

# Bryant Y. Li

<https://linkedin.com/in/bryant-li-0793591a9>, Google Scholar

Email : [bryantli@lbl.gov](mailto:bryantli@lbl.gov) and [bryant.li@berkeley.edu](mailto:bryant.li@berkeley.edu)

Mobile : 2018513233

## EDUCATION

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- **The University of California, Berkeley (UCB)** Berkeley, California  
*Ph.D. & M.S. - Materials Science and Engineering; Applied Data Science GPA: 3.8* *Aug. 2021 – May. 2026*
- **The Pennsylvania State University (PSU)** University Park, Pennsylvania  
*B.S. - Materials Science and Engineering; Minor in Applied Mathematics; GPA: 3.89* *Aug. 2018 – May. 2021*

## EXPERIENCE

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- **Lawrence Berkeley National Laboratory / University of California, Berkeley** Berkeley, California  
*Computational Materials Scientist / Graduate Student Researcher* *Aug 2021 - Present*
  - **High-throughput Workflow Developer:** Developed and optimized high-throughput DFT and MD workflows on High-Performance Computing (HPC) platforms, engineering scalable data pipelines with MongoDB for large-scale materials informatics and AI/ML applications.
  - **Machine Learning Interatomic Potential Specialists:** Engineered and validated machine learning interatomic potentials (MLIPs) using first-principles data and active learning, optimizing model accuracy and efficiency for simulating complex phenomena like interfacial reactivity and correlated transport in battery systems.
  - **Data Engineering and Scientific Methodologies:** Applied advanced data science techniques and Python libraries to analyze large-scale simulation data (up to 1M atoms), deriving scientific insights by integrating DFT, MD, Neural Networks, and Monte Carlo methods with experimental validation.
- **PSU Materials Research Institute, Millennium Science Complex** University Park, Pennsylvania  
*Computational Materials Researcher* *Oct 2019 - May 2021*
  - **Multiscale Simulation Engineer:** Conducted multiscale simulations for a U.S. DOE-funded project on fuel cell electrocatalysts, performing thousands of surface DFT calculations to investigate nanoparticle surface phenomena
  - **Data Management:** Developed Python scripts to automate data management and analysis pipelines for Cluster Expansion (CE) and Monte Carlo simulations, designed for research efficiency.

## SKILLS

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- **High Throughput Workflow Stack:** VASP, LAMMPS, atomate2, fireworks, pymatgen, ASE, jobflow, custodian, mp-api, Quantum ESPRESSO, ICET
- **Python/Data Science Stack:** Pandas, NumPy, SciPy, Matplotlib, scikit-learn, TensorFlow, PyTorch, MongoDB
- **MLIP Stack:** python-ACE, GRACE, MACE, NequIP, QET, M3GNET, CHGNet, DeepMD-kit
- **Software:** Git, Anaconda, Docker, Jupyter Notebooks, Visual Studio Code, MATLAB, Ovito, VESTA, Robo3T, SLURM

## SELECTED PUBLICATIONS

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- Li, B., et al. (2025). *An Atomistic Study of Reactivity in Solid-State Electrolyte Interphase....* **The Journal of Physical Chemistry C**: Led the development of a DFT and MLIP computational framework to successfully predict thermodynamically stable products and reaction dynamics at the Li/Li<sub>7</sub>P<sub>3</sub>S<sub>11</sub> solid-state battery interface.
- Ganose, A.M., ..., Li, B., ..., et al. (2025). *Atomate2: Modular Workflows for Materials Science. Digital Discovery*: Contributed to the development of atomate2, a next-generation high-throughput computational framework designed to accelerate materials screening and the generation of large-scale property databases for materials informatics.
- Gallant, M.C., ..., Li, B., et al. (2025). *A Cellular Automaton Simulation for Predicting Phase Evolution....* **Chemistry of Materials**: Co-developed a novel cellular automaton simulation to predict phase evolution in solid-state reactions, validating the model against experimental synthesis of complex oxides to optimize reaction conditions.
- Goff, J., Li, B., et al. (2021). *Quantifying Multipoint Ordering in Alloys. Physical Review B*: Introduced a method using cluster order parameters to quantify complex chemical ordering in multicomponent alloys, explaining its critical impact on the stability of Pt/Pd nanoparticle electrocatalysts.

## LEADERSHIP & PROJECT MANAGEMENT

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- **Program Management:** Served as lead project manager for the Persson group's DOE Advanced Battery Materials Research (BMR) program, responsible for directing research progress, authoring quarterly stakeholder reports, and managing NRL High-Performance Computing resources.
- **Consortium Representation:** Represented the research group in the multi-institution DOE Energy Storage Research Alliance (ESRA), aligning group strategy with consortium goals for AI/ML applications in sodium and zinc battery systems.
- **Technical Leadership:** Directed the Machine Learning subgroup, leading weekly meetings to coordinate collaborative code development, architect shared computational workflows, and ensure communication across multiple battery research projects.