Stefano Racioppi, Ph.D.

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Professional Experience

09/2025 – present	Research Associate, University of Cambridge, UK.
	Group: Prof. Chris Pickard.

03/2021 – 08/2025 Postdoctoral researcher, State University of New York at Buffalo, USA.

Group: Prof. Eva Zurek.

07/2023 – 09/2023 Visitor, *Aalto University*, Finland.

Group: Prof. Patrick Rinke.

02/2019 – 02/2021 Postdoctoral researcher, Chalmers University of Technology, Sweden.

Group: Prof. Martin Rahm.

Education

10/2015 - 01/2019	Ph.D. in Industrial Chemistry, University of Milan, Italy. Supervisors: Prof. Angelo Sironi;
	Prof. Piero Macchi; Prof. Pierluigi Mercandelli.
11/2018 - 01/2019	Scholarship recipient, National Interuniversity Consortium of Materials Science and
	Technology (INSTM), University of Milan, Italy. Supervisor: Prof. Angelo Sironi.
10/2016 - 03/2018	Visiting student at the <i>University of Bern</i> , Switzerland. Host: Prof. Piero Macchi.
10/2013 - 10/2015	Master's degree in Industrial Chemistry, University of Padua, Italy. Supervisors: Prof.
	Armando Gennaro; Prof. Laura Orian. Final grade: 110/110
10/2010 - 09/2013	Bachelor's degree in Industrial Chemistry, University of Padua, Italy. Supervisor: Prof.

Research Interests

- Computational and Theoretical Chemistry.
- Quantum Crystallography and Crystal Structure Prediction.

Armando Gennaro.

- High Pressure Chemistry.
- Chemical Bonding.
- Material and Planetary Science.
- Superconductivity.
- Catalysis.

Distinctions

03/2025 Marie Skłodowska-Curie Actions (MSCA) Postdoctoral Fellowship, Seal of Excellence 2025.

09/2019 Best Ph.D. Thesis Award by the Italian Crystallographic Association (AIC).

07/2017 Outstanding poster prize offered by Wiley – Journal of Computational Chemistry and International Journal of Quantum Chemistry. Workshop on ab Initio Modeling in Solid State Chemistry with CRYSTAL, *Minneapolis*, USA.

Memberships

2025 – present	Trinity Hall College of Cambridge
2024 – present	European Crystallographic Association (ECA).
2021 – present	American Physical Society (APS).
2018 – present	Italian Crystallographic Association (AIC).
2018 - 2019	Swiss Society for Crystallographic (SGK/SSCr).

External Fundings, Proposals and Scholarships

05/2025	Awarded: Proposal experiment, Structural Determination of Al_2O_3 at super-Earth Conditions
	- AlO Discovery, Omega Laser Facility at the Laboratory for Laser Energetics (LLE); 1 shot
	day awarded. PI: Dr. Racioppi S.
04/2025	Awarded: Beamtime at the European Synchrotron Radiation Facility (ESRF); Diagnosing
	Electride Formation in Ca up to 40 GPa with XANES and SCXRD; 18 shifts awarded. PI: Dr.
	Chin, D. A.; co-PI: Dr. Racioppi, S.
04/2024	Awarded: 34 th European Crystallographic Meeting Scholarship; € 300.

09/2023 – 08/2026 Contributed to the proposal's writing: *High Energy Density Quantum Matter - Renewal*; Award number: LOI-0000043230; Source of Support: Department of Energy (DOE); PI: Dr. Collins G.W.; Co-PIs: Dr. Deemyad S.; Dr. Hemley R. J.; Dr. Zurek E.; \$ 2,100,000.

02/2023 Awarded: Beamtime at Laser-lab Europe LENS; *Tuning Electrical Conductivity of Aurophilic Interactions by Compression* (ID: 22396); PIs: Dr. Racioppi S., Dr. Poreba T.; Prof. Macchi P.

09/2022 Awarded: Seed Funding Project; Center for Matter at Atomic Pressure (CMAP); \$ 3500.

Awarded: Travelling Grant; Ettore Majorana Foundation; € 600.

10/2018 – 01/2019 Awarded: Scholarship; National Interuniversity Consortium of Materials Science and

Technology (INSTM); PI: Prof. Sironi A.; € 5862.

