

CONTACT INFORMATION	216A, Old R.A Hostel, IIT Kanpur Kanpur, UP 208016, India	+91-8090686789 suhasnahas@gmail.com
RESEARCH INTERESTS	<ul style="list-style-type: none"><li>• Dirac materials: graphene, topological insulators, weyl semimetals and related topological phases of materials</li><li>• Group theory in condensed matter physics</li><li>• Electron correlations and magnetism in materials</li><li>• Algorithms for abinitio thermodynamics : cluster expansion, simulated annealing/quenching and evolutionary algorithms</li></ul>	
EDUCATION	<b>Indian Institute of Technology</b> , Kanpur. <ul style="list-style-type: none"><li>• M Tech, <i>Materials Science and Engineering</i>, May 2015. <b>GPA : 9.5/10</b></li></ul> <b>National Institute of Technology</b> , Calicut. <ul style="list-style-type: none"><li>• B Tech, <i>Mechanical Engineering</i>, May 2013. <b>GPA : 7.0/10</b></li></ul> <b>St. Antony's Public School</b> , Kottayam. <ul style="list-style-type: none"><li>• Std XII, <i>AISSCE</i>, March 2009. <b>Score : 90.0/100</b></li></ul> <b>Infant Jesus Higher Secondary School</b> , Kollam. <ul style="list-style-type: none"><li>• Std X, <i>ICSE</i>, March 2007. <b>Score : 90.7/100</b></li></ul>	
WORK EXPERIENCE	<b>09/2015 - present :</b> <ul style="list-style-type: none"><li>• Project Engineer, <i>Computational Materials Science Lab</i>, IIT Kanpur</li></ul>	
POSTGRADUATE RESEARCH	<ul style="list-style-type: none"><li>• <b>Dissertation:</b> A computational study on predicting the structure of amorphous semiconductors using evolutionary algorithms</li><li>• Electric field induced gap modification in ultrathin blue phosphorous</li><li>• Chemical functionalization of black phosphorene</li><li>• Polymorphism in phosphorene and arsenene using evolutionary algorithms</li></ul>	
UNDERGRADUATE RESEARCH	<ul style="list-style-type: none"><li>• <b>Dissertation:</b> Graphene reinforcement on metal and polymer matrices</li></ul>	
REFEREED JOURNAL PUBLICATIONS	<ul style="list-style-type: none"><li>• <b>Nahas, S.;</b> Ghosh, B.; Bhowmick, S.; Agarwal, A. "First-Principles cluster expansion study of functionalization of black phosphorene via fluorination and oxidation" – <i>Phys. Rev. B</i> (2016)</li><li>• Ghosh, B.; <b>Nahas, S.;</b> Bhowmick, S.; Agarwal, A. "Electric field induced gap modification in ultrathin blue phosphorus" – <i>Phys. Rev. B</i> (2015)</li></ul>	
SUBMITTED JOURNAL PUBLICATIONS	<ul style="list-style-type: none"><li>• <b>Nahas, S.;</b> Gaur, A.; Bhowmick, S. "First-Principles prediction of amorphous phases using evolutionary algorithms" – under review in <i>Journal of Chemical Physics</i> – <i>arXiv</i> (2016)</li></ul>	
PAPERS IN PREPARATION	<ul style="list-style-type: none"><li>• <b>Nahas, S.;</b> Bajaj, A.; Bhowmick, S. "New polymorphs of phosphorene and arsenene"</li></ul>	

RELEVANT  
COURSES

- Electronic structure of materials
- Computational materials science
- Nanostructures and nanomaterials
- Symmetry and tensor properties of materials
- Mathematics and computational methods
- Transport phenomena
- Thermodynamics of materials
- Structure and characterization of materials

RESEARCH  
SKILLS

- ***Electronic Structure Codes:***
  - ⇒ **VASP & Quantum Espresso:** Density functional theory, Hybrid functionals, Density functional perturbation theory (Phonons) and Abinitio molecular dynamics (CPMD & BOMD)
  - ⇒ **Wannier90:** Maximally localised wannier functions from DFT
  - ⇒ **BerkeleyGW:** Quasi-particle Energies based on GW approximation, Dielectric matrix and Optical Responses.
  - ⇒ **CASINO:** Variational monte-carlo, Diffusion monte-carlo
- ***First-Principles Thermodynamics Codes:***
  - ⇒ **Universal Structure Prediction Xtallography:** Evolutionary algorithms for predicting stable/metastable phases from *first principles* in all dimensions and transition pathways between different phases.
  - ⇒ **MIT Abinitio Phase Stability:** Cluster Expansion for determining the stability of alloys and defects from *first principles*
  - ⇒ **Eazy Monte Carlo Code:** Determining the entire phase diagram for varying compositions of alloys and defects from *first principles*
- ***Basic Computational Skills:***
  - ⇒ **Programming Languages:** Fortran, Python
  - ⇒ **Scripting Languages:** BASH
  - ⇒ **Numerical Computing Tools:** R, MATLAB
  - ⇒ **Typesetting Tools:** L<sup>A</sup>T<sub>E</sub>X, Microsoft Office
  - ⇒ **Plotting Tools:** Gnuplot, Matplotlib
  - ⇒ **Operating Systems:** Linux, Windows

INTERNATIONAL  
WORKSHOPS

- **S N Bose - JAIST School on Quantum Monte Carlo** Mar, 2015
  - ⇒ Hands-on training in CASINO code
  - ⇒ Talks on the theoretical developments of Quantum Monte Carlo
- **8<sup>th</sup> USPEX Workshop** Jan, 2015
  - ⇒ Hands-on training in Universal Structure Prediction Xtallography code
  - ⇒ Talks on the modern methods of crystal structure prediction

## REFERENCES

### **Dr. Somnath Bhowmick**

Assistant Professor

Phone: +91-512-259-7161

Dept. of Materials Science and Engineering E-mail:[bsomnath@iitk.ac.in](mailto:bsomnath@iitk.ac.in)

Indian Institute of Technology, Kanpur

### **Dr. Amit Agarwal**

Assistant Professor

Phone: +91-512-259-6981

Dept. of Physics

E-mail:[amitag@iitk.ac.in](mailto:amitag@iitk.ac.in)

Indian Institute of Technology, Kanpur

### **Dr. Anshu Gaur**

Assistant Professor

Phone: +91-512-259-7600

Dept. of Materials Science and Engineering

E-mail:[agaur@iitk.ac.in](mailto:agaur@iitk.ac.in)

Indian Institute of Technology, Kanpur