Shane M. Parker, PhD (September 05, 2024)

Frank Hovorka Assistant Professor of Chemistry Case Western Reserve University Department of Chemistry 10900 Euclid Ave., Cleveland, OH 44106

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positions

2021 –	Frank Hovorka Assistant Professor of Chemistry Case Western Reserve University	Cleveland, OH
2019 – 2021	Assistant Professor of Chemistry Case Western Reserve University	Cleveland, OH
2014 – 2019	Postdoctoral Fellow (as Arnold O. Beckman Postdoctoral Fellow 2016 2019) University of California Irvine Advisor: Professor Filipp Furche	Irvine, CA
2009 – 2014	PhD, Theoretical Chemistry Northwestern University <i>Advisor</i> : Professor Mark Ratner and Professor Tamar Seideman	Evanston, IL
2008 – 2009	Fulbright Fellow Technische Universität München Advisor: Professor Notker Rösch	Munich, Germany
2004 – 2008	B.S., Chemistry & B.S., Mathematics University of Florida Advisor: Professor N. Yngve Öhrn	Gainesville, FL

professional affiliations

- Member of the American Chemical Society
- Member of the American Physical Society

awards

2021	Finalist, John S. Diekhoff Award for Distinguished Graduate Student Teaching, CWRU
2021 - 2024	Frank Hovorka Assistant Professor of Chemistry, CWRU
2020 - 2021	Glennan Fellowship, CWRU Pedagogical Award
2020	STAIR Mentor Fellows Program, CWRU
2019	Nominated, John S. Diekhoff Award for Distinguished Graduate Student Teaching, CWRU
2016 - 2019	Arnold O. Beckman Postdoctoral Fellowship
2010 - 2013	Department of Energy Office of Science Graduate Fellowship
2009	Participant, 59th Lindau Nobel Laureate Meeting
2008 - 2009	Fulbright Fellowship Technische Universität München
2006	Anderson Scholar with Highest Distinction, UF academic achievement and uninterrupted study

publications corresponding authorship denoted with *

Case Western Reserve University

26 W. B. Martin, R. E. Warburton, <u>S. M. Parker</u>, V. O. Rodionov, On the characterization of γ-graphyne, *Nature Synthesis* **2024**, doi: 10.1038/s44160-024-00642-1, chemRxiv:2023.xl3gp.v3

25 Z. Zhou, S. M. Parker*,

Converging TDDFT calculations in 5 iterations with minimal auxiliary preconditioning, *Journal of Chemical Theory and Computation* **2024**, 20, 6738-6746, doi: 10.1021/acs.jctc.4c00577, arXiv:2404.17133

