Eric Van Dornshuld, Ph.D.

Department of Chemistry, Mississippi State University, Mississippi State, MS 39762 | Office: Hand 1130 | Office Phone: +1.662.325.8452 | edornshuld@chemistry.msstate.edu | dornshuld.chemistry.msstate.edu

EXPERIENCE

Mississippi State University, Mississippi State, MS Assistant Clinical Professor Instructor Post-Doctoral Teaching/Research Associate	2018-present 2017-2018 2014-2017
The University of Mississippi, University, MS Research Assistant Teaching Assistant	2009–2014 2009–2010
The University of North Florida, Jacksonville, FL Research Assistant Teaching Assistant	2006–2009 2007–2008
EDUCATION	
Ph.D. Chemistry The University of Mississippi, University, MS Dissertation: "Characterizing non-covalent interactions and peptide bond formation with electronic structure theory" Advisor: Prof. Gregory S. Tschumper	August 2014
B.S. Chemistry The University of North Florida, Jacksonville, FL Advisor: Prof. Robert Vergenz (retired)	May 2009
AWARDS	
ACS Student Chapter Awards – Ranked <i>Outstanding</i> ChemDawg Recognition Award for "Outstanding effort and contributions to the Department of Chemistry"	2021 2020
MS ACS Local Section – ACS ChemLuminary Award for "Best National Chemistry Week Event Organized By a Student Group"	2019
MS ACS Local Section – ACS Student Chapter of the Year (SMACS)	2018
ACS Graduate Research Award	2014
Dissertation Fellowship Award	2014
GRANTS	
ACS Innovative Project Grant	2019

PUBLICATIONS

- 1. Olive, L. N., E. V. Dornshuld and C. E. Webster. The curious case of DMSO: A CCSD(T)/CBS(aQ56+d) benchmark and DFT study *J. Chem. Phys.* **2021**. 114304, doi:10.1002/jcc.23522
- 2. Frey, N. C., E. V. Dornshuld and C. E. Webster. Benchmarking the Fluxional Processes of Organometallic Piano-Stool Complexes *Molecules*. **2021**. *26*, 2310, doi:10.3390/molecules26082310
- 3. Bhavsar-Jog, Y., E. V. Dornshuld, T.A. Brooks, G. S. Tschumper, and R. M. Wadkins. Co-Localization of DNA i-Motif-Forming Sequences and 5-Hydroxymethyl-cytosines in Human Embryonic Stem Cells. *Molecules.* **2019**. *24*, 3619, doi:10.3390/molecules24193619.

- 4. Dornshuld, E. V. and G. S. Tschumper. Big Changes for Small Noncovalent Dimers: Revisiting the Potential Energy Surfaces of (P₂)₂ and (PCCP)₂ with CCSD(T) Optimizations and Vibrational Frequencies. *J. Chem. Theory Comput.* **2016**. *12*, 4, 1534–1541, doi:10.1021/acs.jctc.5b01105.
- 5. Dornshuld, E. V., R. A. Vergenz, and G. S. Tschumper. Peptide bond formation via glycine condensation in the gas phase. *J. Phys. Chem. B.* **2014**. *118* (29), 8583–8590, doi:10.1021/jp504924c.
- 6. Dornshuld, E. V., C. M. Holy and G. S. Tschumper. Homogeneous and heterogeneous non-covalent dimers of formaldehyde and thioformaldehyde: structures, energetics, and vibrational frequencies. *J. Phys. Chem. A.* **2014**. *118* (18), 3376–3385, doi:10.1021/jp502588h.
- 7. Bhavsar-Jog, Y., E. V. Dornshuld, T.A. Brooks, G. S. Tschumper, and R. M. Wadkins. Epigenetic modification, dehydration, and molecular crowding effects on the thermodynamics of i-motif structure formation from C-rich DNA. *Biochemistry*. **2014**. *53* (10), 1586–1594, doi:10.1021/bi401523b.
- 8. Dornshuld, E. V. and G. S. Tschumper. Characterization of the potential energy surfaces of two small but challenging noncovalent dimers: (P₂)₂ and (PCCP)₂. *J. Comput. Chem.* **2014**. *35* (6), 479–487, doi:10.1002/jcc.23522.
- 9. Reddy, V., R. Kota, E. V. Dornshuld, D. L. Mattern, G. S. Tschumper, D. Jiang, and A. Dass. Interstaple dithiol cross-linking in Au₂₅(SR)₁₈ nanomolecules: a combined mass spectrometric and computational study. *J. Am. Chem. Soc.*. **2011**. *133* (50), 20258–20266, doi:10.1021/ja206436x.
- 10. Dornshuld, E. V., M. Zhang, T. Keith Hollis, and C. E. Webster. Predicting ¹⁹⁵Pt-NMR chemical shifts in organometallic compounds with non-relativistic density functional approaches. (in preparation)

