

Manas Sharma

PHD (PHYSICS)

■ manas.sharma@uni-jena.de	1	₩ February 2nd, 1995	1	manas.bragitoff.com	Ç	github.com/manassharma07 in
linkedin.com/in/manassharma07	١	(b) 0000-0002-5346-62	280) 💆 @manassharma07	,	🗷 Manas-Sharma-5 🞓 Google
Sc	h	olar • @PhysWhiz		🗇 phys.whiz 🔗 bragii	tof	f.com

Research Interests

My passion lies in **developing codes and methods** for modeling various phenonmena and processes in materials science such as **light-matter interactions**, adsorbate-substrate interactions, and **thin film deposition**. During my Ph.D., I focused on improving and accelerating **quantum embedding techniques** for molecular and periodic systems. I implemented **density functional theory (DFT)-based embedding** methods and coupled them with **real-time time-dependent DFT** (RT-TDDFT) and **wavefunction theory** (WFT) methods. Notably, I calculated excitation energies of solvated molecules and adsorption energies of molecule-periodic systems using WFT-in-DFT, achieving high accuracy at a fraction of the computational cost. My work also extended to **molecule-in-periodic embedding** and **high harmonic generation** via RT-TDDFT.

In my postdoctoral research, I have diversified into machine-learned interatomic potentials (MLIPs), particularly graph neural network (GNN)-based potentials, to study chemical vapor deposition (CVD) or physical vapor deposition (PVD) of thin films on substrates. To explore deposition processes at the atomic scale, I use DFT calculations and nudged elastic band (NEB) methods to study reaction pathways and energy barriers, as well as ab initio molecular dynamics (AIMD) to study dynamic evolution. Using the data from these simulations, I develop MLIPs trained on high-fidelity data and also leverage foundation models to extend their applicability.

Beyond simulations, I have a keen interest in **developing computational tools** for researchers. I have built a performant DFT code as well as a neural network library from scratch, optimized for parallelization and GPU acceleration. Additionally, I have experience creating GUIs for materials modeling, visualization, and input file generation.

Outside research, I enjoy creating YouTube tutorials, web & Android apps, and computer software/libraries for researchers and students.

About me

I am a postdoctoral researcher in the Chemical Engineering department at the Indian Institute of Science (IISc), jointly advised by Prof. Ananth Govind Rajan and Prof. Sudeep Punnathanam.

Previously, I obtained my PhD (in Physics) from Friedrich Schiller University Jena (FSU) in Germany, with the highest distinction of summa cum laude. I was supervised by Prof. Dr. Marek Sierka during my PhD.

I possess strong verbal, presentation, and written communication skills as demonstrated by extensive **participation in >17 conferences** (**7 talks**; **10 posters**) as well as publishing **7 scientific articles**.

Prior to joining FSU, I obtained a Master's in Physics from the University of Delhi (India), where I worked with <u>Dr. Debabrata Mishra</u> and published three papers. I obtained my Bachelor's in Physics (Hons) from the University of Delhi as well.

I am originally from a small, but beautiful town Haldwani, a few minutes from Nainital in India.

Awards and Achievements

	Top Viewed Article , Paper investigating high harmonic generation in molecular thin films was one of the			
2025	most viewed article among the works published in Advanced Optical Materials between January 1 to	_		
	December 31 2023			
2024	Summa Cum Laude, Grade awarded for PhD Thesis	Germany		
2024	Travel and Accommodation Bursary, Awarded for presenting my work at Turbomole 2024 Conference	UK		
2024	3.5 Million Views and 19.9k Subscribers , on YouTube channel, where I share my knowledge on			
2024	computational materials science, physics and scientific computing	Online		
2022	Travel Bursary, Awarded for presenting a talk at ETSF 2022 Conference	Germany		
2021	Best Oral Presentation, Awarded for my talk at EMRS Fall 2021 Meeting	Germany		
2019	Highly Downloaded Paper , My paper on CrysX was among the top 10 percent of most downloaded papers			
	published in Journal of Applied Crystallography between January 2018 and December 2019	India		
2017	Rank #3, Ranked at the 3rd position in the Physics department of my college during M.Sc. Physics	India		
2016	DC Arora Scholarship , Awarded for being a meritorious student during B.Sc. Physics	India		
2016	Rank #2, Ranked at the 2nd position in the Physics department of my college during B.Sc. Physics	India		

