

Daniel Tabor

Texas A&M University
Department of Chemistry
College Station, TX 77843

Research Group Website: <https://dtaborgroup.com>
ORCID: [0000-0002-8680-6667](https://orcid.org/0000-0002-8680-6667)

Phone: (979)-862-8045
Email: daniel.tabor@chem.tamu.edu
daniel_tabor@tamu.edu
GitHub: [Tabor-Research-Group](https://github.com/Tabor-Research-Group)

Experience

- 2019- Assistant Professor, Department of Chemistry, Texas A&M University
- 2016-2019 Postdoctoral Fellow, Harvard University, Aspuru-Guzik Group
- 2011-2016 Research Assistant, University of Wisconsin–Madison, Sibert Group
- 2009-2011 Undergraduate Research Assistant, University of Texas at Austin, Stanton Group

Education

2016 Ph.D. Chemistry, University of Wisconsin–Madison

Advisor: Edwin L. Sibert III

Thesis: Extracting Structural Information from the OH and CH Stretch Spectral Regions with a Local Mode Approach

2011 B.S. Chemistry University of Texas at Austin

Advisor: John F. Stanton

Undergraduate Thesis: High-Accuracy Extrapolated Ab Initio Thermochemistry of Closed and Open Shell Molecules

Publications

*Independent Career, Corresponding Authors Indicated with **

Current Preprints

14. N.P.D. Sawaya,* D. Marti-Dafcik, Y. Ho, **D.P. Tabor**, D. Bernal, A.B. Magann, S. Premaratne, P. Dubey, A. Matsuura, N. Bishop, W.A. de Jong, S. Benjamin, O.D. Parekh, N. Tubman, K. Klymko,* and D. Camps,* HamLib: A library of Hamiltonians for benchmarking quantum algorithms and hardware. arXiv preprint: arXiv:2306.13126 (2023).

Published in Peer-Reviewed Journal

13. R.W. Neisser, J.P. Davis, M.E. Alfieri, H. Harkins, A.S. Petit,* **D.P. Tabor***,* and N.M. Kidwell,* Photophysical Outcomes of Water-Solvated Heterocycles: Single-Conformation Ultraviolet and Infrared Spectroscopy of Microsolvated 2-Phenylpyrrole *J. Phys. Chem. A* **127** (50), 10540–10554 (2023).
12. T. Ma, E. Fox, M. Ai, C.-H. Li, K.A.N. Sachinthan, K. Mohanty, **D.P. Tabor***,* E.B. Pentzer,* and J.L. Lutkenhaus,* Charge Transfer in Spatially Defined Organic Radical Polymers, *Chem. Mater.* **35**, 21, 9346-9351(2023).
11. C.-H. Li and **D.P. Tabor***,* Generative Organic Electronic Molecular Design Informed by Quantum Chemistry. *Chem. Sci.* **14**, 11045-11055 (2023). DOI: 10.1039/D3SC03781A
ChemRxiv Preprint (2023). DOI: 10.26434/chemrxiv-2023-bgcjg-v2
10. T.-H. Chao, S. Rekhi, J. Mittal, and **D.P. Tabor***,* Data-Driven Models for Predicting Intrinsically Disordered Protein Polymer Physics Directly from Composition or Sequence. *Mol. Syst. Des. Eng.* **8**, 1146-1155 (2023). DOI: <https://doi.org/10.1039/D3ME00053B>. ChemRxiv Preprint: 10.26434/chemrxiv-2023-wrnq1 (2023).