Publications

05/2018

Total number of publications: 24 (16 as first author, 6 as corresponding author (*)).

Total number of citations: 212.

H-index: 9.

- (25.) Ocampo, I. K.; Kim, D.; Smith, R. F.; Coppari, F.; **Racioppi, S.**; Zurek, E.; Rygg, J. R.; Chin, D. A.; Millot, M.; Eggert, J. H.; Duffy, T. S.; Atomic-level structure and pressure-density response of Fe-O compounds at deep Earth and exoplanetary interior conditions, *under review* in *Physics of the Earth and Planetary Interiors*.
- 24. Coleman, A. L.; Singh, S.; Lockard, T.; Ocampo, I. K. Lazicki, A. E.; Gorman, M. G.; **Racioppi, S.**; Krygier, A. G.; Wehrenberg, C. E.; Ahmad, R.; Hamel, S.; Han, S.; Ginnane, M. K.; Swift, D. C.; Bonev, S. A.; Zurek, E.; Duffy, T. S.; Eggert, J. H.; McNaney, J.; Smith, R. F. Body-centered cubic phase transformation in gold at TPa pressures. *Accepted* in *PRL*.
- 23. **Racioppi, S.***; Zurek, E. Using Topology to Predict Electrides in the Solid State, *Arxiv*, arXiv:2508.04548, *Accepted* in *J. Phys. Chem. A.*
- 22. **Racioppi, S.***; Zurek, E. Activation of Semicore Electrons in Alkali Metals and Their Role in the B1–B2 Phase Transition under Pressure, *J. Am. Chem. Soc*, **2025**, *147*, 36, 32745-32751.
- 21. Storm, C. V.; Racioppi, S.; Duff, M. J.; McHardy, J. D.; Zurek E.; McMahon, M. I. Experimental Evidence of Interstitial Electron Density in Transparent Dense Sodium, *Commun. Mater*, **2025**, *6*, 201.
- 20. **Racioppi, S.**; Zurek, E. High-Pressure Electrides: A Quantum Chemical Perspective, *Annu. Rev. Mater. Res.*, **2025**, *55*, 421-442.
- 19. **Racioppi, S.**; Saffarin-Deemyad, I.; Holle, W.; Belli, F.; Ferry, R.; Kenney-Benson, C.; Smith, C. J., Zurek, E.; Deemyad, S. Phase Lithium's low-temperature phase transitions: Insights into quantum lattice dynamics and superconductivity, *PRB*, **2025**, *111*, 054111.
- 18. **Racioppi, S.**; Zurek, E. Looking at High-Pressure Electrides Through the Lens of Quantum Crystallography: The Case of Simple Cubic Calcium, *Acta Cryst.*, **2025**, B81.
- 17. **Racioppi, S.**; De la Roza, A.; Hajinazar, S.; Zurek, E. Powder X-Ray Diffraction Assisted Evolutionary Algorithm for Crystal Structure Prediction, *Digit. Discov.*, **2025**, 4, 73-83.
- 16. **Racioppi, S.**; De la Roza, A.; Hajinazar, S.; Zurek, E. PXRD-Assisted Crystal Structure Predictions, *Acta Cryst.*, **2024**, A80, e396.
- 15. **Racioppi, S.**; P.; Hyldgaard, P.; Rahm, M. Quantifying Atomic Volume, Partial Charge and Electronegativity in Condensed Phases, *J. Phys. Chem. C*, **2024**, 128, 4009-4017.
- 14. **Racioppi, S.***; Storm, C. V.; McMahon, M. I.; Zurek, E. On the Electride Nature of Na-hP4, *Angew. Chem. Int. Ed.* **2023**, e202310802.
- 13. **Racioppi, S.**; Miao, M.; Zurek, E. Intercalating Helium into A-site Vacant Perovskites, *Chem. Mater.* **2023**, 35, 4297-4310.
- 12. Fang, M.; Srikanth Kumar, G.; **Racioppi, S.**; Zhang, H.; Zurek, E.; Lin, Q. Hydrazonyl Sultones as Stable Tautomers of Highly Reactive Nitrile Imines for Fast Bioorthogonal Ligation Reaction, *J. Am. Chem. Soc.* **2023**, 145, 18, 9959-9964.
- 11. **Racioppi, S.**; Lolur, P.; Hyldgaard, P.; Rahm, M. A Density Functional Theory for the Average Electron Energy, *J. Chem. Theory Comput.* **2023**, 19, 799-807.
- 10. Antle, J.; Kimura, M.; **Racioppi, S.**; Lang, M.; Damon, C.; Gatley-Montross, C.; Sánchez B., L.; Miller, D.; Zurek, E.; Brown, A.; Gast, K.; Simpson, S. Applying Density Functional Theory to Common Organic Mechanisms: A Computational Exercise, *J. Chem. Educ.* **2023**, 100, 355-360.
- 9. **Racioppi, S.**; Orian, L.; Gennaro, A.; Isse, A. Solvent Coordination Effect on Copper-Based Molecular Catalysts for Controlled Radical Polymerization, *Catalysts* **2022**, 12, 1656.
- 8. Poręba, T.; **Racioppi, S.***; Garbarino, G.; Morgenroth, W.; Mezouar, M. Investigating the Structural Symmetrization of CsI₃ at High Pressures through Combined X-ray Diffraction Experiments and Theoretical Analysis, *Inorg. Chem.* **2022**, 61, 109777-10985.