TEACHING EXPERIENCE

CH-1213 – Chemistry I CH-3141 – Prof. Chem: Literature

CH-1141 – Prof. Chem: Paths CH-8990 – Prof. Chem: Current Problems (compu-

CH-2141 – Prof. Chem: Tools tational chemistry project support)

ONLINE COURSE DEVELOPMENT

CH-1223 - Chemistry II Online Resources

https://dornshuld.chemistry.msstate.edu/courses/chem2/

Chemistry Student Handbook

https://dornshuld.chemistry.msstate.edu/books/chemistry/

WEBSITE LIFETIME ANALYTICS (03/2020 to 09/2022)

Users: 49,463 from 157 countries

Page Views: 529,138

SERVICE

DEPARTMENT

Faculty Advisor to Chi Chi Sigma (College Chemistry Society)	2021-present
Faculty Co-advisor to Chemistry Graduate Student Association (CGSA)	2020–2022
Faculty Advisor to the MSU Esports	2020–2022
Undergraduate Curriculum Committee Member	2020-present
Distance Education Committee Member	2020-present
Department of Chemistry Website Manager	2018–2020
Undergraduate Curriculum Development	2017-present
Faculty Advisor to the MSU SMACS	2016-present

COLLEGE

College of Arts & Sciences Scholarship Committee

2018-2021

PROFESSIONAL

National Chemistry Week Coordinator (MS Local ACS Section)

2019-present

AUTHORED CURRICULUM

Investigations in Chemistry I CH-1211 Lab Manual, 1st ed. T. Brown, Dornshuld, E. V., W. Nettles, and C. E. Webster Investigations in Chemistry II CH-1221 Lab Manual, 2nd ed. T. Brown, Dornshuld, E. V., W. Nettles, and C. E. Webster CH-1213 Chemistry I Coursebook, 5th ed. Dornshuld, E. V., W. Nettles, and C. E. Webster CH-1223 Chemistry II Coursebook, 6th ed. Dornshuld, E. V., W. Nettles, and C. E. Webster	2022
Investigations in Chemistry II CH-1221 Lab Manual, 1st ed. T. Brown, Dornshuld, E. V., W. Nettles, and C. E. Webster	2021
CH-1213 Chemistry I Coursebook, 4th ed.	
Dornshuld, E. V., W. Nettles, and C. E. Webster	
CH-1223 Chemistry II Coursebook, 5th ed.	
Dornshuld, E. V., W. Nettles, and C. E. Webster	2020
CH-1213 Chemistry I Workbook, 3rd ed. Dornshuld, E. V., W. Nettles, and C. E. Webster	2020
CH-1223 Chemistry II Workbook, 4th ed.	
Dornshuld, E. V., W. Nettles, and C. E. Webster	
CH-1213 Chemistry I Workbook, 2nd ed.	2019
Dornshuld, E. V., W. Nettles, and C. E. Webster	
CH-1223 Chemistry II Workbook, 3rd ed.	
Dornshuld, E. V., W. Nettles, and C. E. Webster	
CH-1213 Chemistry I Workbook, 1st ed.	2018
Dornshuld, E. V., W. Nettles, and C. E. Webster	
CH-1223 Chemistry II Workbook, 2nd ed.	
Dornshuld, E. V., W. Nettles, and C. E. Webster	0017
CH-1223 Chemistry II Workbook, 1st ed.	2017
Dornshuld, E. V.	