9. J. Li, B.-J. Peng, S. Li, **D.P. Tabor**, L. Fang,* and C.M. Schroeder,* Ladder-type conjugated molecules as robust multi-state single-molecule switches, *Chem*, **9**, 2282-2297 (2023). DOI: <https://doi.org/10.1016/j.chempr.2023.05.001>
8. T. Ma, C.-H. Li, R.M. Thakur, **D.P. Tabor**, and J.L. Lutkenhaus,* The role of the Electrolyte in Non-conjugated Radical Polymers for Metal-free Aqueous Energy Storage Electrodes, *Nat. Mater.* **22**, 495-502 (2023). DOI: <https://doi.org/10.1038/s41563-023-01518-z>
7. C.-H. Li and **D.P. Tabor***,* Reorganization Energy Predictions with Graph Neural Networks Informed by Low-Cost Conformers, *J. Phys. Chem. A*, **127**, 3484–3489 (2023). DOI: <https://doi.org/10.1021/acs.jpca.2c09030>
ChemRxiv preprint at: 10.26434/chemrxiv-2022-fk22p
6. N.J. Shuber, **D.P. Tabor**, S.W. North,* Theoretical Investigation of the Ground State Dissociation Pathways of CH₂NO₂, *Chem. Phys.* **568** 111823 (2023). DOI: <https://www.sciencedirect.com/science/article/abs/pii/S0301010423000058>
5. J. Lee, S. Li, X. Ji, S. Che, Y. Cao, **D.P. Tabor***,* L. Fang,* Molecular mechanism of rigidity- and planarity-promoted, state-dependent doping of conjugated ladder-type molecules, *Mater. Chem. Front.* **6**, 3329-3337 (2022). DOI: <https://doi.org/10.1039/D2QM00789D>
4. B. Peterson, M. Alfieri, D. Hood, C. Hettwer, D. Constantino, **D.P. Tabor***,* N.M. Kidwell,* Solvent-Mediated Charge Transfer Dynamics of a Model Brown Carbon Aerosol Chromophore: Photophysics of 1-Phenylpyrrole Induced by Water Solvation, *J. Phys. Chem. A* **126**, 4313–4325 (2022). ChemRxiv Preprint: 10.26434/chemrxiv-2022-4rt78.
3. G. Ma, M. Leng, S. Li, Z. Cao, Y. Cao, **D.P. Tabor***,* L. Fang,* and X. Gu,* Robust Chain Aggregation of Low-Entropy Rigid Ladder Polymer in Solution, *J. Mater. Chem. C* **10**, 13896-13904 (2022). DOI: <https://doi.org/10.1039/D2TC00761D>
2. C.-H. Li and **D.P. Tabor***,* Discovery of lead low-potential radical candidates for organic radical polymer batteries with machine-learning-assisted virtual screening, *J. Mater. Chem. A* **10**, 8273-8282 (2022). DOI: <https://doi.org/10.1039/D2TA00743F>. ChemRxiv Preprint: 10.26434/chemrxiv-2022-hlw2f.
1. N.P.D. Sawaya,* F. Paesani, and **D.P. Tabor***,* Near- and Long-Term Quantum Algorithmic Approaches for Vibrational Spectroscopy, *Phys. Rev. A* **104**, 062419 (2021). DOI: <https://doi.org/10.1103/PhysRevA.104.062419>
Preprint available at: arXiv:2009.05066

