

# Ruibin Liang

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## Professional Experience

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Assistant Professor 2020-present  
Department of Chemistry and Biochemistry  
**Texas Tech University**, Lubbock, TX

Postdoctoral Scholar 2017-2020  
Department of Chemistry, **Stanford University**, Stanford, CA  
Research advisor: **Professor Todd J. Martínez**

Postdoctoral Scholar 2016-2017  
Department of Chemistry, **University of California, Berkeley**, Berkeley, CA  
Research advisor: **Professor William H. Miller**

## Education

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Ph.D. in Theoretical and Computational Chemistry 2016  
**The University of Chicago**, Chicago, IL  
Research advisor: **Professor Gregory A. Voth**

M.S. in Theoretical and Computational Chemistry 2012  
**The University of Chicago**, Chicago, IL

B.S. in Chemical Biology 2011  
**Tsinghua University**, Beijing, China

## Honors and Awards

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Chemical Computing Group Excellence Award 2015  
American Chemical Society National Meeting

The Windt Memorial Fund Graduate Fellowship 2015  
The University of Chicago

Outstanding Young Researcher Award 2015  
From Computational Biophysics to Systems Biology” (CBSB) conference

Zongcheng Zheng Academic Excellent Scholarship 2010  
Tsinghua University

Wang Yong Academic Excellent Scholarship 2009

Tsinghua University

Zengjue Sun Academic Excellent Scholarship

2008

Tsinghua University

## **Research Experience**

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Assistant Professor

2020-present

Department of Chemistry and Biochemistry, **Texas Tech University**, Lubbock, TX

- Developed and applied multiscale simulation methods to study the biochemical processes involved in photoactive biological systems. Research projects include molecular photoswitches as therapeutics in photopharmacology, and organic molecules with aggregation-induced-emission characteristics as bioimaging probes for disease diagnosis.
- Applied *ab initio* simulation to study the enzymatic mechanism of deoxyribonuclease, in collaboration with Prof. Roger B. Sutton at TTUHSC.
- Applied molecular dynamics simulation to study the functional mechanism of melibiose transporter, in collaboration with Prof. Lan Guan at TTUHSC.

Postdoctoral Scholar with **Professor Todd J. Martínez**

2017-2020

Department of Chemistry, **Stanford University**, Stanford CA

- Applied restricted-spin ensemble Kohn-Sham (REKS) method in combination with *ab initio* multiple spawning (AIMS) method to simulate the photodynamics of channelrhodopsin 2 and bacteriorhodopsin.
- Applied quantum mechanics/molecular mechanics (QM/MM) simulation to study hydrogen bond structures in ketosteroid isomerase, resulting in a successful collaboration with the experimentalist Prof. Daniel Herschlag.

Postdoctoral Scholar with **Professor William H. Miller**

2016-2017

Department of Chemistry, **University of California, Berkeley**, Berkeley, CA

- Developed the symmetrical quasi-classical windowing model applied to the Meyer-Miller Hamiltonian (the SQC/MM simulation method) to investigate the photochemical processes in complex conjugated systems, such as ultrafast exciton transfer in organic semiconductors.

Graduate student with **Professor Gregory A. Voth**

2012-2016

Department of Chemistry, **The University of Chicago**, Chicago, IL

- Developed new multiscale reactive molecular dynamics methods to study proton transport mechanism in influenza A virus M2 channel and cytochrome c oxidase, resulting in many successful collaborations with several experimentalists, such as Prof. William F. DeGrado, Prof. Mei Hong and Prof. Mårten Wikström.
- Identified the inaccuracy of the self-consistent charge density functional tight binding method (SCC-DFTB) for simulating hydroxide and proton transport in aqueous and biological systems, which led to improvements of the method.

## **Teaching and Mentoring Experience**

Assistant Professor 2020-present

Department of Chemistry and Biochemistry, **Texas Tech University**, Lubbock, TX

- Taught the course CHEM3310 “Introduction to Biochemistry” for undergraduate students.
- Mentored graduate students’ research project in my research group.

Teaching Assistant for General Chemistry 2011-12

Department of Chemistry, **The University of Chicago**, Chicago, IL

- Prepared and led discussion and laboratory sessions for 30 undergraduate students, graded homework assignments and exams.
- Mentored one Ph.D. student’s research, resulting in one publication on JACS.

Postdoctoral Scholar with **Professor Todd J. Martínez** 2017-2020

Department of Chemistry, **Stanford University**, Stanford CA

- Mentored two Ph.D. students’ research, resulting in three publications on JACS.

