Joshua J. Goings, Ph.D.

CONTACT	Yale University	Phone: (203) 432-0737
Information	Department of Chemistry	E-mail: josh.goings@yale.edu
	225 Prospect St, New Haven, CT 06520	Web: www.joshuagoings.com
Education	Ph.D. University of Washington, Seattle, WA	Theoretical Chemistry 2017
	B.S. Seattle Pacific University, Seattle, WA	Chemistry & Biochemistry 2012
Experience	Postdoctoral Research Associate Yale University (transferred 2018) University of Illinois at Urbana-Champaign Advisor: Prof. Sharon Hammes-Schiffer	2017 – present
	Graduate Research Assistant University of Washington Advisor: Prof. Xiaosong Li	2012 – 2017
	Undergraduate Researcher Seattle Pacific University Advisor: Prof. Daniel P. Schofield	2011 – 2012
	NSF REU Undergraduate Researcher University of Connecticut Advisor: Prof. Michael B. Smith	2010 – 2010

Publications

PEER-REVIEWED 26 Early Photocycle of Slr1694 Blue-Light Using Flavin Photoreceptor Unraveled through Adiabatic Excited-State Quantum Mechanical/Molecular Mechanical Dynamics

J. J. Goings, S. Hammes-Schiffer

J. Amer. Chem. Soc. 2019, Just accepted.

25 Proton-Coupled Electron Transfer Drives Long-Range Proton Translocation in **Bioinspired Systems**

E. Odella[†], B. L. Wadsworth[†], S. J. Mora[†], J. J. Goings[†], M. T. Huynh, D. Gust, T. A. Moore, G. F. Moore, S. Hammes-Schiffer, A. L. Moore J. Amer. Chem. Soc. 2019, 141, 14057-14061.

†Equal contribution.

24 The Chronus Quantum Software Package

D. B. Williams-Young, A. Petrone, S. Sun, T. F. Stetina, P. Lestrange, C. E. Hoyer, D. R. Nascimento, L. Koulias, A. Wildman, J. Kasper, J. J. Goings, F. Ding, A. E. DePrince III, E. F. Valeev, X. Li

WIREs Comput. Mol. Sci. 2019, e1436.

23 Electron-Coupled Double Proton Transfer in the Slr1694 BLUF Photoreceptor: A **Multireference Electronic Structure Study**

E. R. Sayfutyarova, J. J. Goings, S. Hammes-Schiffer J. Phys. Chem. B. 2019, 123, 439-447.

22 Propensity for Proton Relay and Electrostatic Impact of Protein Reorganization in Slr1694 BLUF Photoreceptor

J. J. Goings, C. R. Reinhardt, S. Hammes-Schiffer J. Amer. Chem. Soc. 2018, 140, 15241-15251.

21 Controlling Proton-Coupled Electron Transfer in Bio-Inspired Artificial Photosynthetic Relays

E. Odella, S. J. Mora, B. L. Wadsworth, M. T. Huynh, J. J. Goings, P. A. Liddell, T. L. Groy, M. Gervaldo, L. E. Sereno, D. Gust, T. A. Moore, G. F. Moore, S. Hammes-Schiffer, A. L. Moore

J. Amer. Chem. Soc. 2018, 140, 15450-15460.

20 Orientation-Dependent Imaging of Electronically Excited Quantum Dots*

D. Nguyen, J. J. Goings, H. A. Nguyen, J. Lyding X. Li, M. Gruebele J. Chem. Phys., 2018, 148, 064701.

* 2018 JCP Editors' Choice

19 Current Development of Noncollinear Electronic Structure Theory

J. J. Goings, F. Egidi, X. Li

Int. J. Quantum Chem. 2018, 118, e25398.