Work Prior to Independent Career

27. Q. Wang, Z. Yao, C. Zhao, T. Verhallen, **D.P. Tabor**, M. Liu, F. Ooms, F. Kang, A. Aspuru-Guzik, Y.-S. Hu, M. Wagemaker, and B. Li, Interface Chemistry of an Amide Electrolyte for Highly Reversible Lithium Metal Batteries, *Nat. Commun.* **11**, 4188 (2020). DOI: <https://doi.org/10.1038/s41467-020-17976-x>
26. C.F. Perkinson, **D.P. Tabor**, M. Einzinger, D. Sheberla, H. Utzat, T.-A. Lin, D.N. Congreve, M. Bawendi, A. Aspuru-Guzik, and M.A. Baldo, Discovery of Blue Singlet Exciton Fission Molecules via a High-Throughput Virtual Screening and Experimental Approach, *J. Chem. Phys.* **151**, 121102 (2019). DOI: <https://doi.org/10.1063/1.5114789>
25. **D.P. Tabor**, V. Chiykowski, P. Friederich, Y. Cao, D.J. Dvorak, C.P. Berlinguette, and A. Aspuru-Guzik, Design Rules for High Mobility Xanthene-Based Hole Transport Materials, *Chem. Sci.* **10**, 8360-8366 (2019). DOI: 10.1039/C9SC01491H
24. L. Tong, M.-A. Goulet, **D.P. Tabor**, E.F. Kerr, D. De Porcellinis, E.M. Fell, A. Aspuru-Guzik, R.G. Gordon, and M.J. Aziz, Molecular Engineering of an Alkaline Naphthoquinone Flow Battery, *ACS Energy Lett.* **4**, 1880-1887 (2019). DOI: <https://doi.org/10.1021/acsenergylett.9b01321>
Preprint: ChemRxiv doi:10.26434/chemrxiv.7732472.v1
23. **D.P. Tabor**†, R. Gómez-Bombarelli†, L. Tong, R.G. Gordon, M.J. Aziz, and A. Aspuru-Guzik, Mapping the Frontiers of Quinone Stability in Aqueous Media: Implications for Organic Aqueous Redox Flow Batteries, *J. Mater. Chem. A* **7**, 12833-12841 (2019). DOI: 10.1039/C9TA03219C.
Preprint: ChemRxiv doi:10.26434/chemrxiv.6990053.v2. †equal contribution.
22. M.-A. Goulet, L. Tong, D. A. Pollack, **D.P. Tabor**, S.A. Odom, A. Aspuru-Guzik, E.E. Kwan, R.G. Gordon and M.J. Aziz, Extending the Lifetime of Organic Flow Batteries via Redox State Management, *J. Am. Chem. Soc.* **141**, 8014-8019 (2019). DOI: 10.1021/jacs.8b13295
21. K. Alberi, M. Buongiorno Nardelli, A. Zakutayev, L. Mitas, S. Curtarolo, A. Jain, M. Fornari, N. Marzari, I. Takeuchi, M. Green, M. Kanatzidis, M. Toney, S. Butenko, B. Meredig, S. Lany, U. Kattner, A. Davydov, E. Toberer, V. Stevanovic, A. Walsh, N.G. Park, A. Aspuru-Guzik, **D.P. Tabor**, J. Nelson, J. Murphy, A. Setlur, J. Gregoire, H. Li, R. Xiao, A. Ludwig, L. Martin, A. Rappe, S.-H. Wei, and J. Perkins, The 2019 Materials by Design Roadmap, *J. Phys. D: Appl. Phys.* **52**, 013001 (2019). DOI: <https://doi.org/10.1088/1361-6463/aad926>
20. V. Chiykowski, Y. Cao, H. Tan, **D.P. Tabor**, E.H. Sargent, A. Aspuru-Guzik, and C.P. Berlinguette, Precise Control of Thermal and Redox Properties of Organic Hole-Transport Materials, *Angew. Chem. Int. Ed.* **57**, 15529 (2018). DOI: 10.1002/anie.201810809
19. D. Kwabi, K. Lin, Y. Ji, E.F. Kerr, M.-A. Goulet, D. De Porcellinis, **D.P. Tabor**, D.A. Pollack, A. Aspuru-Guzik, R.G. Gordon, and M.J. Aziz, Alkaline Quinone Flow Battery with Long Lifetime at pH 12, *Joule* **2**, 1894-1906 (2018). DOI: 10.1016/j.joule.2018.07.005 (Featured Article).
18. N.P.D. Sawaya, D. Rappoport, **D.P. Tabor**, and A. Aspuru-Guzik, Excitonics: A Set of Gates for Molecular Exciton Processing and Signaling, *ACS Nano* **12**, 6410-6420 (2018). DOI: 10.1021/acsnano.8b00584
Preprint: <https://doi.org/10.26434/chemrxiv.5309617.v1>
17. **D.P. Tabor**, Flow Batteries: Approaching Saturation Limits, *Nat. Energy*, **3**, 455-456 (2018). DOI: 10.1038/s41560-018-0169-1 (News and Views).
16. **D.P. Tabor**, L.M. Roch, S.K. Saikin, C. Kreisbeck, D. Sherberla, J.H. Montoya, S. Dwaraknath, M. Aykol, C. Ortiz, H. Tribukait, C. Amador-Bedolla, C.J. Brabec, B. Maruyama, K.A. Persson, and A. Aspuru-Guzik, Accelerating Discovery of Materials for Clean Energy in the Era of Smart Automation, *Nat. Rev. Mater.* **3**, 5-20 (2018). DOI: 10.1038/s41578-018-0005-z