## **Presentations**

International Conference on Catalysis and Chemical Engineering 2022  
San Francisco, CA

*Effects of enzyme-ligand interactions on the photoisomerization of a light-regulated chemotherapeutic drug*  
**(Invited talk)**

Center for Membrane Protein Research 13th Annual Virtual Symposium 2021  
Texas Tech University Health Science Center, Lubbock, TX

*Towards understanding photoactive biomolecular systems through multiscale simulation*  
**(Invited talk)**

Purdue University, Department of Chemistry, West Lafayette, IN 2021  
*Towards understanding photoactive biomolecular systems through multiscale simulation*

**(Invited talk)**

American Chemical Society National Meeting, Atlanta, GA 2021  
*First-principles non-adiabatic dynamics simulation of azobenzene photodynamics in solutions*

**(Contributed talk)**

American Chemical Society National Meeting, Atlanta, GA 2021  
*Light activation mechanism of channelrhodopsin 2*

**(Contributed talk)**

Workshop “Principles of Light-Induced Charge Transfer for Optogenetics” 2021  
*Light activation mechanism of channelrhodopsin 2*

**(Contributed talk)**

Virtual Conference on Theoretical Chemistry 2020

*Light activation mechanism of channelrhodopsin 2*  
**(Contributed talk)**

Texas Tech University, Lubbock, TX 2020  
*Understanding the quantum mechanical effects in biochemical reactions with multiscale simulation*  
**(Invited talk)**

Vanderbilt University, Nashville, TN 2020  
*Understanding the quantum mechanical effects in biochemical reactions with multiscale simulation*  
**(Invited talk)**

Syracuse University, Syracuse, NY 2020  
*Understanding the quantum mechanical effects in biochemical reactions with multiscale simulation*  
**(Invited talk)**

Northeastern University, Boston, MA 2020  
*Understanding the quantum mechanical effects in biochemical reactions with multiscale simulation*  
**(Invited talk)**

University of Alabama, Tuscaloosa, AL 2020  
*Understanding the quantum mechanical effects in biochemical reactions with multiscale simulation*  
**(Invited talk)**

34th William S. Johnson Symposium, Stanford, CA 2019  
*Nonadiabatic photodynamics of retinal protonated Schiff base in channelrhodopsin 2*  
**(poster)**

Gordon Research Conferences on Proteins, Holderness, NH 2019  
*Nonadiabatic photodynamics of retinal protonated Schiff base in channelrhodopsin 2*  
**(Invited talk)**

The 29th International Conference on Photochemistry, Boulder, CO 2019  
*Nonadiabatic photodynamics of retinal protonated Schiff base in channelrhodopsin 2*  
**(contributed talk)**

Northern California Theoretical Chemistry, Berkeley, CA 2019  
*Nonadiabatic photodynamics of retinal protonated Schiff base in channelrhodopsin 2*  
**(poster)**

Institute of Theoretical Physics, Chinese Academy of Sciences, Beijing, China 2019  
*Molecular dynamics simulation of chemical reactions in proteins*  
**(Invited talk)**

University of Waterloo, Waterloo, ON, Canada 2018  
*Molecular dynamics simulation of chemical reactions in proteins*  
**(Invited talk)**

Tsinghua University, Beijing, China 2018  
*Molecular dynamics simulation of chemical reactions in biomolecules*  
**(Invited talk)**

West Coast Theoretical Chemistry Symposium, Stanford, CA 2018  
*Nonadiabatic photodynamics of retinal protonated Schiff base in channelrhodopsin 2*

(poster)

American Chemical Society National Meeting, Boston, MA 2015  
*Mechanism of Proton Permeation through Influenza A M2 Channel*

(poster)

Midwest Theoretical Chemistry Conference, Ann Arbor, MI 2015  
*Multiscale simulations reveal the proton pumping mechanism in cytochrome c oxidase*

(poster)

From Computational Biophysics to Systems Biology (CBSB), Oklahoma City, OK 2015  
*Mechanism of Proton Permeation through Influenza A M2 Channel*

(Invited talk)

Gordon Research Conferences on Protons & Membrane Reactions, Ventura, CA 2014  
*Mechanism of Proton Permeation through Influenza A M2 Channel*

(poster)

Midwest Theoretical Chemistry Conference, Evanston, IL 2014  
*Mechanism of Proton Permeation through Influenza A M2 Channel*

(poster)

## Technical Skills

Programming: Python, C/C++, Fortran, Mathematica, Matlab, Perl, Bash, GPU computing, Tensorflow

Software packages: Schrödinger, CP2K, TeraChem, CHARMM, AMBER, NAMD, LAMMPS, VMD, etc.

Computational chemistry methods: structure-based ligand design, homology modeling, docking simulation, molecular dynamics, free energy calculation, reactive force-field development, quantum chemistry calculation, quantum dynamics, non-adiabatic dynamics simulation, etc.

## Publications

#co-first author \*corresponding author

1. **Liang R\*** & Bakhtiiari A. Multiscale simulation unravels the light-regulated reversible inhibition of dihydrofolate reductase by phototrexate. (Submitted)
2. Katsube S, **Liang R,\*** Amin A, Hariharan P, & Guan L\* (2022) Molecular basis for the cation selectivity of Salmonella typhimurium melibiose permease. *J. Mol. Biol.*:167598.
3. **Liang R\*** & Bakhtiiari A (2022) Effects of Enzyme–Ligand Interactions on the Photoisomerization of a Light-Regulated Chemotherapeutic Drug. *J. Phys. Chem. B* 126:2382-2393.
4. **Liang R,\*** Das D, Bakhtiiari A (2021). Protein confinement fine-tunes the aggregation-induced emission in the human serum albumin. *Phys. Chem. Chem. Phys.* 2021, 23, 26263-26272.
5. **Liang R\*** (2021) First-principles non-adiabatic dynamics simulation of azobenzene photodynamics in solutions. *J. Chem. Theory Comput.* 17 (5), 3019-3030.
6. **Liang R,** Yu JK, Meisner J, Liu F, & Martínez TJ (2021) Electrostatic Control of Photoisomerization in Channelrhodopsin 2 *J. Am. Chem. Soc.* 143, 5425-5437.