ORAL PRESENTATIONS

- 1. Dornshuld, E. V., M. Zhang, T. K. Hollis, and C. E. Webster. Characterizing ¹⁹⁵ Pt NMR chemical shift with computationally tractable non-relativistic density functional approaches. *69th Southeastern Regional Meeting of the American Chemical Society*, Charlotte, NC, November 7–11, 2017.
- 2. Dornshuld, E. V. Computational Chemistry: A Primer (or what you should know when you want have to talk to a computational chemist). *Feeding and Powering the World*, University, MS, June 19, 2017.
- 3. Dornshuld, E. V., M. Zhang, T. K. Hollis, and C. E. Webster. Utilizing computational ¹⁹⁵Pt NMR chemical shifts for the prediction of meridional ligand donor ability in square planar complexes. *Feeding and Powering the World 2016: Building the Network*, University, MS, July 25–26, 2016
- 4. Dornshuld, E. V., R. A. Vergenz, R. Mourad, M. A. Carrasquillo, J. W. Vickers, and H. F. Schaefer. Mechanism for aqueous glycine condensation. *2008 ACS 236th National Meeting*, Philadelphia, PA, August 17–20, 2008.

WORKSHOPS

1. Telluride School on Theoretical Chemistry (TSTC)

July 15-20, 2013

POSTER PRESENTATIONS

1. Dornshuld, E. V., M. Zhang, X. Zhang, T. K. Hollis, and C. E. Webster. Predicting ¹⁹⁵Pt NMR chemical shifts in small Pt(II) and Pt(IV) organometallic compounds with density functional approaches. *67th Southeast/71st Southwest Joint Regional Meeting of the American Chemical Society*, Memphis, TN, November 4–7, 2015.

