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CONTACT INFORMATION	<a href="#">Department of Chemistry</a> <a href="#">University of Louisville</a> Room 251, Chemistry Bld. 3230 S. Brook St. Louisville, KY USA, 40208	<i>Phone:</i> +1 (502) 852-0730 <i>Mobile:</i> +1 (209) 201-2140 <i>E-mail:</i> <a href="mailto:lee.thompson.1@louisville.edu">lee.thompson.1@louisville.edu</a> <i>Website:</i> <a href="http://www.lmthompsongroup.com">www.lmthompsongroup.com</a>
CURRENT POSITION	<b>Assistant Professor</b> <a href="#">Department of Chemistry, University of Louisville</a>	(Aug 2017 to present)
PREVIOUS EMPLOYMENT	<b>Postdoctoral Scholar</b> <a href="#">Laboratory of Prof. Hrant P. Hratchian,</a> <a href="#">Chemistry and Chemical Biology, University of California, Merced</a>	(Jan 2014 to May 2017)
EDUCATION	<b>Doctor of Philosophy in Chemistry (PhD)</b> <a href="#">Imperial College London</a> Supervisor: Prof. Michael J. Bearpark Co-Supervisor: Dr. Jasper J. van Thor Thesis Title: <a href="#">Molecular Vibrations and Chemical Reactivity in Complex Environments</a>	(Oct 2009 to Sep 2013)
	<b>Master of Chemistry (MChem)</b> <a href="#">University of Southampton</a> First Class Honours	(Oct 2004 to Jul 2008)
TEACHING EXPERIENCE	<b>Lecture Courses</b> <ul style="list-style-type: none"><li>• <b>University of Louisville:</b> CHEM 202 <i>General Chemistry II</i> (Fall 2017); CHEM 660 <i>Advanced Physical Chemistry</i> (Fall 2018)</li></ul> <b>Workshops</b> <ul style="list-style-type: none"><li>• <b>NSCCS Workshops:</b> <a href="#">Gaussian for Beginners</a> (20-21<sup>th</sup> Sep 2016); <a href="#">Gaussian for Beginners</a> (16-17<sup>th</sup> Sep 2015); <a href="#">Gaussian for Beginners</a> (17-18<sup>th</sup> Sep 2014); <a href="#">Applied Computational Chemistry</a> (23-24<sup>th</sup> Apr 2013); <a href="#">Gaussian for Beginners</a> (11-12<sup>th</sup> Sep 2012); <a href="#">Applied Computational Chemistry</a> (17-18<sup>th</sup> Apr 2012); <a href="#">Gaussian for Advanced Users</a> (15<sup>th</sup> Sep 2011); <a href="#">Gaussian for Beginners</a> (14<sup>th</sup> Sep 2011)</li><li>• <b>Gaussian Workshops:</b> <a href="#">New Delhi, India</a> (6-10<sup>th</sup> Jan 2014); <a href="#">Wroclaw, Poland</a> (24-28<sup>th</sup> Jun 2013); <a href="#">New Delhi, India</a> (17-21<sup>th</sup> Dec 2012); <a href="#">Santiago de Compostela, Spain</a> (11-15<sup>th</sup> Jul 2011)</li><li>• <b>Imperial College London:</b> <a href="#">Dynamics of Triatomic Systems Workshop - developed code and GUI for module</a> (2011-2012); <a href="#">Transition States Modelling Workshop</a> (2010-2012); <a href="#">Molecular Orbitals and Symmetry Workshop</a> (2009-2012); <a href="#">1st Year Physical Chemistry Lab</a> (16-26<sup>th</sup> Nov 2009)</li></ul>	
SKILLS	Foreign Language Skills: <ul style="list-style-type: none"><li>• Spanish (B2), French (A1)</li></ul> Computing Skills: <ul style="list-style-type: none"><li>• Fortran 77, Fortran 90-08, OpenMP, MATLAB, Mathematica, Maple, Perl, Python, AWK, SED, UNIX shell scripting, T<sub>E</sub>X (L<sup>A</sup>T<sub>E</sub>X, B<sub>I</sub>B<sub>T</sub>E<sub>X</sub>), Vim, Microsoft Office, Git, Gnuplot, Microsoft Windows, Apple OS X, Linux</li></ul>	
PROFESSIONAL MEMBERSHIP	<a href="#">American Chemical Society (ACS)</a> ; <a href="#">Royal Society of Chemistry (RSC)</a> – Associate Member (AMRSC); <a href="#">American Association for the Advancement of Science (AAAS)</a>	

