

WORK EXPERIENCE

- 2022 – Assistant Professor, Tata Institute of Fundamental Research, Mumbai, India
2019 – 2022 Postdoctoral Research, Princeton University, Princeton, USA
2018 – 2019 Research Associate, University of Illinois at Urbana-Champaign, USA

EDUCATION

- 2012 – 2018 Ph. D. Chemistry, University of Illinois at Urbana-Champaign, USA G. P. A. : 3.91/4
2007 – 2012 Integrated Master's in Chemistry, Indian Institute of Technology, Kanpur G. P. A. : 9.30/10

PUBLICATIONS

INDEPENDENT WORK

(* indicates corresponding author)

- [22] A. Bose*, "Adaptive kink filtration: achieving asymptotic size-independence of path integral simulations utilizing the locality of interactions", *The Journal of Chemical Physics* **in press** (2025).
- [21] A. Bose*, "Incorporation of Empirical Gain and Loss Mechanisms in Open Quantum Systems through Path Integral Lindblad Dynamics", *The Journal of Physical Chemistry Letters* **15**, 3363–3368 (2024).
- [20] D. Sharma and A. Bose*, "Impact of Loss Mechanisms on Linear Spectra of Excitonic and Polaritonic Aggregates", *Journal of Chemical Theory and Computation* **20**, 9522–9532 (2024).
- [19] A. Bose*, "Quantum correlation functions through tensor network path integral", *The Journal of Chemical Physics* **159**, 214110 (2023).
- [18] A. Bose*, "QuantumDynamics.jl: A modular approach to simulations of dynamics of open quantum systems", *The Journal of Chemical Physics* **158**, 204113 (2023).
- [17] A. Bose* and P. L. Walters*, "Impact of Solvent on State-to-State Population Transport in Multistate Systems Using Coherences", *Journal of Chemical Theory and Computation* **19**, 4828–4836 (2023).
- [16] A. Bose* and P. L. Walters*, "Impact of Spatial Inhomogeneity on Excitation Energy Transport in the Fenna–Matthews–Olson Complex", *The Journal of Physical Chemistry B* **127**, 7663–7673 (2023).
- [15] A. Bose*, "Pairwise connected tensor network representation of path integrals", *Physical Review B* **105**, 024309 (2022).
- [14] A. Bose*, "Zero-cost corrections to influence functional coefficients from bath response functions", *The Journal of Chemical Physics* **157**, 054107 (2022).
- [13] A. Bose* and P. L. Walters*, "A multisite decomposition of the tensor network path integrals", *The Journal of Chemical Physics* **156**, 024101 (2022).
- [12] A. Bose* and P. L. Walters*, "Effect of temperature gradient on quantum transport", *Physical Chemistry Chemical Physics* **24**, 22431 (2022).
- [11] A. Bose* and P. L. Walters*, "Tensor Network Path Integral Study of Dynamics in B850 LH2 Ring with Atomistically Derived Vibrations", *Journal of Chemical Theory and Computation* **18**, 4095–4108 (2022).

EARLIER WORK

(* indicates corresponding author)

- [10] **A. Bose** and N. Makri*, “Quantum-classical path integral evaluation of reaction rates with a near-equilibrium flux formulation”, *International Journal of Quantum Chemistry* **121**, 10.1002/qua.26618 (2021).
- [9] **A. Bose** and S. Torquato*, “Quantum phase transitions in long-range interacting hyperuniform spin chains in a transverse field”, *Physical Review B* **103**, 014118 (2021).
- [8] **A. Bose** and N. Makri*, “All-Mode Quantum–Classical Path Integral Simulation of Bacteriochlorophyll Dimer Exciton-Vibration Dynamics”, *The Journal of Physical Chemistry B* **124**, 5028–5038 (2020).
- [7] **A. Bose** and N. Makri*, “Coherent State-Based Path Integral Methodology for Computing the Wigner Phase Space Distribution”, *The Journal of Physical Chemistry A* **123**, 4284–4294 (2019).
- [6] **A. Bose** and N. Makri*, “Quasiclassical Correlation Functions from the Wigner Density Using the Stability Matrix”, *Journal of Chemical Information and Modeling* **59**, 2165–2174 (2019).
- [5] **A. Bose** and N. Makri*, “Wigner Distribution by Adiabatic Switching in Normal Mode or Cartesian Coordinates and Molecular Applications”, *Journal of Chemical Theory and Computation* **14**, 5446–5458 (2018).
- [4] **A. Bose** and N. Makri*, “Non-equilibrium reactive flux: A unified framework for slow and fast reaction kinetics”, *The Journal of Chemical Physics* **147**, 152723 (2017).
- [3] **A. Bose** and D. Goswami, “Investigating the science of few-cycle pulses on simple model systems”, in *Advances in Laser Physics and Technology*, edited by M. Mohan, A. K. Maini, A. B. Bhattacharjee, and A. K. Razdan (Cambridge University Press, New Delhi, India, 2015), pp. 37–52.
- [2] **A. Bose** and N. Makri*, “Wigner phase space distribution via classical adiabatic switching”, *The Journal of Chemical Physics* **143**, 114114 (2015).
- [1] **A. Bose** and D. Goswami, “Insignificance of Relative Time Delay between Photons for a Ultrafast Two-Photon Process”, in *2012 International Conference on Fiber Optics and Photonics (PHOTONICS)* (2012), pp. 1–3.

