



# Guangchen Liu

PhD, Materials Engineering | Computational Materials Scientist | DFT · MD · ML · AI

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## Profiles

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[ResearchGate](#)

[ORCID](#)

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## Summary

Ph.D. candidate in Materials Engineering at Worcester Polytechnic Institute, specializing in **machine-learning-accelerated first-principles simulations** for advanced materials design. Research integrates DFT, molecular dynamics, and active learning to elucidate **structure–property relationships in complex oxides, high-entropy materials, and alloy systems**. Developer of multiple open-source frameworks, including *HEA-ML*, *Interface-Maker*, *LCO-Doper*, *HEP-Explorer*, and *Masgent* for data-driven discovery and design of novel materials.

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## Research Interests

- Machine-learning-accelerated atomic simulations for the discovery and design of complex materials.
  - Defect chemistry, doping, and ionic transport in perovskite oxides and high-entropy materials.
  - Computational frameworks for interface modeling and stability prediction.
  - Data-driven alloy design integrating thermodynamics and machine learning.
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## Education

**Worcester Polytechnic Institute, USA**  
Materials Engineering

2021 - Present  
Ph.D.

**Central South University, China**  
Materials Engineering

2018 - 2021  
M.S.

**Central South University, China**  
Powder Materials Science & Engineering

2014 - 2018  
B.S.

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## Research Experience

**IMPD Group | Advisor: Prof. Yu Zhong**  
Research Assistant

2021 - Present  
Worcester Polytechnic Institute, USA

Ph.D. research focuses on integrating first-principles calculations, molecular dynamics, and machine learning to accelerate the discovery of complex oxides, interfaces, and high-entropy materials.

- Developed **machine-learning-accelerated atomic simulation frameworks** integrating ML potentials and active learning to predict stability and conductivity with DFT-level accuracy.
- Developed an **AI agent for materials simulation**, automating structure handling, VASP input generation, workflow preparation & analysis, and rapid property prediction.
- Designed and deployed multiple **open-source frameworks** including *HEA-ML*, *Interface-Maker*, *LCO-Doper*, *HEP-Explorer*, and *Masgent*.
- Authored and co-authored **7 peer-reviewed papers** in *High Entropy Alloys & Materials*, *Materials Today Physics*, *Materials Today Communication*, *Journal of Alloys and Compounds*, etc.

**PPM Group | Advisor: Prof. Lijun Zhang**  
M.S. Researcher

2018 - 2021  
Central South University, China

M.S. work focused on coupling computational thermodynamics with machine learning to optimize aluminum-based casting alloys and understand composition–microstructure–property relationships.

- Developed **Alloy-Supermarket**, an open-source platform integrating Thermo-Calc and machine learning to design Sc-modified A356 Al-Si-Mg casting alloys.
- Authored and co-authored **5 peer-reviewed papers** in *Journal of Materials Science & Technology*, *Materials Today Communications*, *Journal of Materials Informatics*, etc.

<b>Selected Publications</b>	G Liu, S Yang, Y Zhong. Substitutional Effects at A- and B-Sites in High-Entropy ABO <sub>3</sub> Perovskites: Insights from Machine Learning-Accelerated Simulations and Active Learning Chemical Engineering Journal (Submitted)	2025
	G Liu, S Yang, Y Zhong. Unraveling Doping Effects in LaCoO <sub>3</sub> via Machine Learning-Accelerated Atomic Simulations and Active Learning Acta Materialia (Submitted)	2025
	G Liu, S Yang, Y Zhong. A Computational Framework for Interface Design Using Lattice Matching, Machine Learning Potentials, and Active Learning: A Case Study on LaCoO <sub>3</sub> /La <sub>2</sub> NiO <sub>4</sub> Materials Today Physics 🔗 <a href="https://doi.org/10.1016/j.mtphys.2025.101940">https://doi.org/10.1016/j.mtphys.2025.101940</a>	2025
	G Liu, S Yang, Y Zhong. High-Entropy Materials Design by Integrating the First-Principles Calculations and Machine Learning: A Case Study in the Al-Co-Cr-Fe-Ni System High Entropy Alloys & Materials 🔗 <a href="https://doi.org/10.1007/s44210-024-00041-3">https://doi.org/10.1007/s44210-024-00041-3</a>	2024
	W Yi, G Liu, Z Lu, J Gao, L Zhang. Efficient alloy design of Sr-modified A356 alloys driven by computational thermodynamics and machine learning Journal of Materials Science & Technology 🔗 <a href="https://doi.org/10.1016/j.jmst.2021.09.061">https://doi.org/10.1016/j.jmst.2021.09.061</a>	2022
	G Liu, J Gao, C Che, Z Lu, W Yi, L Zhang. Optimization of casting means and heat treatment routines for improving mechanical and corrosion resistance properties of A356-0.54 Sc casting alloy Materials Today Communications 🔗 <a href="https://doi.org/10.1016/j.mtcomm.2020.101227">https://doi.org/10.1016/j.mtcomm.2020.101227</a>	2020