- 24 E. Roy Miller, S. J. Hoehn, A. Kumar, D. Jiang, <u>S. M. Parker</u>*, Ultrafast Photochemistry and Electron Diffraction for Cyclobutanone in the S₂ State: Surface Hopping with Time-Dependent Density Functional Theory, *The Journal of Chemical Physics* **2024**, *161*, 034105, doi: 10.1063/5.0203679, arXiv:2402.10336
- 23 Y. J. Franzke, C. Holzer, J. H. Andersen, T. Begusic, F. Bruder, S. Coriani, F. Della Sala, E. Fabiano, D. A. Fedotov, S. Fürst, S. Gillhuber, R. Grotjahn, M. Kaupp, M. Kehry, M. Krstic, F. Mack, S. Majumdar, B. D. Nguyen, S. M. Parker, F. Pauly, A. Pausch, E. Perlt, G. S. Phun, A. Rajabi, D. Rappoport, B. Samal, T. Schrader, M. Sharma, E. Tapavicza, R. S. Treß, V. Voora, A. Wodynski, J. M. Yu, B. Zerulla, F. Furche, C. Hättig, M. Sierka, D. P. Tew, F. Weigend, Turbomole: Today and Tomorrow, Journal of Chemical Theory and Computation 2023, 19, 6859-6890, doi: 10.1021/acs.jctc.3c00
- 22 Z. Zhou, F. Della Sala, <u>S. M. Parker</u>*, Minimal auxiliary basis set approach for the electronic excitation spectra of organic molecules, *The Journal of Physical Chemistry Letters* **2023**, *14*, 1968-1976, doi: 10.1021/acs.jpclett.2c03698, submitted version
- 21 V. G. Desyatkin, W. B. Martin, A. E. Aliev, N. E. Chapman, A. F. Fonseca, D. S. Galvão, E. Roy Miller, K. H. Stone, Z. Wang, D. Zakhidov, F. T. Limpoco, S. R. Almahdali, <u>S. M. Parker</u>, R. H. Baughman, V. O. Rodionov, Scalable Synthesis and Characterization of Multilayer γ-Graphyne, New Carbon Crystals with a Small Direct Band Gap, *Journal of the American Chemical Society* **2022**, *144*, 17999-18008, doi: 10.1021/jacs.2c06583, arXiv:2301.05291
- 20 Z. Zhou, <u>S. M. Parker</u>*, Accelerating molecular property calculations with semiempirical preconditioning, *The Journal of Chemical Physics* **2021**, *155*, 204111, doi: 10.1063/5.0071013, accepted version
- 19 M. Gupta, M. Singha, D. Rasale, Z. Zhou, S. Bhandari, S. Beasley, J. Sakr, <u>S. M. Parker</u>, R. Spitale, Mutually orthogonal bioconjugation of vinyl nucleosides for RNA metabolic labeling, *Organic Letters* **2021**, *23*, 7183-7187, doi: 10.1021/acs.orglett.1c02584
- 18 J. Jimenez, Z. Zhou, A. L. Rheingold, <u>S. M. Parker</u>, G. Sauvé, Tuning the properties of azadipyrromethene-based near-infrared dyes using intramolecular BO-chelation and peripheral substitutions, *Inorganic Chemistry* **2021**, *60*, 13320-13331, doi: 10.1021/acs.inorgchem.1c01597
- 17 <u>S. M. Parker*</u>, C. J. Schiltz, Surface hopping with cumulative probabilities: even sampling and improved reproducibility, *The Journal of Chemical Physics* **2020**, *153*, 174109, doi: 10.1063/5.0024372, accepted version
- 16 S. Balasubramani, G. P. Chen, S. Coriani, M. Diedenhofen, M. S. Frank, Y. J. Franzke, F. Furche, R. Grotjahn, M. E. Harding, C. Hättig, A. Hellweg, B. Helmich-Paris, C. Holzer, U. Huniar, M. Kaupp, A. Marefat Khah, S. Karbalaei Khani, T. Müller, F. Mack, B. D. Nguyen, S. M. Parker, E. Perlt, D. Rappoport, K. Reiter, S. Roy, M. Rückert, G. Schmitz, M. Sierka, E. Tapavicza, D. P. Tew, C. van Wüllen, V. K. Voora, F. Weigend, A. Wodyński, J. M. Yu, TURBOMOLE: Modular program suite for ab initio quantum-chemical and condensed-matter simulations, *The Journal of Chemical Physics* 2020, 152, 184107, doi: 10.1063/5.0004635

Before Case Western Reserve University

- 15 <u>S. M. Parker</u>, S. Roy, F. Furche, Multistate hybrid time-dependent density functional theory with surface hopping accurately captures ultrafast thymine photodeactivation, *Physical Chemistry Chemical Physics* **2019**, *21*, 18999-19010, doi: 10.1039/C9CP03127H
- 14 M. Kubota, S. Nainar, S. M. Parker, W. England, F. Furche, R. Spitale, Expanding the Scope of RNA Metabolic Labeling with Vinyl Nucleosides and Inverse Electron-Demand Diels-Alder Chemistry, ACS Chemical Biology 2019, 14, 1698-1707, doi: 10.1021/acschembio.9b00079
- 13 S. M. Parker, D. Rappoport, F. Furche, Quadratic response properties from TDDFT: trials and tribulations, *Journal of Chemical Theory and Computation* **2018**, 14, 807-819, doi: 10.1021/acs.jctc.7b01008
- 12 M. Muuronen, S. M. Parker, E. Berardo, A. Le, M. Zwijnenburg, F. Furche, Mechanism of Photocatalytic Water Oxidation on Small TiO₂ Nanoparticles, *Chemical Science* 2017, 8, 2179-2183, doi: 10.1039/C6SC04378J