## PUBLICATIONS

1. Angnes, R.A.; **Thompson, L. M.**; Mashuta, M. S.; Correia, C.R.; Hammond, G. B. Non-Covalent Substrate Directed Enantioselective Heck Desymmetrization of *cis*-Cyclohex-4-ene-1,2-diol: Synthesis of All *cis* Chiral 5-Aryl-cyclohex-3-ene-1,2-diols and Mechanistic Investigation, *Advanced Synthesis & Catalysis* **2018**, in press DOI:10.1002/adsc.201800785.
2. **Thompson, L. M.**; Jarrold, C. C.; Hratchian, H. P. Explaining the  $\text{MoVO}_4^-$  Photoelectron Spectrum: Rationalization of Geometric and Electronic Structure *Journal of Chemical Physics* **2017**, 146(10), 104301.
3. Abdullahi, M. H.; **Thompson, L. M.**; Bearpark, M. J.; Vinader, V.; Afarinkia, K. The Role of Substituents in Retro Diels-Alder Extrusion of  $\text{CO}_2$  from 2(H)-pyrone Cycloadducts *Tetrahedron* **2016**, 72(40), 6021–6024.
4. **Thompson, L. M.**; Hratchian, H. P. Natural Ionization Orbitals for Interpreting Electron Photodetachment Processes *Journal of Chemical Physics* **2016**, 144(20), 204117.
5. **Thompson, L. M.**; Hratchian, H. P.  $\text{MoNbO}_2^-$  Theoretical Photoelectron Spectra Accounting for Spin Contamination in Density Functional Theory *Journal of Physical Chemistry A* **2015**, 119(32), 8744–8751.
6. **Thompson, L. M.**; Hratchian, H. P. Second Derivatives for Approximate Spin Projection Methods *Journal of Chemical Physics* **2015**, 142(5), 054106.
7. **Thompson, L. M.**; Hratchian, H. P. Spin Projection with Double Hybrid Density Functional Theory *Journal of Chemical Physics* **2014**, 141(3), 034108.
8. **Thompson, L. M.**; Lasoroski A.; Champion, P. M.; Sage, J. T.; Frisch, M. J.; van Thor, J. J.; Bearpark, M. J. Analytical Harmonic Vibrational Frequencies for the Green Fluorescent Protein Computed with ONIOM: Chromophore Mode Character and its Response to Environment *Journal of Chemical Theory and Computation* **2014**, 10(2), 751–766.
9. Vreven, T.; **Thompson, L. M.**; Larkin, S. M.; Kirker, I.; Bearpark, M. J. Deconstructing the ONIOM Hessian: Investigating Method Combinations for Transition Structures *Journal of Chemical Theory and Computation* **2012**, 8(12), 4907–4914.
10. Van Thor, J. J.; Lincoln, C. N.; Kellner, B.; Bourdakos, K. N.; **Thompson L. M.**; Bearpark, M. J.; Champion, P. M.; Sage, J. T. Ultrafast vibrational dynamics of parallel excited state proton transfer reactions in the Green Fluorescent Protein *Vibrational Spectroscopy* **2012**, 61, 1–6.