<b>Selected Projects</b>	<b>Masgent</b> An AI agent for materials simulation that streamlines DFT workflows and analysis, fast machine-learning-potential (MLP) simulations, and lightweight ML modeling for materials science. 🔗 <a href="https://github.com/aguang5241/Masgent">https://github.com/aguang5241/Masgent</a>	2025
	<b>HEP-Explorer</b> An application for analyzing substitutional effects in high-entropy perovskites (HEPs), supporting compositional screening, property prediction, and performance optimization. 🔗 <a href="https://github.com/aguang5241/HEP-Explorer">https://github.com/aguang5241/HEP-Explorer</a>	2025
	<b>LCO-Doper</b> An application for analyzing dopant effects on LaCoO <sub>3</sub> (LCO), supporting composition optimization and material performance enhancement. 🔗 <a href="https://github.com/aguang5241/LCO-DOPER">https://github.com/aguang5241/LCO-DOPER</a>	2025
	<b>Interface-Maker</b> An application for generating customizable slabs and interfaces for first-principles simulations. 🔗 <a href="https://github.com/aguang5241/Interface-Maker">https://github.com/aguang5241/Interface-Maker</a>	2025
	<b>HEA-ML</b> An application for designing Al-Fe-Co-Cr-Ni high-entropy alloys by combining first-principles calculations and machine learning. 🔗 <a href="https://github.com/aguang5241/HEA-ML">https://github.com/aguang5241/HEA-ML</a>	2024
	<b>Alloy-Supermarket</b> An application that integrates computational thermodynamics and machine learning for Sc-modified Al-Si-Mg alloy design. 🔗 <a href="https://github.com/aguang5241/Alloy-Supermarket">https://github.com/aguang5241/Alloy-Supermarket</a>	2022

<b>Awards &amp; Honors</b>	<b>National Science Foundation Scholarship</b> CALPHAD	Jun 2023
	<b>Golden Egret Scholarship</b> Xiamen Golden Egret Special Alloy Co., LTD.	Dec 2020
	<b>Best Paper Award - Grand Prize</b> Hunan Education Department	Dec 2020
	<b>Best Poster Award – First Prize</b> Graduate School of Central South University	Dec 2020

<b>Teaching &amp; Mentoring</b>	<b>Major Qualifying Project (MQP)</b> Research Mentor	<b>Aug 2024 - Dec 2024</b> Worcester Polytechnic Institute, USA
	<b>ME 4875 - Introduction To Nanomaterials And Nanotechnology</b> Teaching Assistant	<b>Mar 2024 - May 2024</b> Worcester Polytechnic Institute, USA
	<b>ME 5385 - Metal Additive Manufacturing</b> Teaching Assistant	<b>Jan 2024 - Mar 2024</b> Worcester Polytechnic Institute, USA
	<b>MTE 594-D01 - ST: Applied Machine Learning</b> Teaching Assistant	<b>Mar 2022 - May 2022</b> Worcester Polytechnic Institue, USA

<b>Professional Activities</b>	<ul style="list-style-type: none"> <li><b>Peer Reviewer</b> for <i>npj Computational Materials</i>, <i>Scientific Reports</i>, <i>Computational Condensed Matter</i>, <i>The Journal of Physical Chemistry</i>, and other international journals in computational materials science.</li> <li><b>Member</b>, <i>The Minerals, Metals &amp; Materials Society (TMS)</i>.</li> <li><b>Conference Organizer</b>, edited the Technical Program and coordinated meeting logistics and activities for the international <i>CALPHAD 2023</i> conference.</li> <li><b>Conference Presentations and Posters</b> <ul style="list-style-type: none"> <li>Oral Presentation   <i>MS&amp;T 2025</i></li> <li>Poster Presentation   <i>WPI Graduate Research Innovation Exchange (GRIE) 2025</i></li> <li>Oral Presentation   <i>CALPHAD 2023</i></li> <li>Oral Presentation   <i>3rd World Congress on High-Entropy Alloys 2023</i></li> <li>Oral Presentation   <i>MS&amp;T 2022</i></li> <li>Poster Presentation   <i>Gordon Conference 2022</i></li> </ul> </li> </ul>
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<b>Technical Skills</b>	<b>Comupational Modeling &amp; Simulation</b> DFT (VASP), ASE, Pymatgen, Machine-Learning Potentials, HPC & Workflow Automation
	<b>AI &amp; Machine Learning for Materials</b> Active Learning, Bayesian Optimization, AI Agent Development, Neural Network Potentials
	<b>Materials Informatics &amp; Thermodynamics</b> Thermo-Calc, PyCalphad, CALPHAD
	<b>Programming &amp; Automation</b> Python, C#, C++, Linux, Markdown, Git, API Development
	<b>Visualization &amp; Analysis</b> Matplotlib, Seaborn, Plotly, VESTA, Origin, Pandas, Numpy
	<b>Tools &amp; Platforms</b> VS Code, JupyterLab, Github, PyQt5, Streamlit, Slurm, MS Office

<b>Languages</b>	<b>Mandarin</b> Native	<b>English</b> Fluent
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