15. Z. Yang, L. Tong, **D.P. Tabor**, E.S. Beh, M.-A. Goulet, D. De Porcellinis, A. Aspuru-Guzik, R.G. Gordon, and M.J. Aziz, Alkaline Benzoquinone Aqueous Flow Battery for Large-Scale Storage of Electrical Energy, *Adv. Energy Mater.* **8** 1702056 (2018). DOI: 10.1002/aenm.201702056
14. D.M. Hewett, **D.P. Tabor**, J.L. Fischer, E.L. Sibert III, and T.S. Zwier, IR-induced Fluorescence-Gain Spectroscopy: Conformation-Specific Excited State Infrared Spectra of the Alkylbenzenes, *J. Phys. Chem. Lett.* **8**, 5296-5300 (2017). DOI: 10.1021/acs.jpcclett.7b02276
13. D.M. Hewett, S. Bocklitz, **D.P. Tabor**, E.L. Sibert III, M. Suhm, and T.S. Zwier, Identifying the First Folded Alkylbenzene: Ultraviolet, Infrared, and Raman Spectroscopy of Pentylbenzene through Decylbenzene, *Chem. Sci.* **8**, 5305 (2017). DOI: 10.1039/C7SC02027A
12. D.J. Bakker, A. Dey, **D.P. Tabor**, Q. Ong, J. Mahé, M.-P. Gageot, E.L. Sibert III, and A.M. Rijs, Fingerprints of Inter- and Intramolecular Hydrogen Bonding in Saligenin-Water Clusters Uncovered by Mid- and Far-Infrared Spectroscopy, *Phys. Chem. Chem. Phys.* **19**, 20343-20356 (2017). DOI: 10.1039/C7CP01951C
11. P.R. Franke, **D.P. Tabor**, C.P. Moradi, G.E. Doublerly, J. Agarwal, H.F. Schaefer III, and E.L. Sibert III, Infrared Laser Spectroscopy of the *n*-Propyl and *i*-Propyl Radicals: Stretch-Bend Fermi Coupling in the Alkyl CH Stretch Region, *J. Chem. Phys.* **145**, 224304 (2016). DOI: 10.1063/1.4971239
10. J.A. Korn, **D.P. Tabor**, E.L. Sibert III, and T.S. Zwier, Conformation-Specific Spectroscopy of Alkyl Benzyl Radicals: Effects of a Radical Center on the CH Stretch Infrared Spectrum of an Alkyl Chain, *J. Chem. Phys.* **145**, 124314 (2016). DOI: 10.1063/1.4963227
9. **D.P. Tabor**, D.M. Hewett, S. Bocklitz, J.A. Korn, A.J. Tomaine, A.K. Ghosh, T.S. Zwier and E.L. Sibert III, Anharmonic Modeling of the Conformation-Specific IR Spectra of Ethyl, *n*-Propyl, and *n*-Butylbenzene, *J. Chem. Phys.* **144**, 224310 (2016). DOI: 10.1063/1.4953181
8. E.L. Sibert III, **D.P. Tabor**, and J.M. Lisy, Modeling The CH Stretch Vibrational Spectroscopy of M^+ [Cyclohexane] ($M=Li, Na, \text{ and } K$) Ions, *J. Phys. Chem. A* **119**, 10293-10299 (2015). DOI: 10.1021/acs.jpca.5b07461 (ACS Editors' Choice).
7. **D.P. Tabor**, R. Kusaka, P.S. Walsh, T.S. Zwier, and E.L. Sibert III, Local Mode Approach to OH Stretch Spectra of Benzene-(H_2O) $_n$ Clusters, $n = 2 - 7$, *J. Phys. Chem. A* **119**, 9917-9930 (2015). DOI: 10.1021/acs.jpca.5b06954
6. **D.P. Tabor**, R. Kusaka, P.S. Walsh, E.L. Sibert III, and T.S. Zwier, Isomer-Specific Spectroscopy of Benzene-(H_2O) $_n$, $n = 6, 7$: Benzene's Role in Re-Shaping Water's Three-Dimensional Networks, *J. Phys. Chem. Lett.* **6**, 1989-1995 (2015). DOI: 10.1021/acs.jpcclett.5b00786
5. Y.-F. Lee, W.-T. Chou, B.A. Johnson, **D.P. Tabor**, E.L. Sibert III, and Y.-P. Lee, Infrared Absorption of CH_3O and CD_3O Radicals Isolated in Solid Para- H_2 , *J. Mol. Spectrosc.* **310**, 57-67 (2015). DOI: 10.1016/j.jms.2014.11.008
4. E.L. Sibert III, **D.P. Tabor**, N.M. Kidwell, J.C. Dean, and T.S. Zwier, Fermi Resonance Effects in the Vibrational Spectroscopy of Methyl and Methoxy Groups, *J. Phys. Chem. A* **118**, 11272-11281 (2014). DOI: 10.1021/jp510142g
3. E.L. Sibert III and **D.P. Tabor**, A Perturbative Description of Non-Adiabatic Effects in Methoxy Vibrations, *Mol. Phys.* **112**, 3138-3143 (2014). DOI: 10.1080/00268976.2014.932455
2. **D.P. Tabor**, M.E. Harding, T. Ichino, and J.F. Stanton, High Accuracy Extrapolated Ab Initio Thermochemistry of the Vinyl, Allyl, and Vinyloxy Radicals, *J. Phys. Chem. A* **116**, 7668-7676 (2012). DOI: 10.1021/jp302527n

Technical Reports

1. Materials Acceleration Platform: Accelerating Advanced Energy Materials Discovery by Integrating High-Throughput Methods with Artificial Intelligence (co-author). View at: [link](#)

Funding Obtained in Independent Career

Budgeted Tabor Group Portion Indicated on Multi-Investigator Grants.