- 2. Dornshuld, E. V., C. M. Holy, and G. S. Tschumper. New insight into $n \to \pi^*$ non-covalent interactions from two simple systems, formaldehyde dimer and thioformaldehyde dimer. *Mississippi Experimental Program to Stimulate Competitive Research State Meeting (EPSCoR)*, April 1, 2014.
- 3. Dornshuld, E. V., C. M. Holy, and G. S. Tschumper. New insight into $n \to \pi^*$ non-covalent interactions from two simple systems, formaldehyde dimer and thioformaldehyde dimer. *25th Austin Symposium on Molecular Structure and Dynamics (ASMD@D)*, March 1–4, 2014.
- Dornshuld, E. V. and G. S. Tschumper. Characterization of the potential energy surfaces of the P₂ dimer and the PCCP dimer. Southeast Theoretical Chemistry Association Annual Meeting (SETCA 2013), May 9–11, 2013.
- 5. Dornshuld, E. V. and G. S. Tschumper. Characterization of the potential energy surfaces of the P₂ dimer and the PCCP dimer. *Mississippi Experimental Program to Stimulate Competitive Research State Meeting (EPSCoR)*, April 18, 2013.
- 6. Dornshuld, E. V. and G. S. Tschumper. Characterization of the potential energy surfaces of the P₂ dimer and the PCCP dimer. *53rd Sanibel Symposium*, February 17–22, 2013.
- 7. Dornshuld, E. V. and G. S. Tschumper. Characterization of the potential energy surfaces of the P₂ dimer and the PCCP dimer. *Mississippi Experimental Program to Stimulate Competitive Research Fall Forum (EPSCoR)*, August 28, 2012.
- 8. Dornshuld, E. V. and G. S. Tschumper. Characterization of the potential energy surfaces of the P₂ and PCCP dimers. *Southeast Theoretical Chemistry Association Annual Meeting (SETCA 2012)*, May 17–19, 2012.
- 9. Dornshuld, E. V. and G. S. Tschumper. Characterization of the potential energy surfaces of the P₂ and PCCP dimers. *Mississippi Experimental Program to Stimulate Competitive Research State Meeting*, April 10, 2012.
- 10. Dornshuld, E. V. and G. S. Tschumper. Characterization of the potential energy surfaces of the P₂ and PCCP dimers. *Mississippi Experimental Program to Stimulate Competitive Research Fall Forum (EPSCoR)*, September 20, 2011.
- 11. Dornshuld, E. V. and G. S. Tschumper. The molecular structures of the P₂ and PCCP dimers. *Southeast Theoretical Chemistry Association Annual Meeting (SETCA 2011)*, May 13–14, 2011.
- 12. Dornshuld, E. V. and G. S. Tschumper. The molecular structures of the P₂ and PCCP dimers. *Mississippi Experimental Program to Stimulate Competitive Research State Meeting (EPSCoR)*, April 14, 2011.
- 13. Dornshuld, E. V., R. A. Vergenz, H. F. Schaefer, III, and G. S. Tschumper. Exploring the stepwise mechanism of aqueous glycine condensation. *19th Conference on Current Trends in Computational Chemistry (CCTCC 2010)*, October 29–30, 2010.
- 14. Dornshuld, E. V., R. A. Vergenz, H. F. Schaefer, III, and G. S. Tschumper. Exploring the mechanism for aqueous glycine condensation. *Mississippi Experimental Program to Stimulate Competitive Research State Meeting (EPSCoR)*, April 15, 2010.
- 15. Dornshuld, E. V., R. A. Vergenz, and R. Mourad. Mechanism for aqueous glycine condensation. *48th Sanibel Symposium*, February 21–26, 2008.
- 16. Bruno, P., M. Carrasquillo, N. Durtshi, E. V. Dornshuld, and J. Vickers. Do weak methyl-donated hydrogen bonds affect protein folding and stability? *Colorado Protein Stability Conference*, July 19, 2007.

CONTRIBUTED PRESENTATIONS

- 1. Olive, L., E. V. Dornshuld, and C. E. Webster. Curious case of DMSO: A computational study. *52nd Southeastern Undergraduate Research Conference*, Tuscaloosa, AL, January 25, 2020.
- 2. Olive, L., E. V. Dornshuld, and C. E. Webster. Curious case of DMSO: A computational study. *258th ACS National Conference and Exposition*, San Diego, CA, August 25–29 2019.
- 3. Olive, L., E. V. Dornshuld, and C. E. Webster. New insights into assessing the performance of DFT energetics on small oxygen/sulfur containing compounds. *Summer Undergraduate Research Symposium*, Mississippi State, MS, August 2019.

