

Adrian Domínguez-Castro, PhD

Data Science, AI and Machine Learning

Henderson, NV, USA | adriandc1989@gmail.com | **GitHub:** <https://github.com/DrAdrianDC>

Linkedin Profile: <https://www.linkedin.com/in/adrian-dominguez-castro-44b51a221/>

SUMMARY

Machine Learning Engineer and Data Scientist with a PhD in Physics and 10 years of expertise in Quantum Chemistry, Data Science, AI and Machine Learning. Proficient in Python, SQL, and computational simulations, delivering data-driven solutions across science and technology sectors

SKILLS

• Machine Learning • Deep Learning • Data Science • Python • SQL • Statistical Analysis

WORK EXPERIENCE

Freelance Machine Learning Engineer and Data Scientist

Feb 2024-Present

Remote (Based in USA)

- Designed and implemented End-to-End Machine Learning and Data Science projects, showcasing them on GitHub (<https://github.com/DrAdrianDC/Portfolio-for-Data-Science>).

Postdoctoral Research Fellow

Feb 2023-Jan 2024

Vanderbilt University

Nashville, TN, USA

- Designed, developed and implemented a Machine Learning system that predicts physical-chemistry observables with more than 85% accuracy, resulting in 1 publication.
- Managed simulations and projects in Computational Chemistry and Machine Learning.

Doctoral Research Assistant

Oct 2018-Dec 2022

University of Bremen

Bremen, Germany

- Completed 3 multi-year research projects, resulting in 4 publications and improved data science solutions resulting in a 50% reduction in data acquisition and processing time.
- Led data modeling efforts to develop advanced simulations and predictive models for new nanoscience technologies, presenting insights at international scientific meetings. Built teams and forged collaborations across five countries (Germany, Spain, China, USA, France)

Graduate Research Fellow

Sept 2017-Jul 2018

University of Groningen

Groningen, The Netherlands

- Managed research on Theoretical Chemistry, and mentored junior team members. Optimized code for simulations and data science projects.

Visiting Research Scholar

Jan 2016-May 2016

Thompson Rivers University

Kamloops, Canada

- Developed computational modeling, and Data Science for Quantum Chemistry. Built teams and forged collaborations across two countries (Canada and Mexico), resulting in 1 publication.

Associate Junior Researcher

Sept 2015-Jul 2017

Higher Institute for Technology and Applied Sciences (InSTEC)

Havana, Cuba

- Led a team to meet project milestones, optimizing computational modeling and automating pre-processing to reduce time by 50%, while developing advanced predictive models for Computational Chemistry, resulting in 2 publications

Assistant Professor

Sept 2014-Jul 2015

Technological University of Havana (CUJAE)

Havana, Cuba

- Delivered Lectures and Tutorials of Calculus, Differential Equations, and Numerical Mathematics (1st and 2nd years Electrical Engineering)

EDUCATION

Ph.D. (Dr. rer. nat.)

University of Bremen

Dissertation: Theoretical investigation on light- driven ultrafast dynamics in nanoscale

(Grade: magna cum laude)

Oct 2018 - Dec 2022

Bremen, Germany

B.Sc. Radiochemistry

Higher Institute for Technology and Applied Sciences (InSTEC)

Dissertation: Essential amino acids interacting with Fumonisin B1: A Theoretical Approach

(Grade: summa cum laude)

Sept 2008 - Jun 2014

Havana, Cuba

CERTIFICATIONS

- Machine Learning by Stanford University & DeepLearning.AI on Coursera.
- AI for Everyone by Stanford University & DeepLearning.AI on Coursera.

PORTFOLIO DATA SCIENCE AND MACHINE LEARNING

Explore the projects in details at <https://github.com/DrAdrianDC/Portfolio-for-Data-Science>.

- **Project 0: Project on Data Cleaning**
A comprehensive process of cleaning and preparing raw data for analysis and modeling.
Tools & Techniques: Python, pandas, numpy, matplotlib, plotly, seaborn, scikit-learn
- **Project 1: Breast Cancer Classification**
Classification problem using Support Vector Machine (SVM) in biomedical research.
Tools & Techniques: Python, Machine Learning, SVM, scikit-learn, numpy, pandas, matplotlib, seaborn
- **Project 2: Wine Quality Prediction**
Predicting the quality of wines using physicochemical variables.
Tools & Techniques: Python, Machine Learning, XGBoost, Logistic regression, Random Forest, Neural networks, pandas, scikit-learn, tensorflow, keras, numpy, matplotlib, data visualization, Streamlit web app.
- **Project 3: WTI Oil Prices Anomaly Detection**
Anomaly Detection in West Texas Intermediate (WTI) Crude Oil Prices.
Tools & Techniques: Python, Deep Learning, yfinance, pandas, tensorflow, keras, numpy, matplotlib, data visualization, LSTM autoencoders
- **Project 4: Apple Stock Market Prediction**
Apple stock market prediction using Deep Learning.
Tools & Techniques: Python, yfinance, pandas, tensorflow, keras, numpy, matplotlib, data visualization, Deep Learning, LSTM
- **Project 5: Image Classification**
Image Classification using Transfer Learning.
Tools & Techniques: Python, Machine Learning, tensorflow, keras, seaborn, numpy, matplotlib, data visualization, Deep Learning, Transfer Learning,
- **Project 6: DFT Meets Machine Learning**
Predicting hydrogen adsorption energies on rocksalt complex oxides combining DFT calculations and Machine Learning.
Tools & Techniques: Python, Machine Learning, Deep Learning, Linear Regression, Random Forest, Neural networks, pandas, scikit-learn, tensorflow, keras, numpy, matplotlib
Published in Theoretical Chemistry Accounts journal <https://lnkd.in/ePBzVbsC>
- **Project 7: Clustering**
Clustering using the K-Means algorithm on the famous Iris dataset.
Tools & Techniques: Python, Machine Learning, K-Means, pandas, numpy, matplotlib, seaborn, scikit-learn