PROFESSIONAL SERVICES

- Reviewer in the Journal of Chemical Physics, the Journal of Chemical Theory and Computation, and the Journal of Physical Chemistry

CONFERENCE TALKS AND POSTERS

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| February 2025 | Talk on “Path Integral Methods for Spectra and Dynamics of Open Quantum Systems: Two Short Stories” at the Spectroscopy and Dynamics of Molecules and Clusters conference |
| January 2025 | Talk on “Exciton Transport in Open Quantum Systems” at the Optics Within Life Sciences conference |
| July 2024 | Talk and poster on “Exciton Transport in Open Quantum System: A Path Integral Perspective” at the Molecular Interactions and Dynamics Gordon Research Conference |
| June 2024 | Talk on “Quantum Transport in Biology: a Path Integral Perspective” at the Physics Colloquium of the Institute of Mathematical Sciences |
| February 2024 | Talk on “Excitation Energy Transfer: Quantum Transport in Biological Systems” at the TIFR NSF Colloquium series |
| October 2023 | Poster on “Exciton Transfer: Path Integral Approaches” in the inaugural Physical Chemistry Symposium organized by the Society of Physical Chemistry |
| August 2023 | Talk on “Dynamics of Extended Open Quantum Systems — Tensor Network Path Integral Approaches” at Harish-Chandra Research Institute as a part of their Physics Colloquium series |

- July 2023** Talk on “Excitonic Dynamics: Path Integrals and Tensor Networks” at Kaleidoscope, 2023 at Udaipur
- September 2022** Talk on “Non-Equilibrium Quantum Dynamics: Tensor Network Path Integral Formalism” at QMat 2022 hosted by Indian Institute of Technology Kanpur
- Spring 2021** Talk on “Nuclear Quantum Effects in *ab initio* Water Dynamics” at American Physical Society’s March Meeting
- Summer 2019** Poster on “Classical and Path Integral Methods for Computing the Wigner Distribution” at American Conference of Theoretical Chemistry
- Summer 2019** Poster on “Quantum-Classical Path Integral: Harmonic Backreaction & Blip Decomposition” at American Conference of Theoretical Chemistry
- Summer 2017** Poster on “Quantum Mechanical Rate Calculations in Condensed Phase Reactions” at American Conference of Theoretical Chemistry
- Summer 2017** Talk on “Quantum Mechanical Rate Calculations in Condensed Phase Reactions” at Midwest Theoretical Chemistry Conference
- Spring 2016** Talk on “Wigner Phase Space Distribution via Classical Adiabatic Switching” at American Chemical Society’s Annual Meeting

TEACHING EXPERIENCE

- Spring 2025** Computational Science: A Hands-On Approach
- Fall 2024** Introduction to Thermodynamics and Statistical Mechanics
- Fall 2023** Quantum Mechanics I covering introduction to classical mechanics, basic postulates of quantum mechanics, linear vector spaces and conversions between position and momentum eigenbasis, 1D solvable problems like particle-in-a-box and harmonic oscillators, time-independent and time-dependent perturbation theory, etc.
- Spring 2023** Chemical Dynamics covering Time-dependent perturbation theory, Born-Oppenheimer approximation and non-Born-Oppenheimer processes, Gaussian wavepacket dynamics, quantum thermodynamics, trajectory surface-hopping, etc.