Current

Kureha Corporation Principal Investigator	1/1/2024-12/31/2024	\$ 83,230
Sandia National Laboratories Sandia LDRD Program Title: Cryptand Molecular Cavities to Capture and Chemically Process CO ₂ Co-PI (PI: Timothy S. Zwier, Sandia National Laboratory)	12/2023-09/2024	\$ 72,576
National Science Foundation NSF Award Number: 2339804 Title: CAREER: Development of Adaptive and Efficient Computational Inverse Design Methods for Organic Functional Materials Principal Investigator	12/15/2023-11/30/2028	\$ 625,000
National Science Foundation NSF Award Number: 2303044 Title: NSF Center for the Mechanical Control of Chemistry (Phase II) PI: James Batteas All other investigators in a CCI are Senior Personnel	10/01/2023-09/30/2028	\$ 327,242 (Estimated Group Portion)
Research Corporation for Science Advancement Cottrell Scholar Program Grant No. CS-CSA-2023-054 Title: Intelligent Optimization of Organic Photophysical Chemical Spaces Principal Investigator	07/01/2023-06/30/2026	\$100,000
Robert A. Welch Foundation Grant No. A-2049-20230405 Title: Accelerating the Prediction and Analysis of Vibrational and Electronic Spectroscopy Principal Investigator	06/01/2023-05/31/2026	\$300,000
National Science Foundation Grant No. DMR-2119672 Title: Collaborative Research: DMREF: Accelerated Design of Redox-Active Polymers for Metal-Free Batteries (Co-PI with Jodie Lutkenhaus (lead), Juan de Pablo, and Stuart Rowan)	10/01/2021-09/30/2025	\$429,587 (Estimated Group Portion)

Former

National Science Foundation Grant CMMI-1930277 Data Science Supplement to: Cellulose Nanocrystal-enabled Manufacturing of Carbon Nanotube Carbon fiber Polymer Composites (Subcontract for Data Science Supplement. PIs: Amir Asadi, Lisa Perez)	08/09/2019-08/31/2023	\$70,000
Robert A. Welch Foundation Grant No. A-2049-20200401 Title: Mapping the Structure and Formation of Aerosols Through Theoretical Spectroscopy and Multiscale Simulation Principal Investigator	06/01/2020-05/31/2023	\$240,000
Texas A&M T3: Triads for Transformation Closed-Loop Discovery Of Radical Biopolymers For Organic Battery Electrodes (Co-PI with Karen Wooley and Jodie Lutkenhaus)	01/01/2020-12/31/2021	\$10,000
Texas A&M Institute for Data Science: Data Science Career Initiation Fellow Program Title: Scientific Machine Learning for Chemical Physics Based Materials Design Principal Investigator	05/01/2021-12/31/2022	\$10,000

Honors and Awards*During Independent Career*

- 2023 Montague-Texas A&M Center for Teaching Excellence Scholar Award
- 2023 Cottrell Scholar
- 2022 Selected for “Hot-topic” Talk at Sanibel Symposium
- 2021 Texas A&M Institute for Data Science Career Initiation Fellow

Prior to Independent Career

- 2016 Richard and Joan Hartl Excellence in Research Award for Physical Chemistry, University of Wisconsin–Madison
- 2015 Poster Award, Physical Division, ACS National Meeting, Boston, Massachusetts
- 2015 Poster Award, Midwest Theoretical Chemistry Conference, Ann Arbor, Michigan
- 2013 Department of Chemistry Outstanding Teaching Assistant Award, University of Wisconsin–Madison
- 2011 Hirschfelder Graduate Fellowship, University of Wisconsin–Madison
- 2010-2011 Beckman Scholar

Presentations

Invited Seminars, Independent Career

1. University of Kentucky Department of Chemistry, November 2023.
2. University of Utah Department of Chemistry, October 2023.
3. Louisiana State University Department of Chemistry, April 2023.
4. Southern Methodist University Department of Chemistry, March 2023.
5. University of Mississippi Department of Chemistry, March 2023.
6. Texas A&M University–Kingsville Department of Chemistry, October 2022.
7. Duquesne University Department of Chemistry and Biochemistry, September 2022.
8. Los Alamos National Laboratory Theoretical Division and Center for Integrated Nanotechnologies (CINT), April 2022.
9. University of California-Merced Department of Chemistry and Biochemistry, April 2022.
10. University of Florida Theoretical Group Virtual Seminar Series. February 2021. Virtual talk given on Zoom.

