University of Chicago, Pritzker School of Molecular Engineering, 5640 S. Ellis Ave., Chicago, IL, 60637, USA

<u>arpank@uchicago.edu</u> | <u>arpan.kundu@gmail.com</u> | +1 773 717 3702 Home Page | Google Scholar | ResearchGate | ORCID | Linked-in | GitHub

EDUCATION

Humboldt-Universität zu Berlin

Berlin, Germany

Ph. D. (with "summa cum laude") in physical and theoretical chemistry

April 2018

Thesis: Ab initio prediction of isotherms for pure and mixed gas adsorption in metal-organic frameworks.

Adviser: Prof. Joachim Sauer

Indian Institute of Technology (IIT) Kanpur

Kanpur, India

M.Sc in Chemistry (GPA 9.7/10, 1st rank)

May 2011

Thesis: Dynamical behavior of supercritical aqueous solutions: orientational motion and angular jumps.

Adviser: Prof. Amalendu Chandra

Jadavpur University

Kolkata, India

B.Sc with honours in Chemistry (1st class with distinction, Marks 77.7%, 6th rank)

May 2009

RESEARCH INTERESTS

• Development and application of *ab initio* methods for (i) electron-phonon interactions in disordered solids and polymers (ii) spin-phonon interactions for quantum technologies, (iii) gas conversion technologies.

RESEARCH EXPERIENCE

University of Chicago, Pritzker School of Molecular Engineering (PME)

Chicago, IL, USA

Post doctoral scholar, Adviser: Prof. Giulia Galli

Feb 2019 – Present

- Combining density functional theory (DFT) with (i) molecular dynamics (MD) (ii) stochastic methods, for studying electron-phonon interactions in crystalline and amorphous solids.
- Developed a python interface for coupling (i) i-PI: an universal force engine and (ii) Qbox: an efficient DFT solver.
- Developed a python package named <u>pyEPFD</u> for performing electron-phonon calculations using (i) frozen phonon, (ii) stochastic methods.

Humboldt-Universität zu Berlin, Institute für Chemie

Berlin, Germany

Ph.D student and post doctoral researcher, Adviser: Prof. Joachim Sauer

Nov 2012 – Jan 2019

- Development of an ab initio methodology for the prediction of pure and mixed gas adsorption in porous solids using quantum chemistry and statistical mechanics.
- Development of a FORTRAN code named <u>LGCMC</u> for simulating pure and mixed gas adsorption isotherm using grand canonical Monte Carlo simulations on a coarse-grained lattice of adsorption sites.

Indian Institute of Science, Solid State and Structural Chemistry Unit

Bangalore, India

Research fellow

Aug 2011 – July 2012

• Investigation of transport of water molecules within carbon nanotubes using force field based molecular dynamics simulations.

Indian Institute of Technology (IIT) Kanpur, Department of Chemistry

Kanpur, India

M.Sc project student

Jan 2011 – Apr 2011

• Investigation of orientational relaxation and hydrogen bond breaking dynamics of supercritical water and aqueous solutions using force field based molecular dynamics simulations.

Indian Institute of Technology (IIT) Kharagpur, Department of Civil Engineering

Kharagpur, India

Summer research student

May 2010-July 2010

 Wet synthesis of As⁰ and Au⁰ nanoparticles (of tunable dimension) and their application in surface-enhanced Raman scattering (SERS).

