming Languages: Python (Pandas, NumPy, scikit-learn, PyTorch, Matplotlib, FastMCP), Bash, Julia
- Tools & Platforms: High-Performance Computing (Slurm, SGE, LSF), Google Cloud Platform, Git/GitHub
- Computational Methods: AI evals, genomic Language Models, empirical ML research, algorithm design and optimization, large-scale genomic data analysis, machine learning pipelines
- Research Skills: Study design, scientific writing, grant writing

Languages: English (Fluent), French (Fluent), German (Fluent), Japanese (Elementary), Greek (Native)

Honors and Awards

- Center for Molecular Carcinogenesis and Toxicology Symposium Abstract Award (1st place)
- Alumni Society Award, Penn State College of Medicine, 2024 (5% acceptance rate)
- University Graduate Fellow (1% acceptance rate)
- Huck Distinguished Fellow, Penn State, 2023 (3% acceptance rate)

Patents

- Georgakopoulos-Soares, I., **Mouratidis, I.**, Provatas, K. (2025). System and Method for Extracting Neomers. U.S. Provisional Patent.
- Ahituv, N., Yizhar-Barnea, O., Georgakopoulos-Soares, I., **Mouratidis, I.**, Hemberg, M. (2023). Systems for mutation caller and methods of using the same. WO2024103003A3.
- Ahituv, N., Yizhar-Barnea, O., Georgakopoulos-Soares, I., **Mouratidis, I.**, Hemberg, M. (2022). Compositions comprising nullomers and methods of using the same for cancer detection and diagnosis. WO2022235718A3.

Selected Publications

1. Georgakopoulos-Soares, I.*, Yizhar-Barnea, O.*, **Mouratidis, I.***, et al. (2025). Leveraging sequences missing from the human genome to diagnose cancer. *Communications Medicine*, 5(1).
2. Wang, G.*, **Mouratidis, I.***, Provatas, K., et al. (2025). ZSeeker: an optimized algorithm for Z-DNA detection in genomic sequences. *Briefings in Bioinformatics*, 26(3).
3. **Mouratidis, I.***, Konnaris, M. A.*, Chantzi, N.*, Chan, C. S. Y.*, et al. (2025). Identification of the shortest species-specific oligonucleotide sequences. *Genome Research*, 35(2).
4. **Mouratidis, I.***, Baltoumas, F. A.*, Chantzi, N.*, et al. (2024). kmerDB: A database encompassing the set of genomic and proteomic sequence information for each species. *Computational and Structural Biotechnology Journal*, 23.
5. **Mouratidis, I.**, Chantzi, N., Khan, U., et al. (2023). Frequentmers - a novel way to look at metagenomic next generation sequencing data and an application in detecting liver cirrhosis. *BMC Genomics*, 24(1).
6. **Mouratidis, I.**, Chan, C. S. Y., Chantzi, N., et al. (2023). Quasi-prime peptides: identification of the shortest peptide sequences unique to a species. *NAR Genomics and Bioinformatics*, 5(2).

Complete list of published work can be found in [Google Scholar](#).

* Equally contributing authors