

John A. Keith

CONTACT INFORMATION	Department of Chemical and Petroleum Engineering (CHE) Swanson School of Engineering University of Pittsburgh 3700 O'Hara Street Pittsburgh, PA 15261 USA	Office: 804 Benedum Hall Telephone: (412) 624-7016 Fax: (412) 624-9639 E-mail: jakeith[at]pitt.edu URL: http://klic.pitt.edu
RESEARCH INTERESTS	Computational chemistry for catalysis and sustainability. Specific applications in solar fuels, solvation modeling, first principles quantum chemistry, and reactive potentials for modeling the nanoscale.	
EDUCATION	California Institute of Technology , Pasadena, CA USA Ph.D., Chemistry, September 2007 <ul style="list-style-type: none">• Dissertation: "Computational Insight into Homogeneous Organopalladium Catalysis"• Advisor: William A. Goddard, III Wesleyan University , Middletown, CT USA B.A., Chemistry with High Honors, May, 2001	
ACADEMIC EXPERIENCE	University of Pittsburgh , Pittsburgh, Pennsylvania USA Department of Chemical and Petroleum Engineering <ul style="list-style-type: none">• R. K. Mellon Faculty Fellow in Energy• Tenure-track Assistant Professor in CHE Princeton University , Princeton, New Jersey USA Department of Mechanical and Aerospace Engineering <ul style="list-style-type: none">• Associate Research Scholar• Instructor APC509: Methods and Concepts in Electronic Structure Theory	Sep. 2013 – present Sep. 2013 – present Nov. 2010 – Jul. 2013 Feb. 2013 – May. 2013
	Universität Ulm , Ulm, GERMANY Institut für Elektrochemie <ul style="list-style-type: none">• Alexander von Humboldt Postdoctoral Fellow	Oct. 2007 – Oct. 2010
HONORS AND AWARDS	<ul style="list-style-type: none">• NSF-CAREER Award• Journal of Materials Chemistry A - Emerging Investigator• Pittsburgh Business Times: Who's Who in Energy• insidehighered.com: Mover and Shaker• Alexander von Humboldt Postdoctoral Fellowship• Phi Beta Kappa• American Chemical Society: Connecticut Valley Regional Award• Bradley Prize for Outstanding Undergraduate Thesis in Chemistry• American Chemical Society Analytical Chemistry Award	2017 2017 2014 – 2016 2013 2008 – 2010 2001 2001 2001 2000
PATENTS	<ul style="list-style-type: none">• Peilin Liao, John A. Keith, Emily A. Carter. Reducing Overpotentials in Hematite Photoanode Photoelectrochemical Cells by Doping. Princeton Docket No. 12-2755-1. Filed 11/1/2012	
MEDIA	<ul style="list-style-type: none">• Pittsburgh WESA: Researcher Finds Possible Way to Make CO₂ Into Energy (3 Feb. 2015) http://bit.ly/1WfC9aU• Pittsburgh Business Times: Pitt researcher receives funding to study carbon dioxide recycling (24 Jul. 2015) http://bit.ly/1TTY48W	

- **Quoted in Science Magazine:** There's too much carbon dioxide in the air. Why not turn it back into fuel? (10 Sep. 2015) <http://bit.ly/1K1xrZQ>

UNIVERSITY OF
PITTSBURGH /
SWANSON SCHOOL
OF ENGINEERING
(SSoE) SERVICE

- Pitt STRIVE/SSoE Diversity Committee Representative for CHE Sep. 2016 – present
- Pitt CHE Graduate student recruiting coordinator. Sep. 2013 – present
Contributed to increasing Ph.D. student enrollment of underrepresented minority students. Recognized by Associate Dean of Diversity Affairs, Dr. Sylvanus Wosu
- Pitt SSoE Committee for updating graduate applications June 2016
- Coordinator for Pitt Classroom Connection (outreach program with the Pittsburgh Science and Technology Academy) Jan. 2015 – Jun. 2015

