Umberto Raucci - Resume

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Research Interests

ab initio molecular dynamics, nonadiabatic molecular dynamics, QM/MM, photochemistry, mechanochemistry, reaction discovery, neural network potentials, heterogeneous catalysis, natural user interfaces for quantum chemistry

Research Experience

2021-	Postdoctoral Researcher - Advisor: Prof. Michele Parrinello
present	Italian Institute of Technology, Genova, Italy
2019-2021	Postdoctoral Researcher - Advisor: Prof. Todd J. Martínez
	Stanford University, CA, USA
10/2018-	Postdoctoral Researcher - Advisor: Prof. Carlo Adamo
12/2018	Chimie Paris Tech, PSL University collaboration with L'Oréal Paris, Paris, France
2016-2018	Postdoctoral Researcher - Advisor: Prof. Nadia Rega
	University of Naples Federico II, Naples, Italy
05/2015-	Visiting Student - Advisor: Prof. Carlo Adamo
07/2015	Chimie Paris Tech, PSL University, Paris, France

Academic Education

2013-2016	PhD in Chemical Sciences
	University of Naples Federico II, Naples, Italy
2009-2012	Master in Chemical Sciences (110/110)
	University of Naples Federico II, Naples, Italy
2006-2009	Bachelor in Chemistry (summa cum laude)
	University of Naples Federico II, Naples, Italy

Participation in workshops and training courses

2022	PyTorch for Deep Learning with Python Bootcamp, Udemy (online)
2022	Complete Python Bootcamp from Zero to Hero in Python, Udemy (online)
2020	Build Websites from Scratch with HTML and CSS, Udemy (online)
2018	Theoretical Chemistry Summer School, Center for Quantum Molecular Design,
	Stanford University, Stanford, California (USA)
2016	VII National School of Photochemistry, Department of Chemistry "G.Ciamician",
	University of Bologna, Bologna, Italy

Scientific Software Contributions

2021 2021 2020	TeraChem Web Services: web-based interface for quantum chemistry MolAR: iOS mobile application for visualizing molecules in augmeted reality ChemVox: Alexa skill for voice-controlled quantum chemistry
	Awards
2021	Wiley Outstanding Postdoc Award, awarded by the American Chemical Society (COMP Division)
2021	Best Flash Communication Award at Sygenta Simposium 2021, awarded by the Swiss Chemical Society and Helvetica Chimica Acta
2017	Guido Barone Award for the best Ph.D thesis, awarded by the Italian Chemical Society (Campania Division)
2016	Best Poster Communication Award at IV National Meeting of the Theoretical and Computational Chemistry division of the Italian Chemical Society
2013	Luigi Gomez Paloma Award for the best master thesis, awarded by the Italian Chemical Society (Campania Division)
	Teaching Experience
2017	Instructor at the Gaussian Workshop, Santiago De Compostela, Spain
2017	Coordination Compound Chemistry for master students in Chemical Sciences at the University of Naples Federico II – Assistance for lectures
2013-2018	Computational Chemistry for master students in Chemical Sciences at the University of Naples Federico II – Assistance for lectures and laboratory
2016-2018	Topics in Physical Chemistry for master students in Chemical Sciences at the University of Naples Federico II – Assistance for lectures and laboratory
2016-2018	Chemical Physics for bachelor students in Food science at the University of Naples Federico II – Assistance for lectures
	Mentoring of one PhD and three master students resulted in publications [5], [10], [15], [19].
	Professional Services
	Journal Reviewer
2018- present	Mol. Catal., Theor. Chem. Acc., Phys. Chem. Chem. Phys., ChemistrySelect, Comput. Biol. Chem, J. Org. Chem.
2020	Meeting Organization Member of the organization committee of VCTC 2020
2018-2019	Italian Chemistry Olympiad for high school students
2016-2018	Open Day, Department of Chemical Sciences, University of Naples Federico II
2017 2016	Staff member of the XXVI National Meeting of the Italian Chemical Society Staff member of the XLIV National meeting of the Physical Chemistry division of the Italian Chemical Society

Invited Seminar

Modeling Artificial Leaf

Heinrich Heine University Düsseldorf, Germany - Host: Prof. Jan Meisner Molecular Dynamics Accelerated Reaction Discovery
 University of Helsinki, Finland - Host: Prof. Theo Kurten Enhanced Sampling Aided Reaction Discovery
 Stanford University, CA, USA - Host: Prof. Todd Martinez A Stairway to Heaven of Photoinduced Charge Transfer Reactions
 KAUST University, Saudi Arabia - Host: Prof. Luigi Cavallo

Conference Contributions (selection)