7. **Liang R**, & Martínez TJ. Chemical modification fine tunes the photodynamics of retinal protonated Schiff-base in methanol solution. (In preparation)
8. Yu JK, Bannwarth C, **Liang R**, Hohenstein EG, Martínez TJ (2020) Nonadiabatic Dynamics Simulation of the Wavelength-Dependent Photochemistry of Azobenzene Excited to the  $\pi\pi^*$  and  $\pi\pi^*$  Excited States. *J. Am. Chem. Soc.* 142 (49), 20680-20690.
9. Pinney MM, Mokhtari D, Akiva E, Yabukarski F, Sanchez DM, **Liang R**, Doukov T, Martinez TJ, Babbitt PC, & Herschlag D. Parallel Molecular Mechanisms for Enzyme Temperature Adaptation. *Science*. 2021, 371, eaay2784.
10. Yu JK, **Liang R**, Liu F, & Martínez TJ (2019) First-Principles Characterization of the Elusive I Fluorescent State and the Structural Evolution of Retinal Protonated Schiff Base in Bacteriorhodopsin. *J. Am. Chem. Soc.* 141(45):18193-18203.
11. Watkins LC, **Liang R**, Swanson JMJ, DeGrado WF, & Voth GA (2019) Proton Induced Conformational and Hydration Dynamics in the Influenza A M2 Channel. *J. Am. Chem. Soc.* 141(29):11667-11676.
12. **Liang R**, Liu F, & Martínez TJ (2019) Nonadiabatic Photodynamics of Retinal Protonated Schiff Base in Channelrhodopsin 2. *J. Phys. Chem. Lett.* 10(11):2862-2868.
13. **Liang R**,# Cotton SJ,# Binder R,# Hegger R, Burghardt I, & Miller WH (2018) The symmetrical quasi-classical approach to electronically nonadiabatic dynamics applied to ultrafast exciton migration processes in semiconducting polymers. *J. Chem. Phys.* 149(4):044101. (**Editor's Pick, Cover Article**)
14. Cotton SJ, **Liang R**, & Miller WH (2017) On the adiabatic representation of Meyer-Miller electronic-nuclear dynamics. *J. Chem. Phys.* 147:064112. (**Editor's Choice**)
15. Pinney MM, Natarajan A, Yabukarski F, Sanchez DM, Liu F, **Liang R**, Doukov T, Schwans JP, Martinez TJ, & Herschlag D (2018) Structural Coupling Throughout the Active Site Hydrogen Bond Networks of Ketosteroid Isomerase and Photoactive Yellow Protein. *J. Am. Chem. Soc.* 140(31):9827-9843.
16. **Liang R**, Swanson JMJ, Wikström M, & Voth GA (2017) Understanding the essential proton-pumping kinetic gates and decoupling mutations in cytochrome c oxidase. *Proc. Natl. Acad. Sci. U. S. A.* 114:5924-5929.
17. **Liang R**, Swanson JMJ, Madsen JJ, Hong M, DeGrado WF, & Voth GA (2016) Acid activation mechanism of the influenza A M2 proton channel. *Proc. Natl. Acad. Sci. U. S. A.* 113:E6955-E6964.
18. **Liang R**, Swanson JMJ, Peng Y, Wikström M, & Voth GA (2016) Multiscale simulations reveal key features of the proton-pumping mechanism in cytochrome c oxidase. *Proc. Natl. Acad. Sci. U. S. A.* 113(27):7420-7425.
19. Lee S,# **Liang R**,# Voth GA, & Swanson JMJ (2016) Computationally Efficient Multiscale Reactive Molecular Dynamics to Describe Amino Acid Deprotonation in Proteins. *J. Chem. Theory Comput.* 12(2):879-891.
20. **Liang R**, Li H, Swanson JMJ, & Voth GA (2014) Multiscale simulation reveals a multifaceted mechanism of proton permeation through the influenza A M2 proton channel. *Proc. Natl. Acad. Sci. U. S. A.* 111(26):9396-9401.
21. **Liang R**, Swanson JMJ, & Voth GA (2014) Benchmark Study of the SCC-DFTB Approach for a Biomolecular Proton Channel. *J. Chem. Theory Comput.* 10(1):451-462.
22. Choi TH,# **Liang R**,# Maupin CM, & Voth GA (2013) Application of the SCC-DFTB Method to Hydroxide Water Clusters and Aqueous Hydroxide Solutions. *J. Phys. Chem. B* 117(17):5165-5179.