Invited Talks, Independent Career

11. Machine-learning-driven approaches for organic electronic materials discovery and design, 265th ACS National Meeting. Physical Chemistry Division, March 2023, San Francisco, CA.
12. Data-driven methods for accelerating condensed phase vibrational spectroscopy modeling, 265th ACS National Meeting. Physical Chemistry Division, March 2023, Indianapolis, IN.
13. Data-Driven Methods for Accelerating Vibrational Spectroscopy Modeling at the Medium to Large Scale, Texas A&M High-Performance Research Computing Conference. May 2022. College Station, TX.
14. Data-Driven Methods for Accelerating Vibrational Spectroscopy Modeling at the Medium to Large Scale, ACS Southwest Regional Meeting. October 2021. Austin, TX.
15. Active Search for Organic Functional Materials with Machine Learning, Southwest Theoretical and Computational Chemistry Meeting. October, 2019. Norman, OK.

Contributed Talks, Independent Career

16. Learning Molecular Hamiltonians Directly from Spectra, 76th International Symposium on Molecular Spectroscopy, June 2023, Urbana, IL.
17. Assigning the vibrational spectra of clusters directly from the spectrum, 265th ACS National Meeting. Physical Chemistry Division, March 2023, Indianapolis, IN.
18. Developing Physically Motivated, Data-Driven Models for the Accurate Prediction of Disordered Protein Polymer Physics, 265th ACS National Meeting. Computers in Chemistry Division, March 2023, Indianapolis, IN.
19. Multiscale Simulation and Computational Design of Non-Conjugated Radical Polymers for Energy Storage Applications, 264th ACS National Meeting. Polymeric Materials: Science and Engineering Division, August 2022, Chicago, IL.

20. Inverse Infrared Spectroscopy via a Data-Driven Approach, 264th ACS National Meeting. Computers in Chemistry Division, August 2022, Chicago, IL.
21. Identification of the key molecular and polymeric levers for maximizing the conductivity of non-conjugated radical polymers via molecular simulation, 263rd ACS National Meeting. Physical Division, March 2022, San Diego, CA.
22. Data-Driven Methods for Accelerating Vibrational Spectroscopy Modeling at the Medium to Large Scale, Sani-bel Symposium. February 2022. St. Simon's Island, GA.
23. Structure and Infrared Spectra of New Aerosol Particle Formation Seed Clusters, 74th International Symposium on Molecular Spectroscopy. June 2021 (Virtual Talk).
24. Theoretical models and computational design of conductive non-conjugated radical polymers for energy storage, 261st ACS National Meeting. Computers in Chemistry Division, April 2021 (Virtual Talk).
25. Active Multi-objective Search for High-performance Organic Functional Materials, 2020 Prairie View A&M/Texas A&M Joint Data Blitz Symposium, Feb. 2020. Prairie View, TX.

Poster Presentations, Independent Career

26. Data-Driven Methods for Accelerating Condensed Phase Vibrational Spectroscopy Modeling, Gordon Conference on Liquids. August 2023, Holderness, NH.
27. Active, Multi-Objective Search for Organic Energy Storage Materials, Virtual Conference on Theoretical Chemistry. July 2020. Virtual Conference held in place of the American Conference on Theoretical Chemistry.

Invited Talks, Prior Independent Career

1. Flow Battery Electrolyte Discovery, JCESR Redoxmer Workshop, May 2019. Boston, MA.
2. Organic Functional Materials Discovery: The Importance of New Optimization Algorithms, 257th ACS National Meeting. Computers in Chemistry Division. March 2019. Orlando, FL
3. Organic Flow Battery Electrolyte Discovery: Importance of New Optimization Algorithms, European Energy Research Alliance Autonomous Materials Development Platforms Workshop. October 2018. Brussels, Belgium.
4. Accelerated Discovery of Materials for Organic Redox Flow Batteries, Mission Innovation Business Day. May 2018. Malmö, Sweden.
5. Modeling Vibrational Spectra with a Local Mode Approach, University of Wisconsin-Madison Departmental Awards Seminar. May 2016. Madison, WI.

