

Christian Venturella

Ph.D. Student - Theoretical Chemistry, Yale University, New Haven, CT

christian.venturella@yale.edu, (917) 767-5548
website - github - LinkedIn

INSTITUTIONS

Ph.D. Student, Theoretical Chemistry, Yale University

SEP 2022 — CURRENT

Bachelors in Chemistry, Princeton University

SEP 2017 — MAY 2021

Program in Applied and Computational Mathematics

Program in Applications of Computing

Program in Materials Science and Engineering

AWARDS AND HONORS

Undergraduate Awards

- summa cum laude (2021)
- American Institute of Chemists Student Award (2021)
- Sigma Xi Research Society (2021)
- Fulbright Semifinalist (2021)

RESEARCH AND WORK EXPERIENCE

Ph.D. Research / Theoretical Chemistry

AUG 2022 — CURRENT

The Zhu Group

Yale University, New Haven, CT

- Formulated machine learning method for predicting many-body quantum chemistry
- Constructed symmetry adapted intrinsic atomic orbital basis transformations for modeling quantum many body properties in a local orbital representation
- Generated and maintained a database of 10,000+ GW and coupled cluster simulations in massively parallel fashion for molecular and nanomaterial systems
- Engineered workflow for extracting photoemission spectrum, density matrix, energy, and excited states from imaginary-frequency self-energy ML predictions
- Developed graph attention network architectures for predicting many-body physics by targeting the self-energy over DFT orbital graphs. Achieved improved performance, heightened data efficiency, and improved generalizability over existing methods on similar benchmarks.

Software Engineering and Data Science / Educational Data Analytics

JUL 2021 — JUL 2022

LinkIt!

New York City, NY

- Engineered a data analysis system for generation of regular administrative reports in PDF and web formats from database views using SQL, pandas, plotly dash, and MikTex; outperformed a legacy Excel system by 100-fold
- Automated process of computing linear regressions on statewide educational metrics for over 500 school districts
- Designed and implemented a pipeline for estimating the probability a student will meet state education standards with calibrated support vector machines
- Created an interactive Dash website for visualizing and validating machine learning model performance on live educational data grouped by grade level and academic subject

Undergraduate Thesis Research / Computational Chemistry

SEP 2020 — APR 2021

The Scholes Group

Princeton University, Princeton, NJ

- Modeled catalyst empirical reaction efficiency as a function of simulation-obtained substrate binding free energies using GPU-accelerated molecular dynamics free energy calculations
- Computed visible spectrum of catalytic intermediate in agreement with experimental spectroscopy using Time Dependent Density Functional Theory
- Employed the weighted histogram analysis method to compute potentials of mean force from umbrella-sampled molecular dynamics trajectories

Summer Research Internship / Polymer Chemistry

JUN 2019 — AUG 2019

The Lau Group

University of Strathclyde, Glasgow, UK

- Modeled fluorescence time series data with Ward-Tordai equation for surface adsorption kinetics
- Implemented nonlinear regression and numerical integration algorithms for experimental data fitting to integro-differential equations
- Synthesized polymer nanosheets from peptide-like oligomers
- Developed and refined protocol to immobilize nanosheets on gold surfaces with thiol linkages for transmission electron microscopy

PUBLICATIONS

- Unified Deep Learning Framework for Many-Body Quantum Chemistry via Green's Functions** **AUG 2024**
Venturella, C.; Li, J.; Hillenbrand, C.; Peralta, X. L.; Liu, J.; Zhu, T. Unified Deep Learning Framework for Many-Body Quantum Chemistry via Green's Functions; 2024. arXiv:2407.20384
- Machine learning many-body Green's function for molecular excitation spectra** **OCT 2023**
C. Venturella, C. Hillenbrand, J. Li, and T. Zhu, "Machine Learning Many-Body Green's Functions for Molecular Excitation Spectra", J. Chem. Theory Comput. 2024, 20, 1, 143–154, arXiv:2310.09911
- Modeling Nonnatural Flavoenzyme Catalysis with Molecular Dynamics and Quantum Chemical Methods** **2021**
Princeton Undergraduate Thesis, Advisor(s): Gregory Scholes, Daniel Oblinsky

TEACHING EXPERIENCE

- Teaching Fellow** **SEP 2022 — Current**
Yale University Chemistry Department *New Haven, CT*
- General Chemistry I SEP 2022 — JUN 2023
 - Physical Chemistry I (Thermodynamics and Kinetics) SEP 2023 — CURRENT
- Undergraduate Teaching Assistant** **SEP 2020 — MAY 2021**
Princeton University Chemistry Department *Princeton, NJ*
- Organic Chemistry I SEP 2020 — DEC 2020
 - Organic Chemistry II JAN 2021 — MAY 2021
- Trenton Arts Program Volunteer** **SEP 2018 — MAY 2021**
Princeton University Music Department *Princeton, NJ*
- Private Music Instruction

SKILLS

- Computational Chemistry**
Python for chemical modeling and informatics (pyscf, geometric, ASE, rdkit), cp2k, Gaussian
- Data Science and Data Engineering**
Python for ML (scipy, sklearn, PyTorch, PyTorch Geometric, TensorFlow), python for data analytics and visualization (pandas, matplotlib, plotly, xlwings, dash, cvx, geopandas), SQL, MATLAB, R, Microsoft Excel
- Mathematical Modeling**
Kernel Ridge Regression, Gaussian Process Regression, Neural Networks, Geometric Deep Learning, Convex Optimization, Classifier Models, Model Calibration
- Word Processing**
L^AT_EX, HTML, Microsoft Word
- Languages**
English (Native), Spanish (Intermediate)