PUBLICATIONS

Adrian Dominguez-Castro. *DFT and machine learning for predicting hydrogen adsorption energies on rocksalt complex oxides.* **Theor Chem Acc** **143**, **50** (2024) doi:[10.1007/s00214-024-03124-x](https://doi.org/10.1007/s00214-024-03124-x)

Michael Wagstaffe, **Adrian Dominguez-Castro**, Lukas Wenthaus, Steffen Palutke, Dmytro Kutnyakhov, Michael Heber, Federico Pressacco, Siarhei Dziarzhyski, Helena Gleißner, Verena Kristin Gupta, Harald Redlin, Adriel Dominguez, Thomas Frauenheim, Angel Rubio, Andreas Stierle, and Heshmat Noei. *Photoinduced Dynamics at the Water/TiO₂ (101) Interface.* **Phys. Rev. Lett.** **130**, **108001** (2023) doi:[10.1103/PhysRevLett.130.108001](https://doi.org/10.1103/PhysRevLett.130.108001)

Adrian Domínguez-Castro and Thomas Frauenheim. *Impact of vibronic coupling effects on light-driven charge transfer in pyrene-functionalized middle and large-sized metalloid gold nanoclusters from Ehrenfest dynamics.* **Phys. Chem. Chem. Phys.**, **2021**, **23**, **17129–17133** doi:[10.1039/d1cp02890a](https://doi.org/10.1039/d1cp02890a)

Adrian Domínguez-Castro, Carlos R. Lien-Medrano, Khaoula Maghrebi, Sabri Messaoudi, Thomas Frauenheim and Arnaud Fihey. *Photoinduced charge-transfer in chromophore- labeled gold nanoclusters: quantum evidence of the critical role of ligands and vibronic couplings.* **Nanoscale**, **2021**, **13**, **6786–6797** doi:[10.1039/d1nr00213a](https://doi.org/10.1039/d1nr00213a)

Michael Wagstaffe, Lukas Wenthaus, **Adrian Dominguez-Castro**, Simon Chung, Guilherme Dalla Lana Semione, Steffen Palutke, Giuseppe Mercurio, Siarhei Dziarzhyski, Harald Redlin, Nicolai Klemke, Yudong Yang, Thomas Frauenheim, Adriel Dominguez, Franz Kärtner, Angel Rubio, Wilfried Wurth, Andreas Stierle, and Heshmat Noei. *Ultrafast Real-Time Dynamics of CO Oxidation over an Oxide Photocatalyst.* **ACS Catal.** **2020**, **10**, **13650–13658** doi:[10.1021/acscatal.0c04098](https://doi.org/10.1021/acscatal.0c04098)

Adrian Domínguez-Castro, Daríel Hernández, and Fernando Guzmán. *Insights into the interactions of biomolecules with small gold clusters: a theoretical study from a DFTB perspective.* **Theor. Chem. Acc.** (2017) **136**:84 doi:[10.1007/s00214-017-2118-7](https://doi.org/10.1007/s00214-017-2118-7)

Adrian Dominguez-Castro, Fernando Guzman, Yasser Novo-Fernandez. *Adsorption on a nanoporous organic polymer for clean energy applications: A multiscale modelling study using Density Functional Tight Binding Approach.* **Computational and Theoretical Chemistry** **1102** (2017) **30–37** doi:[10.1016/j.comptc.2016.12.037](https://doi.org/10.1016/j.comptc.2016.12.037)

R. Castañeda-Arriaga, **A. Domínguez-Castro**, J. Lee, J. R. Alvarez-Idaboy, N. Mora-Diez. *Chemical Repair of Protein Carbon-Centred Radicals: Long-Distance Dynamic Factors.* **Can. J. Chem.**, **2016**, **94** (12): **1119–1126** doi:[10.1139/cjc-2016-0230](https://doi.org/10.1139/cjc-2016-0230).