18 Real-Time Time-Dependent Electronic Structure Theory

J. J. Goings, P. J. Lestrange, X. Li WIREs Comput. Mol. Sci. 2017, e1341.

17 Two-Component Non-Collinear Time-Dependent Spin Density Functional Theory for Excited State Calculations

F. Egidi, S. Sun, J. J. Goings, G. Scalmani, M. J. Frisch, X. Li *J. Chem. Theory Comput.* 2017, 13, 2591–2603.

16 Can Quantized Vibrational Effects Be Obtained from Ehrenfest Mixed Quantum-Classical Dynamics?

J. J. Goings, D. B. Lingerfelt, X. Li *J. Phys. Chem. Lett.* 2016, *7*, 5193–5197.

15 Accelerating Real-Time Time-Dependent Density Functional Theory with a Non-Recursive Chebyshev Expansion of the Quantum Propagator

D. Williams-Young, J. J. Goings, X. Li

J. Chem. Theory Comput. 2016, 12, 5333-5338.

14 Quantum Confinement Effects on Optical Transitions in Nanodiamonds Containing Nitrogen Vacancies

A. Petrone, J. J. Goings, X. Li *Phys. Rev. B.* 2016, *94*, 165402.

13 Real Time Propagation of the Exact Two Component Time-Dependent Density Functional Theory

J. J. Goings, J. M. Kasper, F. Egidi, S. Sun, X. Li *J. Chem. Phys.* 2016, *145*, 104107.

12 A Direct Atomic-Orbital Based Relativistic Two-Component Linear Response Method for Calculating Excited State Fine Structures

F. Egidi, J. J. Goings, M. J. Frisch, X. Li

J. Chem. Theory Comput. 2016, 12, 3711-3718.

11 An Atomic Orbital Based Real-Time Time-Dependent Density Functional Theory for Computing Electronic Circular Dichroism Band Spectra

J. J. Goings, X. Li

7. Chem. Phys. 2016, 144, 234102.

10 Imaging Excited Orbitals of Quantum Dots: Experiment and Electronic Structure Theory

L. Nienhaus † , J. J. Goings † , D. Nguyen, S. Wieghold, J. Lyding, X. Li, M. Gruebele J. Amer. Chem. Soc. 2015, 137, 14743–14750.

†Equal contribution.

9 Approximate Singly Excited States from a Two-Component Hartree-Fock Reference

J. J. Goings, F. Ding, E. R. Davidson, X. Li

J. Chem. Phys. 2015, 143, 144106.

8 Ab Initio Two-Component Ehrenfest Dynamics

F. Ding, J. J. Goings, H. Liu, D. Lingerfelt, X. Li

J. Chem. Phys. 2015, 143, 114105.

7 Energy-Specific Equation-of-Motion Coupled-Cluster Methods for High-Energy Excited States: Application to K-Edge X-ray Absorption Spectroscopy

B. Peng, P. J. Lestrange, J. J. Goings, M. Caricato, X. Li

J. Chem. Theory Comput. 2015, 11, 4146-4153.

6 Stability of the Complex Generalized Hartree-Fock Equations

J. J. Goings, F. Ding., M. J. Frisch, X. Li

J. Chem. Phys. 2015, 142, 154109.

5 Ab Initio Non-Relativistic Spin Dynamics

F. Ding, J. J. Goings, M. J. Frisch, X. Li

J. Chem. Phys. 2014, 141, 214111.

4 Theoretical Characterization of Conduction-Band Electrons in Photodoped and Aluminum-Doped Zinc Oxide (AZO) Quantum Dots

J. J. Goings, A. M. Schimpf, J. W. May, R. W. Johns, D. R. Gamelin, X. Li

J. Phys. Chem. C. 2014, 118, 26584-26590.

3 Assessment of Low-Scaling Approximations to the Equation of Motion Coupled-Cluster Singles and Doubles Equations

J. J. Goings, M. Caricato, M. Frisch, X. Li

J. Chem. Phys. 2014, 141, 164116.

2 Sorption of H₂ to Open Metal Sites in a Metal Organic Framework: A Symmetry Adapted Perturbation Analysis

J. J. Goings, S. Ohlsen, K. Blaisdell, D. Schofield

J. Phys. Chem. A. 2014, 118, 7411-7417.

1 Self-Consistent-Field Using Direct Inversion in Iterative Subspace Method and Quasi-Newton Vectors

J. J. Goings, F. Ding, X. Li

Adv. Quantum Chem. 2014, 68, 77-86.

