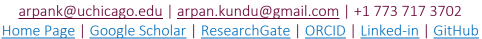
Staff Scientist, University of Chicago, Pritzker School of Molecular Engineering, 5640 S. Ellis Ave., Chicago, IL, 60637, USA





## **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 materials; (ii) spin-phonon interactions in solid-state spin defects, (iii) catalysis and gas conversion technologies.

## RESEARCH EXPERIENCE

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

Chicago, IL, USA

Staff Scientist, PI: Prof. Giulia Galli

May 2024 – Present

Post doctoral scholar, Adviser: Prof. Giulia Galli

Feb 2019 - Apr 2024

- Developed Molecular dynamics and stochastic methods, for electron-phonon interactions from first principles.
- 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 <a href="PyEPFD">PyEPFD</a> for performing electron-phonon calculations.
- Machine learning interatomic potentials and materials' properties.

#### 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 package: <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 MD 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 MD simulations.

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

Kharagpur, India

Summer research student

May 2010-July 2010

• Wet synthesis of As<sup>0</sup> and Au<sup>0</sup> nanoparticles (of tunable dimension) and their application in surface-enhanced Raman scattering (SERS).

#### **PUBLICATIONS**

# \* = Corresponding Author, † = Contributed equally

- **1.** R. Papadopoulos, † B. Masters, † **A. Kundu,** † N. Maldonado, A. Filatov, Y. Liu, T. Kim, G. Galli, A Wüttig.\* "Unlocking microscopic disorder in graphitic carbon with spectroelectrochemistry", *Submitted*, 2024. (co-first author)
- **2.** Berrens, M, **Kundu, A.,** Calegari-Andrade, M. F., Anh-Pham, T., Galli, G., Donadio, D.\* "Nuclear quantum effects on the electronic structure of water and ice", *J. Phys. Chem. Lett.* **2024**, *15*, 6818.
- **3. Kundu, A.\***, Galli, G.\* "Quantum vibronic effects on the excitation energies of the nitrogen-vacancy center in diamond", *J. Phys. Chem. Lett.* **2024**, 15, 802, Preprint: arXiv:2401.06745. (co-corresponding author)
- **4. Kundu**, **A.\***, Galli, G.\* "Quantum vibronic effects on the electronic properties of molecular crystals", <u>J. Chem. Theory</u> *Comput.* **2023**, 19, 13, 4011. (co-corresponding author).
- **5.** 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)
- **6.** Yang, H., Govoni, M.\*, **Kundu, A.**, Galli, G.\* "Computational protocol to evaluate electron-phonon interactions within density matrix perturbation theory", J. Chem. Theory Comput. **2022**, 18, 10, 6031.
- **7. Kundu, A.\***, Song, Y., Galli, G.\* "Influence of nuclear quantum effects on the electronic properties of amorphous carbon", <u>Proc. Natl. Acad. Sci. **2022**</u>, 119, 31, e2203083119. (co-corresponding author), Highlighted on PME News: "<u>A new look at disordered carbon</u>" by Sarah C. P. Williams.
- **8.** Francese, T., **Kundu, A.**, Gygi, F., Galli, G.\* "Quantum simulations of thermally activated delayed fluorescence in an allorganic emitter" *Phys. Chem. Chem. Phys.* **2022**, 24, 10101.
- **9.** 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.
- **10.** 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)
- **11. Kundu, A.**<sup>†</sup>, Sillar<sup>†</sup>, 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).
- **12. 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" <u>J. Phys. Chem. Lett **2017**, 8, 2713</u>.
- **13.** Sillar, K.\*, **Kundu, A.**, Sauer. J. "Ab initio adsorption isotherms for molecules with lateral interactions: CO<sub>2</sub> in metalorganic framework" J. Phys. Chem. C **2017**, 121, 12789.
- **14. Kundu, A.**, Piccini, G., Sillar, K., Sauer, J.\* "Ab initio prediction of adsorption isotherms for small molecules in metalorganic frameworks" *J. Am. Chem. Soc.* **2016**, 138, 14047.
- **15**. 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", <u>New J. Chem. **2014**, 38, 1675</u>.
- **16.** 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", <u>Adv Mat. Lett. **2012**</u>, 3, 177.

## AWARDS AND ACADEMIC ACHIEVEMENTS

Fischer-Nernst award 2018 for best Ph.D. thesis, Department of Chemistry, HU Berlin

Nov 2019 Nov 2013 – Oct 2017

• Successfully defended funding (2 × 2 years) from German Science Foundation.

• 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 1<sup>st</sup> rank in "National Eligibility Test (NET)" examination conducted by CSIR, Govt. of India. Dec 2010
 All India 11<sup>th</sup> 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 vibronic effects on the electronic properties of solid state spin defects" APS March Meeting 2024, Mar 3-8, 2023, Minneapolis, MN, USA.
- **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: 115<sup>th</sup> 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". <u>EUROMOF 2017</u>. 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." <u>STC 2014</u>. 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 15+ manuscripts for journals: *JACS, PNAS, NPJ Comput. Mat. Sci., Commun. Phys., J. Phys. Chem., Phys. Chem. Chem. Phys., Struc. Chem., Crystals, Molecules,* 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. Joachim Sauer

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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.

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Tel. +1 530 752 4042

#### 4. Dr. Michele Ceriotti

Associate Professor, École polytechnique fédérale de Lausanne (EPFL), EPFL STI IMX COSMO, MXG 337 (Bâtiment MXG), Station 12,

CH-1015 Lausanne.

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## 5. Dr. Marco Govoni

Assistant Professor,
Department of Physics, Computer Science,
Mathematics.

University of Modena and Reggio Emilia. Via Campi 213/A, Modena, 41125, Italy.

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Tel: +39 05920 55312

#### 6. Dr. GiovanniMaria Piccini

Associate Professor,
Theoretical & Computational Chemistry, Department of
Chemical & Geological Sciences,
University of Modena and Reggio Emilia,
Via Giuseppe Campi, 103, 41125 Modena MO, Italy.
Email: giovannimaria.piccini@unimore.it