August 5, 2025

Education

University of Delhi (Sri Venkateswara College)

New Delhi, India

Master of Science (MSc) Physics

July 2016 - July 2018 (2 years)

- · Completed a project on density functional theory and material modeling that resulted in two publications in 2019
- · Specialized in Nanoscience
- Ranked 3rd in the Physics department of my college

University of Delhi (Acharya Narendra Dev College)

New Delhi, India

B.Sc (Honours) Physics

July 2013 - July 2016 (3 years)

- Aggregate percentage: 87.08
- Ranked second in the Physics department of my college (Acharya Narendra Dev College)
- Earned the D.C. Arora meritorious Scholarship, Acharya Narendra Dev College (Apr 2016)
- Was an active member of the Physics society there and held various posts

Work Experience

Indian Institute of Science

Bengaluru, India

Postdoctoral Researcher

1st August 2024 - present

- I develop graph neural network-based machine learning interatomic potentials (MLIPs) for investigating thin film deposition and for materials design.
- Perform extensive first-principles (DFT, AIMD and NEB) calculations for reaction pathways and transition state barriers.
- Perform molecular dynamics simulations powered by foundation MLIPs for various properties such as radial distribution functions, density, and
- Served as a teaching assistant for a number of courses:
 - Quantum-Mechanical Modeling of Nanomaterials (CH 253): Conducted tutorials on Quantum ESPRESSO and graded assignments and student exams.
 - AICTE Faculty Development Program (FDP) titled "Application of Machine Learning for Solving Challenging Engineering Problems": Conducted hands-on tutorials on training ML models.
 - Currently serving as a teaching assistant for an NPTEL course on "Machine Learning for Core Engineering Disciplines" where I contribute by creating question papers, assignments, tutorials and reviewing course materials and lectures.

Friedrich Schiller University Jena

Jena, Germany

3rd June 2019 - 4th April 2024 (4 yrs 10

mos)

- Wissenschaftlicher Mitarbeiter (Scientific Employee)
- I developed computationally efficient ab-initio methods and codes to study the properties of hybrid systems. I mainly programmed in Fortran and Python for this role
- Furthermore, I served as a teaching assistant for a number of courses:
 - Modelling and Simulation (Curriculum included basics of quantum mechanics, overview of standard algorithms utilized in quantum chemistry codes such as geometry optimization, and so on.)
 - Basics of Stochastics and Experimental Design
 - Algorithms for Scientific Computing (Curriculum included: Python programming, introduction to machine learning, theory and applications of classical ML models such as SVM, decision trees, and so on.)

As a teaching assistant, I created exercise sheets and examination papers, conducted weekly tutorials, and graded answer sheets. Additionally, I also deveoped course materials for the course on machine learning and Python.

Phys Whiz (Self-Employed)

Content Creator and Founder

January 2016 - present

- · Founded Phys Whiz to create content like tutorials and lectures on physics, computational material science, and scientific computing
- Currently at ~ 18 k subscribers on <code>YouTube</code> and ~ 22 k followers on <code>Instagram</code>
- Acquired various practical skills through this endeavour like image and video editing, graphic designing, animations, basic social media marketing and management skills
- The YouTube channel has garnered 3.2 Million views (approximately)

BragitOff.com (Self-Employed)

Blogger and Founder

June 2014 - present

- Owner and author of <u>bragitoff.com</u>, an educational blog, especially for physics students with a focus on computational material science, numerical techniques, and scientific computing
- The blog gets decent traffic with > 30k monthly views
- Searching 'bragitoff.com' on Google Scholar gives 28 unique results that have cited various blog posts of mine