PUBLICATIONS

- * = Corresponding Author , † = Equal contribution
- **1. Kundu, A.***, Galli, G.* "Quantum vibronic effects on the excitation energies of the nitrogen-vacancy center in diamond", *Submitted*, 2024. (co-corresponding author)
- **2. Kundu, A.***, Galli, G.* "Quantum vibronic effects on the electronic properties of molecular crystals", <u>J. Chem. Theory Comput. **2023**</u>, 19, 13, 4011. (co-corresponding author).
- **3.** Sillar, K.*, **Kundu, A.***, Sauer. J. "Ab initio prediction of adsorption selectivities for binary gas mixtures on a heterogeneous metal-organic framework surface", <u>J. Phys. Chem. C 2023</u>, 127, 27, 13317. (co-corresponding author)
- **4.** Yang, H., Govoni, M.*, **Kundu, A.**, Galli, G.* "Computational protocol to evaluate electron-phonon interactions within density matrix perturbation theory", <u>J. Chem. Theory Comput. **2022**</u>, 18, 10, 6031.
- **5. Kundu, A.***, Song, Y., Galli, G.* "Influence of nuclear quantum effects on the electronic properties of amorphous carbon", Proc. Natl. Acad. Sci. **2022**, 119, 31, e2203083119. (co-corresponding author), Highlighted on PME News: "A new look at disordered carbon" by Sarah C. P. Williams.
- **6.** Francese, T., **Kundu, A.**, Gygi, F., Galli, G.* "Quantum simulations of thermally activated delayed fluorescence in an allorganic emitter" <u>Phys. Chem. Chem. Phys. **2022**</u>, 24, 10101.
- **7.** Yang, H., Govoni, M.*, **Kundu, A.**, Galli, G.* "Combined first-principles calculations of electron-electron and electron-phonon self energies in condensed systems" <u>J. Chem. Theory Comput. **2021**</u>, 17, 12, 7468.
- **8. Kundu, A.***, Govoni, M., Yang, H., Ceriotti, M., Gygi, F., Galli, G.* "Quantum vibronic effects on the electronic properties of solid and molecular carbon" Phys. Rev. Materials **2021**, 5, L070801. (co-corresponding author)
- **9. Kundu, A.**†, Sillar†, K., Sauer, J.* "Predicting adsorption selectivities from pure gas isotherms for gas mixtures in metalorganic frameworks" Chem. Sci. **2020**, 11, 643 (Featured in front cover).
- **10. Kundu, A.**, Sillar, K., Sauer, J.* "Ab initio prediction of adsorption isotherms for gas mixtures by grand canonical Monte Carlo on a lattice of sites" J. Phys. Chem. Lett **2017**, 8, 2713.
- **11.** Sillar, K.*, **Kundu, A.**, Sauer. J. "Ab initio adsorption isotherms for molecules with lateral interactions: CO₂ in metalorganic framework" J. Phys. Chem. C **2017**, 121, 12789.
- **12. Kundu, A.**, Piccini, G., Sillar, K., Sauer, J.* "Ab initio prediction of adsorption isotherms for small molecules in metalorganic frameworks" <u>J. Am. Chem. Soc. **2016**</u>, 138, 14047.
- **13**. Pal, A.*, Saha, S., Maji, S. K., Sahoo, R., Kundu, M., **Kundu, A.** "Galvanic replacement of As(0) nanoparticles by Au(III) for nanogold fabrication and SERS application", New J. Chem. **2014**, 38, 1675.
- **14.** Pal, A.*, Saha, S., Maji S. K., Kundu, M., **Kundu, A.** "Wet chemical synthesis of spherical arsenic nanoparticles by a simple reduction method and its characterization", Adv Mat. Lett. **2012**, 3, 177.

AWARDS AND ACADEMIC ACHIEVEMENTS

•	Fischer-Nernst award 2018 for best Ph.D. thesis, Department of Chemistry, HU Berlin	Nov 2019
•	Successfully defended funding (2 \times 2 years) from German Science Foundation.	Nov 2013 – Oct 2017
•	IMPRS Ph.D. scholarship, FHI Berlin	Nov 2012 - Oct 2013
•	Shyama Prasad Mukherjee Fellowship (SPMF), CSIR: Govt. of India.	Sep 2011
•	The proficiency medal for best M.Sc. project work, IIT Kanpur.	May 2011
•	The general proficiency medal for best academic performance in M.Sc, IIT Kanpur.	May 2011
•	All India 1st rank in "National Eligibility Test (NET)" examination conducted by CSIR, Govt. of	India. Dec 2010
•	All India 11 th rank in "Joint Admission test for M.Sc (JAM)" examination conducted by IITs.	May 2009

TEACHING/ MENTORING EXPERIENCE

University of Chicago, Pritzker School of Molecular Engineering (PME)

Chicago, IL, USA

Mentoring 1 master's and 2 grad students

December 2022 - Present

• Mentored an undergraduate researcher

June 2020 - Oct 2021

Humboldt-Universität zu Berlin

Berlin, Germany

• Taught physical Chemistry 2: a practical course for master's students.

