Antonios P. Sarikas

☑ antonios.sarikas@gmail.com



Computational Chemist

If the implementation is easy to explain, it may be a good idea.



2024-TBA PhD, Physical & Computational Chemistry, Department of Chemistry, University of Crete

2022–2024 MSc, Physical & Computational Chemistry, Department of Chemistry, University of Crete

9.86 (Excellent)

2018-2022 BSc, Chemistry, Department of Chemistry, University of Crete

8.64 (Excellent)

First of the class - obtained in 3.5/4.0 years

Master thesis

Title From Potential Energy Surface to Gas Adsorption via Deep Learning

Supervisor George E. Froudakis

Description Developing a deep learning based method for predicting gas adsorption in porous materials using

3D energy images.

Repository https://github.com/adosar/master_thesis

Bachelor thesis

Title Screening of MOFs for H_2 Storage via Machine Learning

Supervisor George E. Froudakis

Description Developing a machine learning based method for fast screening of large databases, in order to

identify top performing MOFs for hydrogen storage.

Scholarships — Awards — Highlights

2025 Excellence Award "Professor Zoe Dimitriadis Prize", University of Crete

2025 Best MSc Thesis Award (3rd place), Federation of European Materials Societies

2025 Postgraduate Fellowship, Vasileios Apollon Anagnostakis Bequest Fellowship

2025 Best MSc Thesis Award (1st place), Hellenic Society for the Science and Technology of Condensed Matter

2025 Paper featured in $N\alpha ture\ Portfolio\ Nobel\ Chemistry\ 2025\ Collection$. "Gas adsorption meets deep learning: voxelizing the potential energy surface of metal-organic frameworks"

2025 Paper ranked 9th in *Scientific Reports 2024 Materials Science Top 100*. "Gas adsorption meets deep learning: voxelizing the potential energy surface of metal-organic frameworks".

2024 Postgraduate Fellowship, Independent Power Transmission Operator S.A. Fellowship

2024 Postgraduate Fellowship, Maria Michail Manasaki Bequest Fellowship

- 2023 Chatzimarinaki Award, Department of Chemistry, University of Crete
- 2022 Undergraduate Scholarship, Stamatiou Foundation
- Undergraduate Scholarship, Stamatiou Foundation 2021
- 2020 Undergraduate Scholarship, Stamatiou Foundation

¶ Publications

- Aydin Ozcan, François-Xavier Coudert, Sven M. J. Rogge, Greta Heydenrych, Dong Fan, Antonios P. Sarikas, Seda Keskin, Guillaume Maurin, George E. Froudakis, Stefan Wuttke, and Ilknur Erucar. "Artificial Intelligence Paradigms for Next-Generation Metal-Organic Framework Research". In: Journal of the American Chemical Society (June 2025). DOI: 10.1021/jacs.5c08214.
- Antonios P. Sarikas, Konstantinos Gkagkas, and George E. Froudakis. "Gas adsorption meets deep learning: [2] voxelizing the potential energy surface of metal-organic frameworks". In: Scientific Reports 14.1 (Jan. 2024). DOI: 10.1038/s41598-023-50309-8.
- Antonios P. Sarikas, Konstantinos Gkagkas, and George E. Froudakis. "Gas adsorption meets geometric deep learning: points, set and match". In: Scientific Reports 14.1 (Nov. 2024), p. 27360. poi: 10.1038/s41598-024-76319-8.
- Antonios P. Sarikas, George S. Fanourgakis, and George E. Froudakis. "Metal-organic frameworks in the age of machine learning". In: Reticular Chemistry and Applications. De Gruyter, Jan. 2023. poi: 10.1515/ 9781501524721.
- Antonios P. Sarikas, George S. Fanourgakis, Konstantinos Gkagkas, and George E. Froudakis. "Comparison of machine learning approaches for the identification of top-performing materials for hydrogen storage". In: Sustainable Chemistry for the Environment 5 (Mar. 2023), p. 100056. poi: 10.1016/j.scenv.2023. 100056.
- Antonios P. Sarikas, George S. Fanourgakis, Emmanuel Tylianakis, Konstantinos Gkagkas, and George E. [6] Froudakis. "Comparison of Energy-Based Machine Learning Descriptors for Gas Adsorption". In: The Journal of Physical Chemistry C (Oct. 2023). DOI: 10.1021/acs.jpcc.3c04223.

Peer-review activities

- 2025 Reviewer for Scientific Reports
- 2025 Reviewer for Communications Chemistry
- Reviewer for Journal of Cheminformatics





A dsorb - python package for deep learning on molecular point clouds

- O Homepage
- Documentation

A web app for predicting properties of porous materials via deep learning Aldsorb-online O Click to open the app



 \bigcap $\epsilon\lambda$ python package for parallel calculation of energy voxels

- O Homepage
- Documentation

RetNet A 3D ConvNet for Reticular Chemistry

Homepage

Presentations

- ₱ 2025 Young Scientist Symposium (EuroMOF2025), Heraklion
- 2025 EuroMOF2025, Heraklion
- 2025 FEMS Masters Thesis Award, Granada
- 2025 FEMS EUROMAT 2025, Granada
- **⊉** 2025 3C Summer School V2, Heraklion
- **2** 2024 COST Action EU4MOFs, Istanbul
- 2023 Chatzimarinaki Seminar, Heraklion
- 2023 1st Mediterranean Conference on Porous Materials, Rethymnon
- 2022 Chatzimarinaki Seminar, Heraklion
- 2022 XXXVI Pan-Hellenic conference on Solid-State Physics and Materials Science, Heraklion

Workshops

- 2024 Computational Materials Science, University of Thessaly, Online
- 2023 Computational Materials Science, National Kapodistrinan University of Athens, Online
- 2022 Computational Materials Science, University of Ioannina, Online
- 2021 Computational Materials Science, University of Patras, Online
- 2021 DCMS Materials 4.0 Summer School, TU Dresden, Online

Computer skills

Operating Fedora Linux, Ubuntu, Windows

Systems

Programming Python, Fortran

Languages

Markup LATEX, Markdown

Languages

Other Tools Git, Github, Sphinx, Vim, MSOffice

Working experience

- 2025 Coordinator for 3C Summer School V2, Department of Chemistry, University of Crete
- 2023, 2025 Teaching assistant on the Laboratory of Physical-Chemistry II, Department of Chemistry, University of Crete
- 2023, 2024 Teaching assistant on the Laboratory of Physical-Chemistry I, Department of Chemistry, University of Crete

A Z Languages

English Proficiency, University of Michigan

Greek Native language