- 11 S. M. Parker, S. Roy, and F. Furche,
 - Unphysical divergences in response theory, The Journal of Chemical Physics 2016, 145, 134105, doi: 10.1063/1.4963749
- 10 V. A. Nasluzov, S. M. Parker, A. Genest, A. M. Shor, E. A. Ivanova-Shor, N. Rösch, Trinuclear tantalum clusters grafted to hydroxylated silica surfaces: A density-functional embedded-cluster study, Kinetics and Catalysis 2015, 56, 631, doi: 10.1134/S0023158415050134
- 9 I. Kim, <u>S. M. Parker</u>, T. Shiozaki, Orbital Optimization in the Active Space Decomposition Model, *Journal of Chemical Theory and Computation* **2015**, *11*, 3636, doi: 10.1021/acs.jctc.5b00429, arXiv:1505.02346
- 8 S. M. Parker, T. Shiozaki,

Active space Decomposition with multiple sites: Density matrix renormalization group algorithm, *The Journal of Chemical Physics* **2014**, *141*, 211102, doi: 10.1063/1.4902991, arXiv:1410.6407

- 7 S. M. Parker, M. Smeu, I. Franco, M. A. Ratner, T. Seideman, Molecular Junctions: Can Pulling Influence Optical Controllability, *Nano Letters* **2014**, *14*, 4587, doi: 10.1021/nl501629c
- 6 S. M. Parker, T. Shiozaki, Quasi-diabatic states from active space decomposition, *Journal of Chemical Theory and Computation* 2014, 10, 3738, doi: 10.1021/ct5004753
- 5 <u>S. M. Parker</u>*, T. Seideman, M. A. Ratner, T. Shiozaki, Model Hamiltonian analysis of singlet fission from first principles, *Journal of Physical Chemistry C* **2014**, *118*, 12700, doi: 10.1021/jp505082a
- doi: 10.1021/jp505082a
 4 <u>S. M. Parker</u>*, T. Seideman, M. A. Ratner, T. Shiozaki,

Active-space decomposition for molecular dimers, The Journal of Chemical Physics 2013, 139, 021108, doi: 10.1063/1.4813827

- 3 Y. Wu, V. A. Karttunen, <u>S. M. Parker</u>, A. Genest, N. Rösch, Olefin Hydrosilylation Catalyzed by a Bis-N-Heterocyclic Carbene Rhodium Complex. A Density Functional Theory Study, *Organometallics* **2013**, *32*, 2363, doi: 10.1021/om301236n
- 2 S. M. Parker, M. A. Ratner, T. Seideman, Simulating strong field control of axial chirality using optimal control theory, *Molecular Physics* 2012, 110, 1941, doi: 10.1080/00268976.2012.695808
- 1 <u>S. M. Parker</u>, M. A. Ratner, T. Seideman, Coherent control of molecular torsion, *The Journal of Chemical Physics* **2011**, *135*, 224301, doi: 10.1063/1.3663710

book chapters

Before Case Western Reserve University

1 S. M. Parker, F. Furche

Frontiers in Quantum Chemistry, edited by M. J. Wójcik, H. Nakatsuji, B. Kirtman, Y. Ozaki, Springer Singapore (2018) Response theory and molecular properties

reports in media and professional journals

2020	Scilight featured article	Photochemistry simulations dramatically improve reliability and reproducibility
2020	JCP Editor's choice	Cumulative surface hopping paper highlighted by AIP
2019	PCCP Editor's choice	Multi-state TDDFT paper selected as outstanding article by Editor-in-Chief