28 contributions in national and international conference proceedings

2022 Cutting-Edge Technologies in Computational Chemistry, contributed poster at the

American Chemical Society Spring Meeting

20 ChemVox: Voice-Controlled Quantum Chemistry, contributed talk at the 2020 TeraChem Developers Meeting

Ab Initio Molecular Dynamics to Simulate Excited State Proton Transfer to Solvent: The Strange Case of a Super Photoacid in Water and Methanol Solution, contributed talk at the XXVI National Meeting of the Italian Chemical Society

Modeling of Proton Coupled Electron Transfer in the Framework of Density Functional Theory, contributed talk at **Pacifichem**

Ab Initio Molecular Dynamics Combined with Different Solvation Models for Simulating Excited State Proton Transfer, contributed talk at **Pacifichem**

Publications

30. Raucci U., Weir H., Sakshuwong S., Seritan S., Hicks C., Vannucci F., Rea F., Martínez T. J., *Interactive Quantum Chemistry Enabled by Machine Learning, Graphical Processing Units, and Cloud Computing*, **Annu. Rev. Phys. Chem.**, 2023, 74, 1

29. Raucci U., Sanchez D. M., Martínez T. J., Parrinello M., Enhanced Sampling Aided Design of Molecular Photoswitches, J. Am. Chem. Soc., 2022, 144, 19265 28. Stricker F., Sanchez D. M., Raucci U., Dolinski N. D., Zayas M. S., Meisner J., Hawker C. J., Martínez T. J., Read de Alaniz J., A Multi-Stage Single Photochrome System for Controlled Photoswitching Responses, Nat. Chem., 2022, 14, 942

- 27. Raucci U., Weir H., Bannwarth C., Sanchez D. M., Martínez T.J., *Chiral Photochemistry of Achiral Molecules*, Nat. Commun., 2022, *13*, 2091
- **26**. Raucci U., Rizzi V., Parrinello M., *Discover, Sample, and Refine: Exploring Chemistry with Enhanced Sampling Techniques*, **J. Phys. Chem. Lett.**, 2022, 13, 1424
- 25. Sakshuwong S., Weir H., <u>Raucci U.</u>, Martínez T.J., <u>Bringing Chemical Structures to Life with Augmented Reality, Machine Learning, and Quantum Chemistry</u>, **J. Chem. Phys.**, 2022, 156, 204801
- 24. Sanchez D.M., Raucci U., Martínez T.J., In Silico Discovery of Multistep

2020

2017

2015

2023

2022

2021

- Chemistry Initiated by a Conical Intersection: The Challenging Case of Donor-Acceptor Stenhouse Adducts, J. Am. Chem. Soc., 2021, 48, 20015
- 23. Raucci U., Valentini A., Pieri E., Weir H.V., Seritan S., Martínez T.J. Voice-Controlled Quantum Chemistry, Nat. Comput. Science, 2021, 1, 42
- **22**. Chiariello M. G., Donati G., <u>Raucci U.</u>, Perrela F., Rega N., <u>Structural Origin and Vibrational Fingerprints of the Ultrafast Excited State Proton Transfer of the Pyranine-Acetate Complex in Aqueous Solution</u>, **J. Phys. Chem. B**, 2021, 36, 10273
- **21**. Chiariello M. G., <u>Raucci U.</u>, Donati G., Rega N., Water Mediated Excited State Proton Transfer of Pyranine-Acetate in Aqueous Solution: Vibrational Fingerprints from Ab-Initio Molecular Dynamics, **J. Phys. Chem. A**, 2021, 17, 3569
- **20**. Coppola F., Cimino P., <u>Raucci U.</u>, Chiariello M. G., Petrone A., Rega N., Exploring the Franck-Condon Region of a Photoexcited Charge Transfer Complex in Solution via Femtosecond Stimulated Raman Spectroscopy: Excited State Electronic Structure Methods to Unveil Non-Radiative Pathways, **Chem. Sci.**, 2021, 12, 8058
- 19. Tirri B., Mazzone G., Ottochian A., Gomar J., <u>Raucci U.</u>, Adamo C., Ciofini., A Combined Monte Carlo/DFT Approach to Simulate UV-vis Spectra of Molecules and Aggregates: Merocyanine Dyes as a Case Study, J. Comput. Chem., 2021, 42, 1054
- 18. Sanchez D.M., <u>Raucci U.</u>, Ferreras K. N, Martínez T.J., <u>Putting Photomechanical Switches to Work: An Ab Initio Multiple Spawning Study of Donor-Acceptor Stenhouse Adducts</u>, J. Phys. Chem. Lett., 2020, 11, 7901
- 17. Raucci U., Savarese M., Adamo C., Ciofini I., Rega N., Modeling the Electron Transfer Chain in an Artificial Photosynthetic Machine, J. Phys. Chem. Lett., 2020, 11, 9738
- **16**. Raucci U., Chiariello M. G., Rega N., Modeling Excited State Proton Transfer to Solvent: a Dynamics Study of a Super-Photoacid with a Hybrid Implicit/Explicit Solvent Model, J. Chem. Theory and Comput., 2020, 16, 7033
- **15**. Raucci U., Perrella F., Donati G., Zoppi N., Petrone A., Rega N., Ab-initio Molecular Dynamics and Hybrid Explicit-Implicit Solvation Model for Aqueous and non Aqueous Solvents: GFP Chromophore in Water and Methanol Solution as Case Study, J. Comput. Chem., 2020, 41, 2228
- 14. <u>Raucci U.</u>, Chiariello M. G., Coppola F., Perrella F., Savarese M., Ciofini I., Rega N., *An Electron Density Based Analysis to Establish the Electronic Adiabaticity of Proton Coupled Electron Transfer Reactions*, **J. Comput. Chem.**, 2020, 41, 1835
- 13. Platella C., <u>Raucci U.</u>, Rega N., D'Atri S., Levati L., Roviello G., Fuggetta M.P., Musumeci D., Montesarchio D., <u>Shedding Light on the Interaction of Polydatin and Resveratrol with G-Quadruplex and Duplex DNA: a Biophysical, Computational and Biological Approach, Int. J. Biol. Macromol., 2020, 151, 1163</u>
- 12. Esposito R., <u>Raucci U.</u>, Cucciolito M. E., Di Guida R., Scamardella C., Rega N., Ruffo F., *Iron(III) Complexes for Highly Efficient and Sustainable Ketalization of Glycerol: A Combined Experimental and Theoretical Study*, **ACS Omega**, 2019, 4, 688
- 11. Chiariello M. G., <u>Raucci U.</u>, Coppola F., Rega N. *Unveiling Anharmonic Coupling by Means of Excited State Ab Initio Dynamics: Application to Diarylethene Photoreactivity*, **Phys. Chem. Chem. Phys.**, 2019, 21, 3606