Contributed Talks, Prior Independent Career

6. Discovery of Organic Flow Battery Electrolytes via a Machine Learning Driven Approach, 256th ACS National Meeting. Energy and Fuels Division. August 2018. Boston, MA.
7. Coupling Generative Models to Virtual Screening: Application to Organic Redox Flow Batteries, 255th ACS National Meeting. Computers in Chemistry Division. March 2018. New Orleans, LA.
8. Influence of Aromatic Molecules on The Structure and Spectroscopy of Water Clusters, 71st International Symposium on Molecular Spectroscopy. June 2016. Urbana, IL.

9. Modeling the Conformation-Specific Infrared Spectroscopy of N-Alkylbenzenes, 71st International Symposium on Molecular Spectroscopy. June 2016. Urbana, IL
10. Theoretical Study of the IR Spectroscopy of Benzene-(Water)_n Clusters, 70th International Symposium on Molecular Spectroscopy. June 2015. Urbana, IL.
11. Vibrational Spectroscopy of Benzene-(Water)_n Clusters with $n = 6, 7$, 70th International Symposium on Molecular Spectroscopy. June 2015. Urbana, IL.
12. Theoretical Study of the Ethyl Radical, 69th International Symposium on Molecular Spectroscopy. June 2014. Urbana, IL.
13. Theoretical Study of the Ethyl Radical, 68th International Symposium on Molecular Spectroscopy. June 2013. Columbus, OH.

Poster Presentations, Prior Independent Career

14. Accelerating the Rate of Discovery of Organic Functional Materials, Flatiron Institute Center for Computational Quantum Physics Quantum Chemistry Workshop. November 2017. New York, NY.
15. High-Throughput Virtual Screening of Flow Battery Electrolytes, American Conference on Theoretical Chemistry. July 2017. Boston, MA.
16. High-Throughput Virtual Screening of Flow Battery Electrolytes, MIT Energy Night. October 2016. Cambridge, MA.
17. Theoretical Study of the IR Spectroscopy of Benzene-(Water)_n Clusters with $n = 3 - 7$, 250th ACS National Meeting. Physical Division. August 2015. Boston, MA.
18. Theoretical Study of the IR Spectroscopy of Benzene-(Water)_n Clusters with $n = 3 - 7$, Midwest Theoretical Chemistry Conference. June 2015. Ann Arbor, MI.
19. Probing Molecular Structure with CH Chromophores” Gordon Research Conference on Vibrational Spectroscopy. August 2012. Biddeford, ME.
20. Towards a Potential Energy Surface for Vibrational Analysis of the Ethyl Radical, Midwest Theoretical Chemistry Conference. June 2012. Madison, WI.
21. Computing Highly Accurate Enthalpies of Formation via Isodesmic and Hypohomodesmotic Schemes, Beckman Research Symposium. August 2011. Irvine, CA.
22. Computing Highly Accurate Enthalpies of Formation via Isodesmic and Hypohomodesmotic Schemes, Dynamics of Molecular Collisions Conference. July 2011. Snowbird, UT.

Teaching and Mentoring

Teaching

Courses Taught

Fall 2021, Fall 2022, Fall 2023 Chemistry 119H (Honors Fundamentals of Chemistry I)

Fall 2020, Fall 2021 Chemistry 631 (Statistical Thermodynamics Graduate Level)

Fall 2019, Fall 2020, Fall 2022, Fall 2023 Chemistry 648 (Principles of Quantum Mechanics Graduate Level)

Mentoring

Current Graduate Students

2022- Hayden Moran

2023 NDSEG Fellow

2021- Katie Felde

2021- Daniel Doria

2021- Tzu-Hsuan Chao

2020- Abigail Moody

2019- Cheng-Han Li

Current Postdoctoral Fellows

2023- Vijay Sundar

Current Undergraduate Students

2024- Tanya Chandra

2024- Manasi Ramkumar

2022- David Chi

2020- Curran Watson

Former Group Members

Graduate Students

2021-2022 Emily Chappie

Current: OpenEye

2020- Jezrielle Annis

2020 DOE CSGF Fellow

Undergraduates

2023 Autumn Kimber

2023 Alejandro Mejias

2023 Fernando Ramirez

2022-23 Samantha Rosal

Current Ph.D. student in computational chemistry, Washington University St. Louis