AUGUST 5, 2025 2

Publications

Here is a link to my Google Scholar

• Optical Gaps of Ionic Materials from GW/BSE-in-DFT and CC2-in-DFT

M. Sharma, and M. Sierka

J. Chem. Theo. Comput. 20, 21, 9592-9605 (2024)

• TURBOMOLE: Today and Tomorrow

Y. J. Franzke, C. Holzer, J. H. Andersen, T. Begušić, F. Bruder, S. Coriani, F. Della Sala, E. Fabiano, D. A. Fedotov, S. Fürst, S. Gillhuber, R. Grotjahn, M. Kaupp, M. Kehry, M. Krstić, F. Mack, S. Majumdar, B. D. Nguyen, S. M. Parker, F. Pauly, A. Pausch, E. Perlt, G. S. Phun, A. Rajabi, D. Rappoport, B. Samal, T. Schrader, **M. Sharma**, E. Tapavicza, R. S. Treß, V. Voora, A. Wodyński, J. M. Yu, B. Zerulla, F. Furche, C. Hättig, M. Sierka, D. P. Tew, and F. Weigend

J. Chem. Theo. Comput. 19, 20, 6859-6890 (2023)

Note: The author list for the above publication was determined using alphabetical order, with the exception of the first two authors and the last five corresponding authors.

• Resonance Effect in Brunel Harmonic Generation in Thin Film Organic Semiconductors

W. Li, A. Saleh, **M. Sharma**, C. Huenecke, M. Sierka, M. Neuhaus, L. Hedewig, B. Bergues, M. Alharbi, H. ALQahtani, A. M. Azeer, S. Graefe, M. F. Kling, A. F. Alharbi, and Z. Wang

Adv. Optical Mater. 2203070 (2023) [ON COVER] @

• Efficient Implementation of Density Functional Theory based Embedding for Molecular and Periodic Systems using Gaussian Basis Functions M. Sharma, and M. Sierka

J. Chem. Theo. Comput. 18, 11, 6892-6904 (2022) [ON SUPPLEMENTARY COVER]

Real-time time-dependent density functional theory using density fitting and the continuous fast multipole method
 C. Mueller, M. Sharma, and M. Sierka
 J. Comput. Chem. 41, 2573–2582 (2020)

• CrysX: crystallographic tools for the Android platform

M. Sharma, and D. Mishra

J. Appl. Cryst. **52**, 1449-1454 (**2019**) [ON COVER PAGE]

First-principles study of the structural and electronic properties of bulk ZnS and small ZnnSn nanoclusters in the framework of the DFT+U
method

M. Sharma, D. Mishra, and J. Kumar Phys. Rev. B 100, 045151 (2019)

• DFT+U study of small ZnO nanoclusters

M. Sharma, and D. Mishra

AIP Conference Proceedings 2142, 110025 (**2019**)

Review Activities

Served as a peer reviewer for the following journals published by the American Chemical Society:

- The Journal of Physical Chemistry A
- ACS Omega

Skills

Programming languages C, C++, C#, Python, FORTRAN, Java, shell scripting

DFT methods development PySCF, TURBOMOLE

Quantum Chemistry packages Quantum ESPRESSO, PySCF, TURBOMOLE, NWChem, Serenity, ORCA, VASP

Molecular Dynamics packages LAMMPS, ASE

Modeling and visualization VESTA, Avogadro, Jmol, VMD, CrysX-3D Viewer, BURAI, Py3Dmol, ASE

Machine/Deep Learning scikit-learn, TensorFlow, PyTorch, Nequip, CrysX-NN

App development Web development Wordpress, HTML

Data processing/wrangling pandas, numpy

Data visualization matplotlib, seaborn

Outreach YouTube videos, Blog posts, Instagram posts

Debugging and profiling ARM forge for FORTRAN, VS code for python

Miscellaneous Video editing, Molecular animations, Graphic designing, Photo editing, Git, MS Excel