- 4. Olive, L., E. V. Dornshuld, and C. E. Webster. Assessing the performance of DFT energetics on small oxygen/sulfur containing compounds. *Feeding and Powering the World 2019: The Next Generation*, University, MS, July 15–16, 2019.
- 5. Frey, N. C., E. V. Dornshuld, F. Aghabozorgi, and C. E. Webster. Characterizing the fluxional behavior in (TMCOT)M(CO)₃ and (COT)Cr(CO)₃ complexes with computational approaches. *Feeding and Powering the World 2019: The Next Generation*, University, MS, July 15–16, 2019.
- 6. Frey, N. C., E. V. Dornshuld, F. Aghabozorgi, and C. E. Webster. Examining the electronic properties of twisted pyrene compounds. *Feeding and Powering the World 2019: The Next Generation*, University, MS, July 15–16, 2019.
- Autry, S., M. Zhang, E. V. Dornshuld, T. K. Hollis, and C. E. Webster. Probing the effects of environment on novel CCC-NHC-Pt(II) pincer complexes. 257th ACS National Meeting and Exposition, Orlando, FL Mar. 31 – Apr. 4, 2019.
- 8. Zhang, M., E. V. Dornshuld, J. C. Bunquin, M. Delferro, T. K. Hollis, and C. E. Webster. The platinum electronic parameter (PtEP): A compliment to TEP. *Feeding and Powering the World 2018: Planning for the Future*, University, MS, July 16–17, 2018.
- 9. Adiraju, K., M. Zhang, E. V. Dornshuld, R. W. Lamb, G. Liang, C. E. Webster, and T. K. Hollis. State-of-the-art nitrogen reduction: Haber-Bosch to homogeneous back to heterogeneous. *Feeding and Powering the World 2018: Planning for the Future*, University, MS, July 16–17, 2018.
- Zhang, M., S. Autry, V. Dixit, E. V. Dornshuld, J. Denny, N. Hammer, C. E. Webster and T. K. Hollis. Synthesis, characterization, and photophysics of CCC-NHC pincer platinum complexes. 255th ACS National Meeting and Exposition, New Orleans, LA, March 18–22, 2018.
- 11. Dixit, V., E. V. Dornshuld, C. E. Webster, and T. K. Hollis. Theoretical study of substituted CCC-NHC palladium and platinum complexes for OLED applications. *255th ACS National Meeting and Exposition*, New Orleans, LA, March 18–22, 2018.
- 12. Hollis, T. K., C. E. Webster, M. Zhang, E. V. Dornshuld, V. Dixit, J. C. Bunquin, and M. Delferro. NHC pincer complex donor ability-PtEP (Platinum (Pt) electronic parameter): A donicity scale incorporating strictly meridional, tridentate ligands. *255th ACS National Meeting and Exposition*, New Orleans, LA, March 18–22, 2018.
- 13. Hollis, T. K., C. E. Webster, M. Zhang, E. V. Dornshuld, J. C. Bunquin, and M. Delferro. NHC pincer complex donor ability? PtEP (platinum (Pt) electronic parameter): A donicity scale incorporating strictly meridional tridentate ligands. *69th Southeastern Regional Meeting of the American Chemical Society*, Charlotte, NC, November 7–11, 2017.
- 14. V. Dixit, M. Zhang, E. V. Dornshuld, T. K. Hollis, C. E. Webster. Theoretical study of the effect of substituents on the optical spectra and ¹⁹⁵Pt NMR shift of CCC-NHC Pt(II) complexes. *69th Southeastern Regional Meeting of the American Chemical Society*, Charlotte, NC, November 7–11, 2017.
- 15. Zhang, M., E. V. Dornshuld, V. Dixit, J. A. Denny, C. E. Webster, and T. K. Hollis. Synthesis and characterization of CCC-NHC pincer platinum complexes. *Feeding and Powering the World 2016: Building the Knowledge Base*, University, MS, July 19–20, 2017.
- 16. Dixit, V. E. V. Dornshuld, and C. E. Webster. Substitution effects on the emission spectra of platinum CCC-NHC pincer complexes. *Feeding and Powering the World 2016: Building the Network*, University, MS, July 25–26, 2016.
- 17. Zhang, M., E. V. Dornshuld, X. Zhang, and C. E. Webster. Predicting metal complex reactivity for energy applications: Donicity of meridional tridentate ligands: ¹⁹⁵Pt NMR of CCC-NHC Pt pincer square planar complexes. *Feeding and Powering the World 2016: Building the Network*, University, MS, July 25–26, 2016.
- 18. Lamb, R. W., E. V. Dornshuld, and C. E. Webster. Computational investigation of linkage isomerization in sulfoxide-containing ruthenium complexes. *67th Southeast/71st Southwest Joint Regional Meeting of the American Chemical Society*, Memphis, TN, November 4–7, 2015.