talks

Case Western Reserve University

September 2024	TDDFT-ris: A semiempirical model and preconditioner for fast and accurate sp (invited)	ectra and nonlinear properties Oxford, UK
	Turbomole User and Developer Meeting	
July 2024	TDDFT-ris: A semiempirical model and preconditioner for fast and accurate sp (invited)	ectra and nonlinear properties Chicago, IL
June 2024	CECAM Reaction Prediction Workshop Towards Photochemistry with the Resonating Mean-Field Method (invited) ACS Mid-Atlantic Regional Meeting	State College, PA
June 2024	Towards Photochemistry with the Resonating Mean-Field Method (invited) Madison, WI	University of Wisconsin-Madison,
	Midwest Theoretical Chemistry Conference 2024	
March 17-18, 2024	(1) TDDFT-ris: A semiempirical model and preconditioner for fast and accurate s probabilities make surface hopping simulations cheaper and more reproducible Meeting of the American Chemical Society	
January 16, 2024	Simulating Photochemistry from First-Principles Quantum Chemistry (invited) University of Akron	Akron, OH
September 11, 2023	Computational Photochemistry (invited) Binghamton University, Department of Physics	Binghamton, NY
July 6, 2023	Fast spectra with the minimal auxiliary basis approach to TDDFT (invited) Rutgers TDDFT Workshop	Rutgers, NJ
June 3, 2023	Fast spectra with the minimal auxiliary basis approach to TDDFT Midwest Theoretical Chemistry Conference 2023	urdue University, West Lafayette, IN
October 2022	Fast Spectra with the minimal auxiliary basis approach to TDDFT (invited) Benasque 9th TDDFT Workshop	Benasque, Spain
August 2022	Accelerating Quantum Chemistry with Semiempirical Preconditioning (invited) CWRU-Tohoku 8th Annual Data Science in Engineering and Life Sciences	
June 2022	Accelerating TDDFT with Semiempirical Preconditioning 52nd MWTCC	Columbus, Ohio
March 2022	Cumulative Surface Hopping: Faster and More Reproducible APS March Meeting 2022	Chicago, Illinois
Before Case Wester	n Reserve University	
August 2018	Excited-State Chemistry with TDDFT (invited) 256th American Chemical Society National Meeting and Exposition	Boston, Massachussetts
August 2018	Ensemble Optimized Time-Dependent Density Functional Theory (invited) 256th American Chemical Society National Meeting and Exposition	Boston, Massachussetts
March 2018	Quadratic response properties from TDDFT: trials and tribulations APS March Meeting	Los Angeles, California
July 2017	Nonlinear properties from TDDFT: trials and tribulations (invited) Excited States: Electronic Structure and Dynamics	Telluride, Colorado
April 2017	Nonlinear properties from TDDFT: trials and tribulations (invited) 253rd American Chemical Society National Meeting and Exposition	San Francisco, California
March 2016	Diagnosis and implications of spurious poles in the quadratic response of ap	proximate electronic structure
	methods	San Diego, California
	251st American Chemical Society National Meeting and Exposition	
June 2015	Non-adiabatic molecular dynamics (invited)	Ottawa, Ontario
1	98th Meeting of the Canadian Society of Chemistry	
June 2014	Model Hamiltonians from the Active-space Decomposition Method (invited) McMaster University, Department of Chemistry	Hamilton, Ontario

pedagogical lectures

Case Western Reserve University

October 2022 TDDFT in Chemistry and Biochemistry (2 lectures)

Benasque 9th TDDFT School

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July 2017 Nonadiabatic molecular dynamics with TDDFT (2 lectures)

Telluride School on Time-dependent Density Functional Theory

Telluride, Colorado

Benasque, Spain

support

funded support

• Project Title: Photochemistry with Resonating Mean-Field

Amount: \$650,000 (100% Parker)

Role: Pl

Source: NSF CAREER, Division of Chemistry (CHE)