Documentation Markdown, LaTeX/Overleaf, Jupyter notebooks, MS Word, MS Powerpoint

Soft Skills Time Management, Teamwork, Problem-solving, Documentation, Engaging Presentation

AUGUST 5, 2025 3

Social Media

LinkedIn https://www.linkedin.com/in/manassharma07

Blog https://bragitoff.com/

YouTube https://www.youtube.com/@PhysWhiz
Twitter https://twitter.com/manassharma07
GitHub https://github.com/manassharma07/

• https://www.instagram.com/phys.whiz

Instagram • https://www.instagram.com/ducktape07/

https://www.instagram.com/crysx_3d/

• https://www.facebook.com/ducktape07

Facebook • https://www.facebook.com/bragitoff/

https://www.facebook.com/physwhizforum/

Google Scholar https://scholar.google.com/citations?user=WYOEL94AAAAJ&hl=en

ORCID https://orcid.org/0000-0002-5346-6280

Researchgate https://www.researchgate.net/profile/Manas-Sharma-5

Conferences, Workshops and Seminars

IISc-Fujitsu Workshop on Quantum Computing, IISc Bengaluru India (<u>link</u>)	23-24 Jan 2025
AI/ML Workshop 2025, IISc Bengaluru India (<u>link</u>)	07-08 Jan 2025
TURBOMOLE Users Meet Developers Conference, Oxford, UK (<u>link</u>)	19 Sep 2024
Mastering DFT: A Hands-on Training Workshop, Janardan Singh Foundation (ONLINE)	27 Jul 2024
17th International Congress of Quantum Chemistry 2023 in Bratislava, Slovakia (<u>link</u>)	25 Jun-01 Jul 2023
VISTA Seminars 2022 (VIRTUAL (<u>link</u>)	30 Nov 2022
BIOVIA Conference 2022 (VIRTUAL (<u>link</u>)	11-13 Oct 2022
ICQNN 2022 Conference in Jena, Germany (<u>link</u>)	05-09 Sep 2022
Psi-K 2022 Conference at EPFL in Lasuanne, Switzerland (<u>link</u>)	22-25 Aug 2022
25th ETSF Workshop on Electronic Excitations 2022 in Leuven, Belgium (<u>link</u>)	13-17 Jun 2022
Turbomole Developers Seminar Series 2021-2022	21 Feb 2022
NOA Spring Meeting 2022 (ONLINE) (<u>link</u>)	21-23 Feb 2022
Computational Methods in Materials Science (CMMS 2021) (ONLINE) (<u>link</u>)	24-25 Sep 2021
57th Symposium on Theoretical Chemistry (STC 2021) (ONLINE) (<u>link</u>)	20-24 Sep 2021
EMRS Fall 2021 meeting (Online) PBEST ORAL PRESENTATION AWARD (link)	20-23 Sep 2021
The Materials and Molecular Modelling Hub++ Annual Conference 2021 (ONLINE) (link)	14-15 Sep 2021
DokDok Lite 2021 in Jena, Germany (<u>link</u>)	01-03 Sep 2021
	12-15 Jul 2021
	07-08 Jun 2021
International Workshop on Computer-Aided Materials Discovery (Weekly Online ZOOM Webinars) (<u>link</u>)	27 May-24 Jun 2021
NOA CRC 1375 Spring Meeting (ONLINE) (link)	15-17 Feb 2021
	11-20 Nov 2020
	17 Nov 2020
·	26-28 Oct 2020
	02-06 Mar 2020
	21 Jan 2020
	16-20 Sep 2019
ICABS 2019 in Bhiwani, India (<u>link</u>)	07-09 Feb 2019
6th ISIF 2017 in New Delhi, India (<u>link</u>)	13 Dec 2017
	Al/ML Workshop 2025, IISc Bengaluru India (link) TURBOMOLE Users Meet Developers Conference, Oxford, UK (link) Mastering DFT: A Hands-on Training Workshop, Janardan Singh Foundation (ONLINE) 17th International Congress of Quantum Chemistry 2023 in Bratislava, Slovakia (link) VISTA Seminars 2022 (VIRTUAL (link) BIOVIA Conference 2022 (VIRTUAL (link) ICQNN 2022 Conference in Jena, Germany (link) Psi-K 2022 Conference at EPFL in Lasuanne, Switzerland (link) 25th ETSF Workshop on Electronic Excitations 2022 in Leuven, Belgium (link) Turbomole Developers Seminar Series 2021-2022 NOA Spring Meeting 2022 (ONLINE) (link) Computational Methods in Materials Science (CMMS 2021) (ONLINE) (link) 57th Symposium on Theoretical Chemistry (STC 2021) (ONLINE) (link) EMRS Fall 2021 meeting (Online) *BEST ORAL PRESENTATION AWARD** (link) The Materials and Molecular Modelling Hub++ Annual Conference 2021 (ONLINE) (link) DokDok Lite 2021 in Jena, Germany (link) International Workshop on Recent Developments in Electronic Structure (ES21) (ONLINE) (link) eSSENCE-eMMC eMeeting MMMM 2021 (Online) (link) International Workshop on Computer-Aided Materials Discovery (Weekly Online ZOOM Webinars) (link) NOA CRC 1375 Spring Meeting (ONLINE) (link) Intel® Software Development Tools for HPC (Webinar) (link) NOA Fall PhD School (ONLINE) (link) NOA Spring School and PI Workshop in Jena, Germany (link) NOA Spring School and PI Workshop in Jena, Germany (link) NOA Seminar by Dr. Heiko Appel (MPI Hamburg) in Jena, Germany (link) ICABS 2019 in Bhiwani, India (link)