HONORS AND AWARDS

- 2018** Certificate of Graduate Specialization in Computational Science and Engineering, UIUC
- 2017** Center for Advanced Theory and Molecular Simulation Travel Award, Department of Chemistry, UIUC
- 2016** Eastman Travel Award, Department of Chemistry, UIUC
- 2015 – 2016** Harry G. Drickamer Fellowship, Department of Chemistry, UIUC
- 2014 – 2015** Robert Carr Fellowship, Department of Chemistry, UIUC
- 2013 – 2014** Walter Brown Fellowship, Department of Chemistry, UIUC
- 2007 – 2012** KVPY Fellowship, Department of Science and Technology, Government of India: Fellowship instituted to promote excellence in pure science
- 2012** IIT Kanpur, Best Master’s Thesis Dissertation
- 2008 – 2009** IIT Kanpur, Academic Excellence Award
- 2006** All India Rank 82 in the 5th National Cyber Olympiad
- 2004** Qualified for the Indian National Mathematical Olympiad with qualifying rank 18; among 3 students from 10th standard to qualify
- 2004** All India Rank 31 in Senior Mathematical Olympiad
- 2003** All India Rank 28 in Junior Mathematical Olympiad

RESEARCH EXPERIENCE

ASSISTANT PROFESSOR

TIFR

- Developed a method for making the scaling of path integral simulations asymptotically independent of the system size.²²
- Developed an approach for incorporating empirical loss mechanisms in numerically exact path integral simulations.^{21,20}
- Developed new tensor network algorithm for non-perturbatively simulating thermal correlation functions of open quantum systems.¹⁹
- Designed and developed an **open-source Julia package** for simulations of dynamics in quantum non-adiabatic systems.¹⁸
- Developed a rigorous technique for identification of dynamical pathways in quantum transport processes in complex systems.^{17,16}

POSTDOCTORAL RESEARCH

PRINCETON UNIVERSITY

- Explored quantum effects of transverse field on spin chains with long-range interactions that have disordered stealthy hyperuniform ground states using density matrix renormalization group (DMRG) with Prof. Salvatore Torquato.⁹ Showed the possibility of generating order from disorder through quantum fluctuations in these spin systems.

INDEPENDENT WORK

- Established a novel pairwise connected tensor network representation for path integrals. This work generalizes the ideas from the MPS representation to a more flexible custom tensor network that manifestly captures the structure of the Feynman-Vernon influence functional, further illustrating the deep fundamental ties between tensor networks and path integrals.¹⁵
- Derived a series representation for the coefficients that capture the influence of the solvent on to the system in terms of the Kubo transform of the so-called “bath response function.”¹⁴
- Developed a new multisite tensor network formalism for simulating extended quantum systems coupled to local dissipative environments like exciton transfer and charge transfer chains coupled with local vibrations. This MS-TNPI is a two-dimensional structure that, when evaluated yields the time propagated reduced density matrix of the entire extended system in the form of a matrix product state.¹³
- Explored the impact of thermal gradients on quantum transport in the Frenkel-Holstein model. Demonstrated for the first time, the possibilities of control using spatially inhomogeneous temperature profiles.¹²
- Studied the excitonic dynamics and absorption spectrum of a B850 ring using MS-TNPI accounting for the effects of atomistically derived protein and vibrational environments.¹¹

PH. D. RESEARCH

UIUC

- Derived an approach to approximating the thermal correlation functions for mixed quantum-classical methods. It is shown that this method can exactly calculate rates of reactions for systems in atomistic environments.¹⁰
- Simulated exciton transfer in a chlorophyll dimer with all local vibrations present. The effects of static disorder on the resultant dynamics were analyzed.⁸
- Formulated a new exact numerical method for calculating multidimensional Wigner distributions for thermalized operators.⁷
- Derived a method for calculating correlation functions using stability matrices under the quasiclassical approximation. A very simple yet accurate approximation which is applicable for multidimensional systems is proposed.⁶
- Applied the adiabatic switching based Wigner method to atomistic Hamiltonians in normal mode coordinates and Cartesian coordinates. Calculated quasiclassical correlation functions and explored the importance of Zero-Point Energy and quantization of the thermal density.⁵

- Constructed a nonequilibrium reactive flux based method for calculating rates of reactions. This method is shown to unify slow and fast reaction dynamics.⁴
- Described an efficient classical trajectory based method for calculating the quantum Wigner phase-space distribution.²

MASTER'S RESEARCH PROJECTS

IITK

- Investigated a simple model of two-photon processes using non-relativistic quantum electrodynamics.¹
- Modelled and studied the impact of the shape of the envelope of a few-cycles laser pulse on the nature of the breakdown of rotating wave approximation in the dynamics of two-level systems.³
- Learned ultrafast spectroscopy with Prof. Debabrata Goswami. Worked on the instrumentation of a 2D nonlinear optical spectroscope using an acousto-optical modulator pulse shaper with colinear pulses.
- Worked with Prof. Martin Gruebele on expressing PGK-FRET proteins for subsequent study of protein dynamics in presence of gold nanorods during the Summer, 2011.