NATIONAL SERVICE

- Physical Chemistry Division Officer for ACS, Energy Subdivision Mar. 2016 – present
- Symposium organizer 2017 Fall ACS meeting, PHYS division: "Spectroscopic and Computational Insights into Solid/Liquid Interfaces for Energy Conversion" *Aug. 2017
- Symposium organizer 2017 Spring ACS meeting, COMP division: "State-of-the-Art Methods for Modeling Materials Chemistry" Apr. 2017
- Conference co-organizer and session coordinator, Midwest Theoretical Chemistry Conference, Pittsburgh, PA Jun. 2016
- Co-chair for AIChE 2015 annual meeting, Catalysis and Reaction Engineering Division: "Computational Catalysis", Salt Lake City, UT Nov. 2015
- Co-chair for AIChE 2015 annual meeting, Computational Molecular Science and Engineering Forum: "Recent Advances in Molecular Simulation Methods", Salt Lake City, UT. Nov. 2015
- Website/publicity co-chair and session organizer 24th North American Catalysis Society Meeting, Pittsburgh, PA. Jun. 2015
- Intel ISEF 2015 - Grand Award Judge May 2015
- Symposium organizer 2015 Spring ACS meeting, CATL division: "Theoretical and Experimental Synergies at the Frontiers of Renewable Energy Catalysis" Mar. 2015
- Co-chair for AIChE 2014 annual meeting, Catalysis and Reaction Engineering Division: "Computational Catalysis V", Atlanta, GA. Nov. 2014
- Co-chair for AIChE 2014 annual meeting, Computational Molecular Science and Engineering Forum: "Recent Advances in Molecular Simulation Methods I & 2", Atlanta, GA. Nov. 2014
- Co-chair for AIChE 2013 annual meeting, Catalysis and Reaction Engineering Division: "Applications of DFT+X in Catalysis II" San Francisco, CA. Nov. 2013

SCIENTIFIC
REVIEWING

AAAS: Sci. Rep.
 ACS: ACS Books; ACS Catal.; Chem. Rev.; Environ. Sci. Technol.; Ind. Eng. Chem. Res.; Inorg. Chem.; J. Am. Chem. Soc.; J. Chem. Theory Comput.; J. Org. Chem.; J. Phys. Chem. C; J. Phys. Chem. Lett; Organometallics
 AIP: J. Chem. Phys.
 APS: Phys. Rev. Lett.
 ECS: J. Electrochem. Soc.
 Elsevier: Acta Mater.; Catal. Today; Chem. Eng. Sci.; Chem. Phys.; Coord. Chem. Rev.; Electrochim. Acta; Electrochem. Commun; J. Catal.; J. Power Sources, Mater. Chem. Phys.; Surf. Sci.
 IOP: J. Phys. Condens. Matter
 NPG: Nature Commun.
 RSC: Catal. Sci. Technol.; Dalton Trans.; Energy Environ. Sci.; J. Mater. Chem. A; Nanoscale; Phys. Chem. Chem. Phys.
 Springer: Electrocatal.; J. Solid State Electrochem.; Theor. Chem. Acc.
 Wiley: Angew. Chem. Int. Ed.; Chem. Eur. J.; Eur. J. Org. Chem.
 Grant DOE ad hoc reviewer; NSF panelist; NSF ad hoc reviewer; ACS-PRF; Ken- proposals: tucky Science & Technology Corporation; NWO (Netherlands Organisation for Scientific Research)

TEACHING

University of Pittsburgh

- CHE 0400 Summer 2017
Reactive Process Engineering OMET:
Undergraduate pillar course in chemical engineering kinetics
- CHE 1017/2017 Spring 2016
Chemical Energy & Nature of the Chemical Bond OMET: 4.00/5.0
Elective course in applications of quantum mechanics in chemistry
- CHE 2101 Fall 2015
Fundamentals of Thermodynamics OMET: 3.44/5.0
Graduate-level chemical engineering thermodynamics & statistical mechanics
- CHE 1017/2017 Spring 2015
Chemical Energy & Nature of the Chemical Bond OMET: 4.27/5.0
Elective course in applications of quantum mechanics in chemistry
- CHE 2101 Fall 2014
Fundamentals of Thermodynamics OMET: 3.50/5.0
Graduate-level chemical engineering thermodynamics
- Center for Simulation and Modeling Workshop May 7 – 9, 2014
Three lectures in quantum chemistry

Princeton University

- APC509 – *Methods and Concepts in Electronic Structure Theory* Spring 2013
Graduate level course in applications of quantum mechanics in chemistry 4.1/5.0