Awards

Chemical Computing Group (CCG) Excellence Award		
National Science Foundation (NSF) Graduate Research Fellow		
Bernard and Claudine Nist Endowed Fellowship in Chemistry		
Peter Salamon Award, Telluride Science Research Center		
Natt-Lingafelter Graduate Fellowship Award		
David T. & Christina L. Wong Scholarship in Chemistry		

ORAL PRESENTATIONS

- Impact of protein conformational changes and electrostatics on a BLUF photoreceptor.
 258th ACS National Meeting, San Diego, CA (2019)
- Engineering a redox-active proton wire with proton-coupled electron transfer. 258th ACS National Meeting, San Diego, CA (2019)
- Can quantized vibrational effects be obtained from Ehrenfest mixed quantum-classical dynamics?

253rd ACS National Meeting, San Francisco, CA (2017)

- Atomic-orbital based real-time TDDFT for circular dichroism spectroscopy.
 251st ACS National Meeting, San Diego, CA (2016)
- Theoretical investigation of magnetic exchange interactions in dilute magnetic semiconductor quantum dots induced by defects.
 Pacifichem 2015, Honolulu, HI (2015)
- Linear Response Complex Generalized Hartree-Fock.
 249th ACS National Meeting, Denver, CO (2015)
- Equation of motion formalism of second order many-body perturbation theory (EOM-MBPT2) and second-order approximate coupled-cluster (CC2).
 248th ACS National Meeting, San Francisco, CA (2014)
- Low-Scaling Approximations to the Equation of Motion Coupled-Cluster Singles and Doubles Equations.
 Excited States and Time-Dependent Electronic Structure Theory, Telluride, CO (2014)

Poster Presentations

- Propensity for Proton Relay and Impact of Protein Reorganization in Slr1694 BLUF Photoreceptor.
 Gordon Research Conference on Computational Chemistry, West Dover, VT (2018)
- Real Time Propagation of the Exact Relativistic Two-Component Equations.

 Theory and Applications of Computational Chemistry (TACC), Seattle, WA (2016)
- Recent advances in real-time TDDFT for the description of optical activity.
 Low-scaling and Unconventional Electronic Structure Techniques Conference (LUEST),
 Telluride, CO (2016)
- Towards a Real-Time Description of Magnetic Systems with Applications to Magnetic Circular Dichroism Spectroscopy.
 251st ACS National Meeting, San Diego, CA (2016)

TEACHING

Teaching Assistant, Chemistry, Yale University

Spring 2019

CHEM 426/526: Computational Chemistry and Biochemistry

- Designed a computational chemistry course for graduate and undergraduate students.
- Gave students hands-on guidance using computational chemistry software.
- Facilitated student access to Yale's high-performance computing center.
- · Performed computational "demos" at the end of each lecture; guest lectured as necessary.

Guest Lecturer, Chemistry, Seattle Pacific University

Spring 2016

CHM 2960: Introduction to Research Methods

- Lecture title: "Molecules and Light (or, Why you see what you see)."
- Introduction to computational spectroscopy for undergraduate chemistry majors.

Guest Lecturer, Chemistry, Seattle Pacific University

Spring 2015

CHM 2960: Introduction to Research Methods

- Lecture title: "Explorations into Molecular Magnetism."
- Introduction to computational chemistry research with undergraduate chemistry majors.

Teaching Assistant, Chemistry, University of Washington

Spring 2014

- CHEM 465/565 Computational Chemistry
- Gave students hands-on guidance using computational chemistry software.
- Guided groups to perform novel computational research projects.

Teaching Assistant, Chemistry, University of Washington

Summer 2013

CHEM 162 General Chemistry

· Taught lab and quiz section to undergraduates.

Teaching Assistant, Chemistry, University of Washington

Winter 2013

CHEM 152: General Chemistry

• Taught two lab and two quiz sections to undergraduates.

Teaching Assistant, Chemistry, University of Washington CHEM 142: General Chemistry Autumn 2012

• Taught two lab and two quiz sections to undergraduates.

OUTREACH

Volunteer, UW Science Explorers

2014 - 2015

Sanislo Elementary, Seattle, WA

- After-school science program for 2nd through 5th grade elementary students.
- Worked one-on-one with students on science projects to encourage interest in STEM.
- Sanislo is the most racially diverse school in WA, according to Seattle Times (Jul 18, 2016).

Technical Blogger, www.joshuagoings.com

2013 - present

- Share code snippets, notes, and derivations with the theoretical chemistry community.
- Receives several hundred unique visitors each month.