Summer 2015, summer 2016

Taught Molecular Modelling (MoMo): a practical course for bachelor's students.

Summer 2013

Supervised a visiting master's student researcher.

Summer 2013

CONFERENCES/WORKSHOPS

Invited and contributed talks

- **Kundu, A.**, Galli, G. "Quantum nuclear vibrations and electronic properties of molecular crystals" APS March Meeting 2023, Mar 5-10, 2023, Las Vegas, NV, USA
- (invited) Kundu, A. (i)(talk) "Influence of electron-phonon interactions on the electronic properties of materials" &
 (ii) Hand-on tutorial on i-PI---Qbox coupling for computing electron-phonon renormalization of materials. NOMAD-ECAM Workshop: Modelling Materials at Realistic time Scales via Optimal Exploitation of Exascale Computers and Artificial Intelligence. Jul 25 -29, 2022, Berlin, Germany.
- **Kundu, A.**, Song, Y., Galli, G. "Revisiting the electronic properties of diamond like amorphous carbon" APS March Meeting 2022, Mar 14-18, 2022, Chicago, IL, USA
- **Kundu, A.**, Govoni, M., Ceriotti, M., Gygi, F., Galli, G. "Finite temperature electronic properties of diamond-like carbon materials" APS March Meeting. Mar 15-19, 2021, Virtual.
- **Kundu, A.**, Govoni, M., Ceriotti, M., Gygi, F., Galli, G. "Finite temperature electronic properties of diamond and diamondoids" APS March Meeting. Mar 2-6, 2020, Denver, CO, USA (Cancelled due to COVID-19).
- Kundu, A, Sillar, K., Sauer, "Ab initio prediction of pure and mixture gas adsorption of small molecules in CPO-27-Mg" <u>Bunsentagung 2016: 115th General Assembly of German Bunsen Society for Physical Chemistry</u>. May 5-7, 2016, Rostock, Germany.
- (invited) Kundu, A, Sillar, K., Sauer, J. "Effects of lateral interactions on adsorption of gas mixtures: an ab initio lattice model study." <u>Doktorandenseminar (Seminar of Ph.D. students) on Adsorption</u>. Aug 25-26, 2015, Magdeburg, Germany.
- **Kundu, A**, Sillar, K., Sauer, J. "Ab initio prediction of adsorption isotherms of pure and gas mixtures in CPO-27-Mg" CECAM Workshop: From the Chemical Bond to the Chemical Plants. Aug 25-29, 2014, Bangalore, India.

Poster Presentations

- **Kundu, A.**, Sillar, K., Sauer, J. "Ab initio prediction of co-adsorption of gases by GCMC simulations on a lattice of sites." IMPRS workshop 2018: From Models to Reality. Feb 19 23, 2018, Tegernsee, Germany.
- **Kundu, A.**, Sillar, K., Sauer, J., "Adsorptive separations of small gas molecules using CPO-27-Mg: Insights from ab initio GCMC simulations". EUROMOF 2017. Oct 29 Nov 1, 2017, Delft, Netherlands.
- **Kundu, A.**, Sillar, K., Sauer, J. "Ab initio prediction of co-adsorption of gases by GCMC simulations on a lattice of sites." WATOC 2017. Aug 27 Sep 1, 2017, Munich, Germany.
- **Kundu, A.**, Sillar, K., Sauer, J. "Ab initio prediction of pure and mixture gas adsorption of small molecules in CPO-27-Mg" Future of Chemical Physics, J. Chem. Phys. (AIP). Aug 31 Sep 2, 2016, Oxford, U.K.
- Sillar, K., **Kundu, A.,** Sauer, J. "Ab initio adsorption thermodynamics of small gas molecules in Mg-MOF-74" <u>EUROMOF 2015</u>. Oct 11-14, Potsdam, Germany.
- **Kundu, A.**, Piccini, G., Sillar, K., Sauer J. "Ab initio adsorption thermodynamics of small gas molecules in Mg-MOF-74", <u>STC 2015</u>. Sep 20 24, 2015, Potsdam, Germany.