Code Development.

Contributions

TURBOMOLE DFT based embedding coupled with WFT and RT-TDDFT methods within the RIPER module of the popular TURBOMOLE package

www.turbomole.org/

Independently Developed Libraries and Android/PC/Web Applications

AUGUST 5, 2025 4

CrysX-NN	An efficient neural network library from scratch that supports parallelization and GPUs	₽ GitHub link
CrysX-3D Viewer	A molecule and crystal viewer that renders high quality visualiza- tions using complex shaders developed using Unity gaming en- gine. Avaliable on Android, Windows, Mac and Linux	₽ Project Home page
CrysX-AR	An Android app for augmented reality visualization of molecules and crystals	₽Google Play Store link
CrysX-Crystallographic Tools	A set of crystallographic tools (XRD pattern simulation, CIF creator/parser, equation of state fitting, etc.) available as an Android	₿Google Play Store link
CrysX-Demo DFT based Embedding	app Online demo of frozen density embedding and projection based embedding	₽Web App link
RIPER-Tools	An online set of tools that allows to create input files for the RIPER module of TURBOMOLE from MaterialsProject or PubChem database	₽Web App link
VASP-GUI	An online GUI for VASP input preparation and parsing/visualizing output files	₽ Web App link
CrysX-CompChem File Converter	Web app that allows inter-conversion between various chemical file formats in current use	₽Web App link
Basis Set Converter	Web app to inter-convert between various basis set formats, powered by Basis Set Exchange	₽Web App link
The Math App	A suite of mathematical tools that has the potential to act as a substitute for Computer softwares like Matlab/Scilab on Android devices	₿ Google Play Store link
ML: Microstructure Classification Demo	A web demo app of a neural network (crysx_nn) model trained to classify microstutures	₽Web App link
ML: MNIST_Plus Digit Classification Demo	A web app that classifies user given handwritten digits using a convolutional network model (PyTorch) trained on a modified MNIST dataset	₽ Web App link

Languages _____

English Bilingual proficiencyHindi Native proficiencyGerman Elementary proficiency

AUGUST 5, 2025