Guillermo Vazquez

Ph.D. Candidate \cdot Computational Material Scientist College Station, TX



□ vzguille.me

Summary

- 8+ years of experience on Python and computational science most popular modules such as: scikit-learn, pandas, Tensorflow, etc. Dexterity in any programming language for computational science (Matlab, C++, javascript, R, FORTRAN, etc.).
- 7+ years of experience on Density Functional Theory (DFT).
- 5+ years of experience of Materials Design from materials simulations and data analytics.
- Fast growing author in academia with 150+ citations in 7 peer-reviewed (1 first author) journal publications during my PhD. Expected to publish 4 more (1 first author) during my last year.

Education

Texas A&M University

Ph.D. in Materials Science & Engineering

Aug. 2019 - Graduation Date: Aug 2024

College Station, TX

Autonomous University of Queretaro

B.S. Nanotechnology Engineering

Aug. 2013 – Aug 2018 Queretaro, Mexico

Experience

Texas A&M University Research Assistant

Aug. 2019 – May 2024 College Station, Texas

- Developed Density Functional Theory (DFT) automation methodology for elastic and stability analysis in alloy spaces.
- Data Analysis and Machine Learning applied to Materials Design.
- Using the CALPHAD method we calculate in a high-throughput manner thousands of data points. We then fit a regression model of custom architecture and use it as a kernel in optimization tasks for Material Design.
- High-Throughput generation of a DFT database for different configuration structures for an alloy system. Then a GNN regression model is trained for the application of otherwise commonly expensive DFT tasks.

Texas A&M University Teaching Assistant

Aug. 2022 – May 2024 College Station, Texas

• Fall 2022 Thermodynamics in Materials Science · Graduate level

- \bullet Spring 2023 and Spring 2024 Thermodynamics of Materials \cdot Undergraduate level
- Created a Thermo-Calc walk-through designed for undergrad engineering students.
- Created and graded challenging and comprehensive homework sets that helped the student approach thermodynamics from a Materials Science point of view.

Selected Publications

Efficient machine-learning model for fast assessment of elastic properties of high-entropy alloys.

Vazquez, G. et al., Singh, P., Sauceda, D., Couperthwaite, R., Britt, N., Youssef, K., Johnson, D.D. and Arróyave, R., 2022. Acta Materialia, 232, p.117924.

A deep neural network regressor for phase constitution estimation in the high entropy alloy system Al-Co-Cr-Fe-Mn-Nb-Ni. Vazquez, G., Chakravarty, S., Gurrola, R. and Arróyave, R., 2023. npj Computational Materials, 9(1), p.68.

Selected Presentations

Cluster Expansion Approximation Accelerated by a Graph Neural Network Regressor. Vazquez, G., Sauceda, D., and Arroyave, R., Orlando, Florida, Mar. 2024

Deep Neural Network Regressor for Phase Fraction Estimation on the High Entropy Alloy System Al-Co-Cr-Fe-Mn-Nb-Ni.Vazquez, G., Chakravarty, S., Gurrola, R. and Arroyave, R., Anaheim, California, Feb. 2022

Technical Skills

Programming Languages: Python, Matlab, C++, javascript, R, FORTRAN, SQL,

HTML/CSS

Framework/OSs: GitHub, Windows, Linux, MacOS

Scientific Software: VASP, QuantumEspresso, VESTA, Xcrysden, Thermo-Calc, Origin,

Gnuplot, Pandas, scikit-learn, pyTorch, TensorFlow, OpenCV

Languages: English, Spanish

Honors & Awards

Datathon 2023 Texas A&M University Major League Hacking DATATHON 2023, 2nd place in Computer Vision Challenge Oct. 2023

Outstanding Teacher Department of Materials Science & Engineering Awards for my outstanding work as a TA in the MSEN 210 Course. May 2023

D3EM Certificate Recipient Accepted for the Data-Enabled Discovery and Design of Energy Materials (D3EM) Certificate. **Jun. 2020**

Prof. Development & Certifications

SATA 2022 - School for Advanced thermodynamics Assessments	Jun. 2022
1st Online VASP Workshop: Introduction to Ab-initio Simulation	Nov. 2021
Computational Materials Science Summer School 2021	Jun. 2021
AFLOW Summer School on Computational Materials Science Across Scales	Jun. 2020