Jakub Martinka

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in jakub-martinka

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EDUCATION

Ph.D. in Modeling of chemical nanostructures and biostructures

10/2022 - present

Thesis: Excited state molecular dynamics with non-adiabatic and spin-orbit effects assisted by machine learning Faculty of Science, Charles University

MSc. in Physical chemistry summa cum laude

10/2020 - 9/2022

Thesis: Non-adiabatic molecular dynamics of photochemical processes accelerated by machine learning

Faculty of Science, Charles University

BSc. in Biochemistry

10/2017 - 9/2020

Thesis: Molecular modeling of the interaction between cytochrome P450s and their substrates/products

Faculty of Science, Charles University

EXPERIENCE

J. Heyrovsky Institute of Physical Chemistry, CAS CR

12/2018 – present

- Worked at the department of Theoretical Chemistry. Contributed to Newton-X (C++), MLatom (Python). Experience with quantum chemistry packages: *Turbomole, Molpro, Molcas, Columbus, Gaussian, Molden*.

UniCredit Bank Czech Republic and Slovakia, a.s.

9/2020 – present

- Created, edited and maintained intranet pages.

JavaScript, HTML, CSS, SharePoint

Private lessons: I CAN IB, Top Learning s.r.o.

9/2015 – present

- Tutored in Math, Physics, and Chemistry.

Československá obchodní banka

7/2022 - 11/2022

- Worked with content of internet banking and edited JavaScript of PDF forms. JavaScript, SharePoint

Charles University, Faculty of Science, Department of Biochemistry

11/2020 - 5/2021

– Created 3D models of organelles and proteins for a new biochemistry textbook.

Blender, VMD

Summer schools, Workshops & Visits

Molecular Excited States Workshop

9 - 13/6/2025

- Got invited to workshop devoted to the theoretical modeling of the excited states of chemical molecules. My role is to present practical tutorials for machine learning applications in computational chemistry (Toruń, Poland).

Research visit (Xiamen University, China)

21/10 - 10/11/2024

 Completed a two-week stay with prof. Dral's research group, focusing on the application of ML in non-adiabatic molecular dynamics, including the implementation of the Fewest Switches Surface Hopping method in MLatom (Python).

Machine Learning for Chemistry Summer School

9/9 - 13/9/2024

Attended a summer school organized by KIT covering: Molecular descriptors and ML based molecular property prediction,
 GNNs, MLPs, Molecular synthesis prediction, Self-driving labs. (Karlsruhe, Germany)

Research visit (Aix-Marseille Université, France)

22/1 - 27/1/2024

 Participated in a week-long stay with prof. Barbatti's research group, focusing on the application of ML techniques in non-adiabatic molecular dynamics - learning and sign correction of nonadiabatic couplings (Python).

Modern Wavefunction Based Methods in Electronic Structure Theory

27/8 - 2/9/2023

Attended a summer school organized by Max-Planck-Institut f
ür Kohlenforschung, focusing on the theory/implementation
of quantum chemistry methods. (Pisa, Italy)

Autumn School of Quantum Chemistry

3 - 7/10/2020

Participated in hands-on sessions and implemented quantum chemical methods - Hartree-Fock, Configuration Interaction,
 Møller-Plesset perturbation theory on top of Psi4 (in Python). (Prague, Czechia)

Swiss Equivariant Learning Workshop

11 - 14/7/2022

- Attended a summer school organized by EPFL covering: equivariant machine learning techniques. (Lausanne, Switzerland)

Science Popularization Workshop

2022

 Participated in a workshop on the popularization of science organized by the J. Heyrovsky Institute of Physical Chemistry with communication experts from the Czech Academy of Sciences. (Prague, Czechia)

Excited States and Nonadiabatic Dynamics CyberTraining Workshop

14 - 26/6/2021

 Completed a remote workshop orginised by State University of New York at Buffalo focusing on non-adiabatic molecular dynamics methods, concluding with a small project and its presentation. (remote due to COVID-19)

Conferences

13th Triennial Congress of the WATOC (Oslo, Norway)	22 - 27/6/2025
11th Triennial Congress of the ISTCP (Qingdao, China)	13 - 18/10/2024
WTC Mariapfarr - Machine Learning for Chemistry (TU Graz, Austria)	27/2 - 1/3/2024
The International Symposium on ML in QCh (Uppsala University, Sweden)	29/11 - 1/12/2023
17th International Congress of Quantum Chemistry (Bratislava, Slovakia)	26/6 - 1/7/2023
ICQC Satellite meeting on Strong Correlation in Molecules (Znojmo, Czechia)	20 - 23/6/2023

Talks & Popularization

ChemQuest (evaluation of high-school student's project)	2024-5
Open science (leading a research intership for 5 high-school students)	2024-5
AI in context (talk covering applications of ML in Chemistry)	26/10/2023
Science fairs (presentation of group/department research)	2023 – 5
Ask a scientist (project focused on answering nonscientific questions)	since $5/2023$
Student conferences at J. H. Inst. (best talk awards)	2023-5
FameLab – Presentation & Masterclass training	$30/9/2022 \ \& \ 10 - 11/9/2022$
Falling Walls Lab – Presentation	13/9/2022

Publications

- [1] Martinka J., Zhang L., Hou Y.-F., Martyka M., Pittner J., Barbatti M. and Dral P. O., "A Descriptor Is All You Need: Accurate Machine Learning of Nonadiabatic Coupling Vectors", *submitted to JPCL* (2025).
- [2] Martinka J., Pederzoli M., Barbatti M., Dral P. O. and Pittner J., "A Simple Approach to Rotationally Invariant Machine Learning of a Vector Quantity", *Journal of Chemical Physics*, **161**, 174104 (2024).

SKILLS & INTERESTS

Grants: GAUK 2024 (project 6224), POINT 2024, GACR (project 23-06364S), AMULET project

Language: English C1 German A2 Japanese A0 Czech (Native)

Computer: C/C++ Python UNIX/bash/vim HTML/CSS git/GitHub LaTeX

Interests: Science popularization, History of physics, Travelling, Chess, Rock climbing, Football, FOSS