

# AAMOD ATRE

In Pursuit | M. Sc. Quantum Science and Technology

## About Me

1. **INTERESTED** in understanding molecular and optical interactions & their dynamics across various timescales
2. **ENJOY** delving into the theoretical underpinnings of a problem and exploring the system computationally
3. **INTEND** to pursue the development of theoretical frameworks in atomic, molecular and optical sciences

## Education History

<b>Post-Graduate</b> -	<b>Technical University of Munich</b> M. Sc. Quantum Science and Technology	<b>Oct 2022 - Present</b>
<b>Undergraduate</b> GPA 8.65/10.0	<b>Birla Institute of Technology and Science, Pilani</b> B.E. Chemical Engg.   Minor : Physics	<b>Aug 2017 - Aug 2021</b>
<b>Higher Secondary</b> Percentage : 96.5 %	<b>FIITJEE Junior College, Kukatpally, Hyderabad</b> Grades XI & XII	<b>Apr 2015 - Aug 2017</b>
<b>Senior Secondary</b> GPA : 10.0/10.0	<b>Delhi Public School, Nacharam, Hyderabad</b> Grades IX & X	<b>Mar 2013 - Mar 2015</b>

## Work Experience

**Remote Research Assistantship** **Oct. 2021 - Oct. 2022**  
*Dr. Aaron Kelly | Max Planck Institute for the Structure and Dynamics of Matter, Hamburg*

Benchmarking a semiclassical mapping-based dynamics method (spin-PLDM) by modelling the interactions between an atomic subsystem and cavity-modified field modes.

**Remote Research Internship** **July 2021 - Sept. 2021**  
*Dr. Pengfei Huo | University of Rochester, New York*

Compared the performance of standard and spin-based partially linearized density matrix (PLDM) algorithms in calculating linear absorption spectra of a bi-exciton coupled dimer model.

**Undergraduate Thesis | Research Internship** **Jan. 2021 - June 2021**  
*Dr. Jeremy Richardson | ETH Zürich*

Gauged the efficacy of a spin-mapping-based semiclassical dynamics technique by studying the population and coherence dynamics of exciton relaxation in a 1D polymer chain. Formulated a spin-mapping for tight-binding polymer chains which conform to  $SU(2)$  symmetry.

**Research Internship** **May 2019 - July 2019**  
*Prof. Bibek Dash | CSIR Institute of Minerals and Materials Technology, India*

DFT-based computational designing of triazole-based molecular precursors for selective  $CO_2$  capture. Statistically determined the optimum functional-basis combination to model the  $CO_2$ -triazole interactions with DFT. Studied  $CO_2$  interactions with the aromatic building blocks to propose a new triazole moiety design.

## Technical Skills

Programming		Software Packages	Language
Python	MATLAB	LAMMPS	English (Native/Bilingual)
Fortran	Maple	Quantum Espresso	German (Elementary - A2)
C	Mathematica	Gaussian 09	French (Elementary)
			Marathi   Hindi (Native)

## Undergraduate Teaching Experience

**Teaching Assistant** **Aug. 2020 - Dec. 2020**  
*Dr. S. D. Manjare | Process Design Principles - I*

**Teaching Assistant** **Aug. 2020 - Dec. 2020**  
*Dr. Radhika Vathsan | Quantum Mechanics - II*

Standardized Tests		
GRE General Test : 335/340   Quant : 168/170   Verbal : 167/170   Analytical Writing : 5/6		Aug. 2021
TOEFL : 115/120   Reading : 30/30   Listening : 30/30   Speaking : 25/30   Writing : 29/30		Sept. 2021
Academic Interests		
<ol style="list-style-type: none"> <li>1. Dynamics of many-body, optical and open quantum systems.</li> <li>2. Development of techniques in theoretical and computational chemistry.</li> <li>3. Development of molecular and optical tools towards quantum technology.</li> </ol>		
Relevant Academic Credit Courses		
Quantum Mechanics I & II Statistical Mechanics Nonlinear Dynamics and Chaos Theory	Solid State Physics Atomic and Molecular Physics	Statistical and Molecular thermodynamics Transport Phenomenon
Featured Undergraduate Projects		
<b>Designing Lithium-based metal organic frameworks for hydrogen production</b>		Jan. 2020 - June 2021
<i>Design Project   Dr. Paramita Haldar</i>		
<i>Ab initio</i> computational study of graphene-based Li-MOFs, employing density functional theory implementation in Quantum Espresso for electronic structure calculations & reactive molecular dynamics within LAMMPS framework to study hydrogen evolution amongst proposed models.		
<b>Study of cavity QED formalism and modern quantum control techniques</b>		Aug. 2020 - Dec. 2020
<i>Study Project   Dr. Raghunath Ratabole</i>		
Literature survey of modern qubit implementations, entangled state preparation and quantum gate implementations in molecule-coupled cavity systems. Study of field quantization, cavity QED formalism and the applications of Jaynes-Cummings model.		
<b>Modelling kinetics of photo-catalytic reactions involved in waste-water treatment</b>		Aug. 2019 - Dec. 2019
<i>Study Project   Dr. Sharad Sontakke</i>		
Modelled $TiO_2$ -based photocatalytic degradation of phenol and extraction of $Cr$ and $Cu$ ions with MATLAB. Optimized $TiO_2$ catalyst concentration and reaction rates for varying contamination levels.		
<b>Study of metal-organic frameworks as tools for adsorptive <math>CO_2</math> capture</b>		Jan. 2019 - May. 2019
<i>Study Project   Dr. Richa Singhal</i>		
Literature review of thermodynamic and electronic properties of MOFs. Studied the methodologies and developments in the field of $CO_2$ capture, focusing on MOFs.		
<b>Other Academic Projects</b>		Aug. 2018 - Dec. 2020
<i>BITS Pilani</i>		
<ul style="list-style-type: none"> <li>· Simulation of Hamiltonian maps and transport in structured fluids</li> <li>· Fugacity coefficient estimation of pure <math>CO_2</math> and <math>H_2O</math> using Van der Waals and Soave-Redlich-Kwong EOS modelling</li> <li>· Derivation of various thermodynamic relations for a grand canonical ensemble of <math>NO_2</math></li> </ul>		
Extracurricular Activities		
<b>Quantum Computing</b>	· Cleared Quantum Open Source Foundation (QOSF) cohort 4 assesment task	Sept. 2021
	· Completed IBM Quantum's Global Summer School	June 2020, July 2022
<b>Positions of Responsibility</b>	· Core Member : <i>Kala - Fine Arts Club</i>	Mar. 2018 - May 2020
<b>Hobbies</b>	· Long distance running, swimming and squash sports	
	· Avid reader of fiction and non fiction literature	