Status: current

Start Date: 06/01/2023 End Date: 05/30/2028

Award ID: CHE-2236959

• Project Title: Chemistry on Computers at CWRU

Amount: \$6,500

Role: PI

Source: Case Western Reserve University

Status: completed

Start Date: 07/2020 **End Date:** 06/2021

student fellowships

• student: Ericka Miller

title: Implementing an efficient, stable, and open-source version of the state-averaged Resonating Hartree-Fock method

for use in photochemical applications

fellowship: Molecular Sciences Software Fellowship

amount: \$53,092

dates - 07/01/2023 - 06/30/2024

collaborators

Case Western Reserve University

- Prof. Genevieve Sauve (CWRU Chemistry)
- Prof. Valentin Rodionov (CWRU Chemistry)
- Prof. Clemens Burda (CWRU Chemistry)
- Prof. Carlos Crespo (CWRU Chemistry)
- Prof. Divita Mathur (CWRU Chemistry)
- Prof. Matthew Bertin (CWRU Chemistry)
- Prof. Metin Karayilan (CWRU Chemistry)

External

- Prof. Robert Spitale (University of California, Irvine Department of Chemistry)
- Prof. Fabio Della Sala (Center for Biomolecular Nanotechnologies, Istituto Italiano di Tecnologia; Institute for Microelectronics and Microsystems (CNR-IMM))

service

- Member, Faculty Search Committee (AY2023-24)
- Member, Artificial Intelligence Screening Committee (2023-24)
- Member, Graduate Program Review Committee (Fall 2023)
- Panelist, NSF CAREER Writing Workshop (2023)
- Co-founder and Co-organizer, CWRU Chemistry Research Symposium (2023–)
- Panel Reviewer, Squire Scholarship Interview Panel (2023)

- Member, Graduate Admissions Committee (2019–)
- Co-chair, Colloquium Committee (2019–)
- Panel Reviewer, Expanding Horizons Initiative (2023)
- Member, Faculty Search Committee (AY2021-22)
- Member, Curriculum Committee (Fall 2020 (Sabbatical Replacement))
- Member, Chemistry Executive Committee (2019–2021)

professional service

peer review

- Physical Chemistry Chemical Physics (12)
- Journal of Physical Chemistry Letters (6)
- The Journal of Chemical Physics (4)
- Journal of Computational and Theoretical Chemistry (3)
- Wiley Interdisciplinary Reviews (1)
- Nanoscale Advances (1)
- Inorganica Chimica Acta (1)
- The European Physical Journal B (1)

grant review panels

- Department of Energy
- National Science Foundation

courses taught

- CHEM446/337: Quantum Mechanics I (3 credits) Fall 2019, Fall 2021, Fall 2023
- CHEM447: **Modern Physical Chemistry (co-taught)** (3 credits) Fall 2019, Fall 2022
- CHEM336: Physical Chemistry II (3 credits) Spring 2020, Spring 2021, Spring 2023
- CHEM442/342: **Computational Chemistry** (3 credits) Fall 2020, Fall 2022

advisees

postdoctoral researchers

- Dr. Maybe You (2024)
- Dr. Sean Hoehn (2022 2024) now at AkzoNobel
- Dr. Srijana Bhandari (2020 2022) now at PNNL

Ph.D. students

- Ericka Miller (2019)
- Abhijith Kumar (2021)
- Dehua Jiang (2022)
- Praseetha Prakash (2023)
- John Zhou (2019 2023) now at Changping National Laboratory

undergraduate students

- Thomas-Ethan Kaji (2022)
- William Kattner (2022 2023) now at Medpace
- Annabella Debernardo (2019 2021) now at UIUC
- Colin Schiltz (2019 2021) now at UC Irvine
- Krista Schoonover (2020 2021) now at Texas A&M

student awards

2023	Ericka Miller Best Presentation at 2023 Midwest Theoretical Chemistry Conference
2023	Ericka Miller Best Presentation at 2023 CWRU Chemistry Research Symposium
2023	Ericka Miller Molecular Sciences Software Institute (MolSSI) Software Fellowship