2022-2023 Jae Trinh

2022 Avi Bedi (2022 REU Student from Butler University)

2021-2022 Zachary Graham

2021 Daniel Hernandez

2021 Tyler Paxton

2021 Logan Hendricks

2021 David A. González Narváez (2021 Chemistry Department NSF REU Student)

Current Ph.D. student in theoretical chemistry, Columbia University

2021 Olivia McNally (2021 Materials Science and Engineering Department NSF REU Student, co-advised with Prof. Emily Pentzer)

2021 Brittany DeNicholas (2021 Materials Science and Engineering Department NSF REU Student, co-advised with Prof. Emily Pentzer)

2021 Jia Li (2021 REU Student, co-advised with Prof. Amir Asadi)

2020-2021 Emily Chappie (Remote Visiting Scholar from the Kidwell Group at William & Mary)

2020-2022 Brianna Bishop

2021 Goldwater Scholar

Postdoctoral Fellows

2020-2022 Shi Li

Now at Argonne National Laboratory

Service

University and Department Service

2022-2024 Department of Chemistry Graduate Admissions Committee

2022- Chair of Texas A&M Local ACS Section

2022- Facilitator for Texas A&M Graduate Mentoring Academy

2021- Director, Undergraduate Chemistry Honors Program

2021 College of Science Information Technology Director Search Committee Member

2021 Laboratory for Molecular Simulation Research Scientist Search Committee Member

2019-2022, 2023-2024 Faculty Candidate Search Committee Member

2020 Hullabaloo U Instructor

2020- Department Proactive Recruiting Operations (PROps) Committee Member

2019-2023 Department Information Technology Committee Member

Scientific Committees and Professional Activities

- 2022-2023 Co-editor (with Zhou Lin) of Special Issue on Machine Learning in Spectroscopy for the Journal of Molecular Spectroscopy
- 2022 Poster Session Judge for Physical Division at Spring 2022 ACS Meeting
- 2022 Mail-in Reviewer for U.S. Department of Energy
- 2022 Reviewer for TAMU Qatar Proposals
- 2022 Co-organizer (with Kelvin Lee) of the Machine Learning in Spectroscopy Minisymposium at the International Symposium for Molecular Spectroscopy
- 2021 Mail-in Reviewer for U.S. Department of Energy
- 2021 Panel Reviewer and ad hoc Reviewer for National Science Foundation
- 2020 Participant, JCESR ML/AI Virtual Workshops
- 2019 Invited Participant, NSF Summit on Big Data and Cyberinfrastructure for Materials Research, Chicago, Illinois
- 2019 Panel Reviewer for U.S. Department of Energy
- 2019 Participant, Cottrell Scholars Collaborative/ACS New Faculty Workshop, Pasadena, CA
- 2018 Panel Reviewer: Sectoral Fund CONACYT-Secretariat of Energy-Energy Sustainability, Mexico
- 2018 Invited Participant, ARPA-E Machine Learning Workshop, Falls View, Virginia
- 2017 Participant, Mission Innovation Clean Energy Materials Innovation Challenge Workshop, Mexico City, Mexico

Scientific Refereeing of Publications

Referee: *ACS Appl. Polym. Mater.*, *ACS Cent. Sci.*, *ACS Omega*, *Adv. Mater.*, *Adv. Theor. Sim.*, *Batter. Supercaps*, *Cell Rep. Phys. Sci.*, *Chem. Mater.*, *Chem. Sci.*, *Energy Env. Sci.*, *J. Mater. Chem. A*, *J. Phys. Chem. A*, *J. Phys. Chem. C*, *Joule*, *Mater. Horiz.*, *Mol. Phys.*, *Nanoscale Horiz.*, *Nat. Energy*, *Phys. Chem. Chem. Phys.*, *RSC Adv.*, *Trends Chem.* (61 total manuscripts since 2019)

Outreach and Related Activities

- 2022, 2023 Co-organizer of Chemistry High School Summer Camp Focused on Light and Matter Interactions for High School Students (co-organized with Lei Fang)
- 2022 Young Researcher Conference Co-organizer for Alliance for Diversity in Science and Engineering (co-organizer with Osvaldo Gutierrez, Emily Pentzer, and Andy Thomas)
- 2018-2019 Board Member of Alliance for Diversity in Science and Engineering

Professional Society Memberships

2009- Member, American Chemical Society

2016- Member, Materials Research Society

2020- Member, American Physical Society

Last updated: January 25, 2024