- 7. Della Pergola, R.; Garlaschelli, L.; Macchi, P.; Ruffo, R.; Racioppi, S.; Sironi, A. From Small Metal Clusters to Molecular Nanoarchitectures with Core-Shell Structure: Synthesis, Redox Fingerprint, Theoretical Analysis and Solid-State Structure of [Co₃₈As₁₂(CO)₅₀]⁴⁻, *Inorg. Chem.* **2022**, 61, 9888-9896.
- Kumar, G. S.; Racioppi, S.; Zurek, E.; Lin, Q. Superfast Tetrazole-BCN Cycloaddition Reaction for 6. Bioorthogonal Protein Labeling on Live Cells, J. Am. Chem. Soc. 2022, 144, 57–62.
- 5. Racioppi, S.; Rahm M. In-Situ Electronegativity and the Bridging of Chemical Bonding Concepts, *Chem.* Eur. J. 2021, 27, 18156-18167.
- Racioppi, S.*; Sironi, A.; Macchi, P. On Generalized Partition Methods for Interaction Energies, Phys. 4. Chem. Chem. Phys. 2020, 24291-24298.
- Racioppi, S.; Andrzejewski, M.; Colombo, V.; Sironi, A.; Macchi, P. Different Metallophilic Attitudes 3. Revealed by Compression, Inorg. Chem. 2020, 59, 2223–2227.
- Racioppi, S.*; Della Pergola, R.; Colombo V.; Sironi, A.; Macchi, P. Electron Density Analysis of Metal 2. Clusters with Semi-Interstitial Main Group Atoms. Chemical Bonding in [Co₆X(CO)₁₆] Species, J. Phys. Chem. A 2018, 122, 5004-5015.
- Della Pergola, R.; Sironi, A.; Colombo, V.; Garlaschelli, L.; Racioppi, S.; Sironi, A.; Macchi, P. 1. Periodical trends in [Co₆E(CO)₁₆]⁻ clusters: Structural, synthetic and energy changes produced by substitution of P with As, J. Organomet. Chem. 2017, 849-850, 130-136.

Invited and Contributed Talks and Seminars

Talks

- 24th Biennial International Conference on the Science of Compression in Condensed Matter (SCCM): 06/2025 Computational Methods: Theory, Codes and Applications (FMDA/B).
- Crystal Structure Prediction Workshop, a Practical Approach, Poitier. 06/2025
- 4th European Symposium on Chemical Bonding (CBOND2024). 09/2024
- 34th European Crystallographic Meeting (ECM34); Microsymposium M27: High Pressure 08/2024 Crystallography: Exploring Structure and Method Development at Extreme Conditions.
- Gordon Research Seminar (GRS), Research at High Pressure.* 07/2024
- 28th AIRAPT and 60th EHPRG International Conference on High Pressure Science and Technology; 07/2023 Session: Chemical Bonding.
- American Physical Society (APS) March Meeting; Session G24: Matter at Extreme Conditions: Phase 03/2022
- 09/2021 Italian Crystallographic Association (AIC) XLIX Meeting.*
- 06/2018 International School of Crystallography in Erice; 52nd Course: Quantum Crystallography.

