Dakota Folmsbee

CHEMISTRY GRADUATE STUDENT · UNIVERSITY OF PITTSBURGH

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Education

University of Pittsburgh

Ph.D. STUDENT IN PHYISCAL CHEMISTRY

Aug. 2016 - PRESENT

Pittsburgh, PA

Clarkson University

Aug. 2012 - May 2016

Potsdam NY

Skills

B.S. IN CHEMISTRY

Chemistry NMR, HPLC, Mass Spectrometry, FTIR, UV/Vis

Programming Python, Scikit-Learn, Tensorflow, Keras, PyTorch, Bash, MFX, C++, Julia

General GNU/Linux, VIM, Microsoft Office Suite

Professional Experience

Computational/Physical Chemistry Graduate Student

Jan. 2017 - PRESENT

University of Pittsburgh

- · Instituted fast property prediction models to aid a genetic algorithm in rapid material screening
- Developed a molecular machine learning representation for chemical property predictions

General Chemistry Teaching Assistant/Fellow

Aug. 2016 - Dec. 2017

University of Pittsburgh

- Recitation & Lab Honors General Chemistry
- · Recitation & Lab General Chemistry
- Lab General Chemistry for Engineers

Undergraduate Researcher

Aug. 2013 - May 2016

CLARKSON UNIVERSITY

- Synthesized carriers for cancer detecting molecules and chemotherapy drugs
- Analyzed compounds using techniques such as NMR, TOF-MS, and HPLC
- Researched procedures and applications for Gold nanoparticles and nanorods
- Synthesized Gold nanorods and analyzed with thermogravimetric analysis

General Chemistry Teaching Assistant/Mentor

Aug. 2013 - May. 2016

CLARKSON UNIVERSITY

- Recitation & Lab General Chemistry for Engineers
- Recitation & Lab General Chemistry for Chemistry and Chemical Engineering

Publications.

2020

D. Folmsbee and G. Hutchison. Assessing Conformer Energies using Electronic Structure and Machine Learning Methods. 2020. ChemRxiv. http://doi.org/10.26434/chemrxiv.11920914.v1

2019

D. Folmsbee, S. Upadhyay, A. Dumi, D. Hiener, & D. Mulvey. chemreps/chemreps: Molecular Machine Learning Representations (Version 0.1.1). 2019.Zenodo. http://doi.org/10.5281/zenodo.3333856

Presentation

ACS National Meeting & Expo, Online

Online

POSTER PRESENTATION March 2020

Assessing Conformer Energies: Machine Learning vs Conventional Quantum Chemistry

Covestro Lecture Series, University of Pittsburgh	Pittsburgh, PA
Poster Presentation	Jan. 2020
Assessing Conformer Energies: Machine Learning vs Conventional Quantum Chemistry	
Frederick Kaufman Memorial Lecture Series, University of Pittsburgh	Pittsburgh, PA
POSTER PRESENTATION	Oct. 2019
Assessing Conformer Energies: Machine Learning vs Conventional Quantum Chemistry	
Science 2019, University of Pittsburgh	Pittsburgh, PA
POSTER PRESENTATION Assessing Conformer Energies: Machine Learning vs Conventional Quantum Chemistry	Oct. 2019
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Advancing Research through Computing 2019, University of Pittsburgh POSTER PRESENTATION	Pittsburgh, PA
Rapid Predictive Methods to Aid in Screening of Organic Dielectric Materials	Mar. 2019
Science 2018, University of Pittsburgh	Pittsburgh, PA
POSTER PRESENTATION	Oct. 2018
Rapid Predictive Methods to Aid in Screening of Organic Dielectric Materials	
Frederick Kaufman Memorial Lecture Series, University of Pittsburgh	Pittsburgh, PA
Poster Presentation	Oct. 2018
Rapid Predictive Methods to Aid in Screening of Organic Dielectric Materials	
Covestro Lecture Series, University of Pittsburgh	Pittsburgh, PA
POSTER PRESENTATION	Sept. 2018
Rapid Predictive Methods to Aid in Screening of Organic Dielectric Materials	
Simulators Meeting 2018, Carnegie-Mellon University	Pittsburgh, PA
ORAL PRESENTATION Machine Learning to Aid in Screening for Organic Dielectric Materials	May 2018
Machine Learning to Aid in Screening for Organic Dielectric Materials	
Covestro Lecture Series, University of Pittsburgh Poster Presentation	Pittsburgh, PA Oct. 2017
Genetic Algorithms & Machine Learning for Rapid Materials Screening	OCI. 2017
Frederick Kaufman Memorial Lecture Series, University of Pittsburgh	Pittsburgh, PA
Poster Presentation	Oct. 2017
Genetic Algorithms & Machine Learning for Rapid Materials Screening	
Programming Projects	
chemreps	Aug. 2018 - PRESENT

DEVELOPER

- https://github.com/chemreps/chemreps
- Developed a molecular representation library for machine learning in chemistry.

QM/MM Study Group

July 2018 - Dec. 2018

INSTRUCTOR & ORGANIZER

- https://github.com/shivupa/QMMM_study_group
- Organized and taught various lessons surrounding computational chemistry.

Honors & Awards

- 2017 **Safford Teaching Award,** University of Pittsburgh
- 2017 **First Year Graduate Teaching Assistant Mentor**, University of Pittsburgh
- 2015 Walsh Fellow, Clarkson University