- **Kundu, A.**, Sillar, K., Sauer, J. "Ab initio prediction of adsorption isotherm of pure and mixture of gases in Mg-MOF-74" IMPRS workshop 2015: Micro to Macro. Feb 9 13, 2015, Tegernsee, Germany.
- **Kundu, A.**, Sillar, K., Sauer, J. "Ab initio prediction of adsorption isotherm of pure and mixture of gases in Mg-MOF-74." STC 2014. Sep 14 18, 2014, Vienna, Austria.
- Sillar, K., **Kundu, A.,** Sauer, J. "Pure gas and mixture adsorption of CO₂ and CH₄ in CPO-27-Mg" International MOF Symposium 2013, Sep 16 − 17, 2013, Dresden, Germany.

Workshops / Summer schools attended

- MICCoM workshop and Hands-on Tutorials 2022, Oct 13-14, Argonne National Laboratory, Lemont, Illinois, USA.
- European Summer School in Quantum Chemistry (ESQC) 2015, Sep 6 − 19, 2015, Sicily, Italy.
- CECAM Summer School on "Computational modelling for catalysis". Sep 1 5, 2014, Bangalore, India.
- International workshop on molecular modelling and simulation: natural science meets engineering, Mar 11 12,
 2013, Frankfurt/Main, Germany.

PROFESSIONAL SERVICES

Reviewer: Reviewed 10+ manuscripts for journals: *JACS, PNAS, NPJ Comput. Mat. Sci., Commun. Phys., J. Phys. Chem., Phys. Chem. Chem. Phys., Struc. Chem.,* and for the Czeck Science Foundation (GACR).

Guest Editor: Served as a guest associate editor in *Frontiers in Catalysis*.

Membership: American Physical Society (2019 – Present), International Max Planck Research School (IMPRS) at Max Planck Society (2012-2018).

Conference Organizer: Organized two triennial IMPRS workshop: (i) From micro to macro, 2015 and (ii) From models to reality, 2018

REFERENCES

1. Prof. Dr. Joachim Sauer

Humboldt-Universität zu Berlin, Institut für Chemie, Unter den Linden 6. 10117 Berlin. Email: <u>is@chemie.hu-berlin.de</u> Tel. +49 30 2093 7135

2. Prof. Giulia Galli

University of Chicago,
Pritzker School of Molecular Engineering,
5640 S. Ellis Ave., William Eckhardt Research Center
(ERC) 233, Chicago, IL, 60637.
Email: gagalli@uchicago.edu
Tel. +1 773 702 0515

3. Prof. François Gygi

University of California, Davis,
Department of Computer Science,
3013 Ghausi Hall (Engineering III), Davis, CA, 95616.
Email: fgygi@ucdavis.edu
Tel. +1 530 752 4042

4. Prof. Amalendu Chandra

Indian Institute of Technology Kanpur, Department of Chemistry, Kanpur, UP 208016, India Email: amalen@iitk.ac.in
Tel: +91 512 259 7241

5. Dr. Michele Ceriotti

Associate Professor, École polytechnique fédérale de Lausanne (EPFL), EPFL STI IMX COSMO, MXG 337 (Bâtiment MXG), Station 12, CH-1015 Lausanne. Email: michele.ceriotti@epfl.ch

Tel: +41 21 693 2939

6. Dr. Marco Govoni

Assistant Professor,
Department of Physics, Computer Science,
Mathematics.
University of Modena and Reggio Emilia.
Via Campi 213/A, Modena, 41125, Italy.

Email: mgovoni@unimore.it Tel: +39 05920 55312