## TALKS AND POSTERS

1. **Thompson, L. M.** *Methods for localized electronic structure in photocatalytic mechanisms* Contributed talk presented at 256 ACS National Meeting & Exposition, 23<sup>rd</sup> August 2018, Boston, USA
2. **Thompson, L. M.** *Computational Methodologies for Assignment of Transition-Metal-Oxide Cluster Species in Photoelectron Detachment Spectra* Invited talk presented at the Contemporary Computational Chemistry Symposium held during SERMACS 2017, 9<sup>th</sup> November 2017, Charlotte, USA
3. **Thompson, L. M.** *Exploring the Structure of Transition Metal Oxide Clusters* Invited talk presented at the Brown and Williamson Departmental Seminar, 27<sup>th</sup> January 2017, Louisville, USA

4. **Thompson, L. M.;** Hratchian, H. P. *Simulation of Transition Metal Oxide Cluster Photodetachment Spectra* (Abstract #M38) Poster presented at Theory and Application of Computational Chemistry, 29<sup>th</sup> August 2016, Seattle, USA
5. **Thompson, L. M.;** Hratchian, H. P. *Efficient Simulation of Transition Metal Cluster Photoelectron Spectra Using Approximate Projection* (Abstract #01758) Contributed talk presented at 99th Canadian Chemistry Conference & Exhibition, 9<sup>th</sup> June 2016, Halifax, Canada
6. **Thompson, L. M.;** Hratchian, H. P. *Efficient Modelling of Transition Metal Systems Using Approximate Projection: Development and Applications* (Abstract #141) Contributed talk presented at 251st ACS National Meeting & Exposition, 14<sup>th</sup> March 2016, San Diego, USA
7. **Thompson, L. M.;** Hratchian, H. P. *Transition Metal Oxide Clusters: Accounting for Spin Contamination* (Abstract #639) Contributed talk presented at 250th ACS National Meeting & Exposition, 20<sup>th</sup> August 2015, Boston, USA
8. **Thompson, L. M.;** Hratchian, H. P. *Resolving Multiple Spin Contaminants Using Approximate Projection* (Abstract #362) Contributed talk presented at 250th ACS National Meeting & Exposition, 19<sup>th</sup> August 2015, Boston, USA
9. **Thompson, L. M.;** Hratchian, H. P. *Metal oxide clusters: The need for spin pure states* (Abstract #610) Poster presented at 248th ACS National Meeting & Exposition, 13<sup>th</sup> August 2014, San Francisco, USA
10. **Thompson, L. M.;** Champion, P. M.; Sage, J. T.; Frisch, M. J.; van Thor, J. V.; Bearpark, M. J. *Analytical Harmonic Modes of GFP: Chromophore Response to Environment* (Abstract #48) Contributed talk presented at 247th ACS National Meeting & Exposition, 16<sup>th</sup> March 2014, Dallas, USA
11. **Thompson, L. M.** *Using the ONIOM Method in the Study of Large Molecules* Invited Talk presented at NSCCS Applied Computational Chemistry Workshop for Synthetic Chemists, 24<sup>th</sup> April 2014, London, UK
12. **Thompson, L. M.** *Vibrational Frequencies of Large Molecules Using the ONIOM method* Invited Talk presented at NSCCS Gaussian Workshop for Beginners, 12<sup>th</sup> September 2012, London, UK
13. **Thompson, L. M.;** Forester, A.; Vreven T.; Bearpark, M. J. *Computing and Analysing Vibrational Frequencies of Large Molecules Using the ONIOM method* Poster presented at Computational Molecular Science 2012, 25<sup>th</sup> June 2012, Cirencester, UK
14. **Thompson, L. M.;** Bearpark, M. J. *Vibrational Analysis of Photoactive Proteins using ONIOM* Contributed talk presented at Modelling of Biologically-Inspired Photoactive Systems, 31<sup>st</sup> March 2011, Marseilles, France
15. **Thompson, L. M.;** Forester, A.; Bearpark, M. J. *Vibrational Analysis of Photoactive Proteins using ONIOM* Poster presented at Computational Molecular Science 2010, 28<sup>th</sup> June 2010, Cirencester, UK