2020

2019

- 10. Perrella F., <u>Raucci U.</u>, Chiariello M.G., Chino M., Maglio O., Lombardi A., Rega N., *Unveiling the Structure of a Novel Artificial Heme-Enzyme with Peroxidase-Like Activity: A Theoretical Investigation*, **Biopolymers**, 2018, 109, e23225
- 9. Battista E., Scognamiglio P.L., Di Luise N., <u>Raucci U.</u>, Donati G., Rega N., P.A. Netti, Causa F., *Turn-On Fluorescence Detection of Protein by Molecularly Imprinted Hydrogels Based on Supramolecular Assembly of Peptide Multi-Functional Blocks*, J. Mater. Chem. B, 2018, 6, 1207

2017

8. Savarese M., <u>Raucci U.</u>, Fukuda R., Adamo C., Ehara M., Rega N., Ciofini I., Comparing the Performance of TD-DFT and SAC-CI Methods in the Description of Excited States Potential Energy Surfaces: an Excited State Proton Transfer Reaction as Case Study, J. Comput. Chem., 2017, 38, 1084

2016

- 7. Raucci U., Ciofini I., Adamo C., Rega N., Unveiling the Reactivity of a Synthetic Mimic of the Oxygen Evolving Complex, J. Phys. Chem. Lett., 2016, 7, 5015
- **6**. Savarese M., <u>Raucci U.</u>, P.A. Netti, Adamo C., Rega N., Ciofini I., *A Qualitative Model to Identify Non-Radiative Decay Channels: the Spiropyran as Case Study*, **Theor. Chem. Acc.**, 2016, *135*, 211
- **5**. Cimino P., <u>Raucci U.</u>, Donati G., Chiariello M.G., Schiazza M., Coppola F., Rega N., *On the Different Strength of Photoacids*, **Theor. Chem. Acc.**, 2016, 135, 117

2015

4. Raucci U., Savarese M., Adamo C., Ciofini I., Rega N., Intrinsic and Dynamical Reaction Pathways of an Excited State Proton Transfer, J. Phys. Chem. B, 2015, 119, 2650

2014

- 3. Savarese M., <u>Raucci U.</u>, Adamo C., Netti P.A., Ciofini I., Rega N., *Non Radiative Decay Paths in Rhodamines: New Theoretical Insights*, **Phys. Chem. Phys.**, 2014, 16, 20681
- 2. Savarese M., <u>Raucci U.</u>, Netti P.A., Adamo C., Ciofini I., Rega N., <u>Modeling of Charge Transfer Processes to Understand Photophysical Singatures: The Case of Rhodamine 110, Chem. Phys. Lett.</u>, 2014, 610, 148
- 1. Cusano A.M., Causa F., Della Moglie R., Falco N., Scogliamiglio P.L., Aliberti A., Vecchione R., Battista E., Marasco D., Savarese M., <u>Raucci U.</u>, Rega N., Netti P.A., <u>Integration of Binding Peptide Selection and Multifunctional Particles as Tool-Box for Capture of Soluble Protein in Serum</u>, J. R. Soc. Interface, 2014, 11, 20140718