Seminars

- Cambridge Crystallographic Data Centre (CCDC), Virtual Seminar; Host: Dr. Fabio Montisci.* 05/2025
- High Energy Density Science Center (HEDS) Virtual Seminar Series; Lawrence Livermore National 07/2024 Laboratory (LLNL, USA); Host: Dr. Federica Coppari.*
- 01/2024 University of Milan (Italy), Dep. of Chemistry; Host: Prof. Davide Proserpio.*
- Aalto University (Finland), Dep. of Physics; Host: Prof. Patrick Rinke.* 08/2023
- 06/2023 University of Oviedo (Spain), Dep. of Chemistry; Host: Prof. José Manuel Recio.*
- Italian Young Crystallographers (GCI) Happy Hour Virtual Seminar; Host: Dr. Marta Morana.* 04/2023
- Chicago/DOE Alliance Center (CDA) Virtual Seminar Series; Host: Prof. Russell Hemley.* 03/2022
- * = invited

Conference and Workshop Appointments

- 07/2026 Elected chair of the Gordon Research Seminar (GRS), Research at High Pressure.
- 07/2025 Co-organizer of the "High Energy Density Physics: Quantum Materials and Quantum Phenomena" session at the 24th Biennial Conference on Science of Compression in Condensed Matter (SCCM).
- Co-organizer of a workshop on the crystal structure prediction code XtalOpt (co-organizer: Prof. Eva 06/2025 Zurek) at the Crystal Structure Prediction Workshop, a Practical Approach, held in Poitier, France.
- 05/2025 Moderator at the *Graduate Student Symposium* (GSS) at SUNY Buffalo.
- Co-organizer of a workshop on the crystal structure prediction code XtalOpt (co-organizer: Prof. Eva 06/2022 Zurek) at the International School of Crystallography in Erice; 56th Course: Crystallography under extreme conditions.

O6/2022 Co-organizer of a workshop on computational chemistry techniques (co-organizer: Dr. Julia Contreras-García) at the International School of Crystallography in Erice; 56th Course: Crystallography under extreme conditions.

Teaching and Supervision

2021/2025	Co-supervisor of three PhD students (Morgan Reddington, Masashi Kimura, Isaac Ampersand-
	Kwadwo Akwetey) and supervisor of two undergraduate students (Amelia Kelly, Jarrod DuMond).
01/2024	Special virtual lecture for the students of the general chemistry course at the University of Milan –
	Host: Prof. Davide Proserpio. "History and Evolution of the Electronegativity"
2022 - 2023	Hosted online workshops for the usage of the crystal structure prediction code XtalOpt.
2021	Supervisor of a high school student (Henry Feng) for summertime project.
2017 - 2019	Co-supervisor of two master students (Stephanie Terruzzi and Simona Sorbara).
2018/2019	Teaching assistant, physical chemistry II, 2 nd year degree in Industrial Chemistry.
2015/2016	Teaching assistant, general and inorganic chemistry, 3 rd year degree in Physics.

Computational Developments

Electronegativity in gas-phase: https://github.com/SteRacioppi/Chi-bar-Atomic-and-Molecular-Electronegativity Electronegativity in solid: https://www.scm.com/doc/BAND/Expert Options/Restarts.html

Public Service and Outreach

- 2025 Interviewed by News Center News & information from UB, Core electron bonding may not always require extreme pressure, study finds, regarding our paper "Activation of Semicore Electrons in Alkali Metals and Their Role in the B1-B2 Phase Transition under Pressure" published in JACS.
- Interviewed by *Science News*, <u>A new iron compound hints 'primordial' helium hides in Earth's core</u>, regarding the discovery of helium bearing iron at high pressure.
- Interviewed by News Center News & information from UB, Sodium's high-pressure transformation can tell us about the interiors of stars, planets, regarding our paper "On the Electride Nature of Na-hP4" published in Angewandte Chemie.