FUNDING

Total external funding as of 03/20/2017: \$735,000

- Mascaro Center for Sustainable Innovation (Pitt) Jul. 2017 – Jun. 2018
Seed grant: Toward machine learning blueprints for greener chelants
Role: PI, Amount awarded: **\$50,000**, PI time: 0.0 month
- NSF (CBET-1653392) Feb. 2017 – Jan. 2022
CAREER: SusChEM: Unlocking local solvation environments for energetically efficient hydrogenations with quantum chemistry
Role: PI, Amount awarded: **\$500,000**, PI time: 1.0 month
- Naval Research Laboratory (BAA-N00173-04) Aug. 2016 – Jan. 2017
Quantifying the effect of solvation on anti-corrosion coatings
Role: PI, Amount awarded: **\$50,000**, PI time: 0.5 month
- Naval Research Laboratory (BAA-N00173-03) Oct. 2015 – Apr. 2016
Preventing Corrosion by Controlling Cathodic Reaction Kinetics
Role: PI, Amount awarded: **\$75,000**, PI time: 1 month
- ACS Petroleum Research Fund Jul. 2015 – Jun. 2017
Unraveling Heterocycle-Promoted Hydride Transfer Mechanisms for Energetically Efficient Fuel and Petrochemical Production
Role: PI, Amount awarded **\$110,000**, PI time: 0.4 month
- Pitt Central Research Development Fund (CRDF) Jul. 2015 – Jun. 2017
Towards a Robust and Efficient Computational Modeling Approach for Elucidating Fundamental Photocatalysis
Role: PI, Amount awarded **\$16,000**, PI time: 0 month

STUDENTS AT PITT

Postdocs

- Aude Marjolin (Pitt postdoc) Jan. 2014 – Oct. 2014
Currently program manager for Pittsburgh Quantum Institute
- Victor Oyeyemi (Pitt postdoc) Jul. 2014 – Jun. 2015
Currently data scientist for Bloomberg

Graduate students

- Yasemin Basdogan (Pitt CHE PhD student) Jan. 2016 – present
- Mitchell C. Groenenboom (Pitt CHE PhD student) Jan. 2014 – present
- Karthikeyan Saravanan (Pitt CHE PhD student) Jan. 2014 – present
- Nguyen Vo (Pitt CHE MS student, co-advised with Karl Johnson) Jan. 2015 – Apr. 2017
- Yaqun Zhu (Pitt CHE MS student) Jan. 2014 – Jul. 2015

Special project MS students

- Benjamin Carlson (Pitt CHE) Jan. 2017 – Apr. 2017

Undergraduate students

- Ethan Henderson (Pitt CHE) Jan. 2017 – present
- Angela Leo (Pitt CHE) Jan. 2017 – present
- Charles Hansen (Pitt CHE) Sep. 2016 – Apr. 2017

- Solomon Astley (Pitt Comp Sci.) Sep. 2015 – Jan. 2016
- Kelly Lynch (Pitt CHE) Sep. 2015 – Jan. 2016
- Yinan Kang (Pitt CHE/Economics) Sep. 2014 – Sep. 2015
- Jeffrey Carr (Pitt CHE) Jan. 2014 – Apr. 2015
- Rohith Amruthur (Pitt CHE) Jan. 2014 – Dec. 2014
- Bronson Lockwood (Pitt CHE 1097 student) Jan. 2014 – Apr. 2014
- Blaec Toncini (Pitt CHE 1097 student) Jan. 2014 – Apr. 2014

Visiting students

- Caelin Celani (summer REU student from Wash. & Jeff. college) Jun. 2017 – present
- James Dean (summer REU student from Winthrop University) Jun. 2015 – Jul. 2015
- Dinesh Sundaravadevelu Devarajan (undergraduate visitor)
Currently graduate student at Texas Technological University Jun. 2015 – Jul. 2015
- Eric Gottlieb (visiting CMU chemistry PhD student) Jan. 2014 – Jun. 2015
- Alyssa Shorak (high school student) Jun. 2015 – Sep. 2015
- Gina Wagner (summer REU student from Trine University) Jun. 2014 – Jul. 2014

INVITED
PRESENTATIONS
SINCE ARRIVING AT
PITT

- Philadelphia Conference in Theoretical Chemistry (PCTC), Philadelphia, PA *Aug. 2017.
- 254th ACS meeting; ENFL division session “Innovative Chemistry & Electrocatalysis for Low-Carbon Energy & Fuels: Discovery to Application”, Washington DC *Aug. 2017
- National Energy Technology Laboratory, Pittsburgh, PA Apr. 12, 2017
- Department of Materials Science and Engineering - Carnegie Mellon University Feb. 3, 2017
- Department of Chemistry and the Center for Photochemical Sciences - Bowling Green State University (BGSU) Jan. 18, 2017
- Graduate Chemical Engineering Seminar - Drexel University Nov. 11, 2016
- Army Research Laboratory - Theory department, Aberdeen, MD Oct. 11, 2016
- Pittsburgh-Cleveland Catalysis Society, Pittsburgh, PA Sep. 23, 2016
- 252th ACS meeting; CATL division session “Electrocatalysis for CO₂ reduction”, Philadelphia, PA (presented by Karthikeyan Saravanan) Aug. 24, 2016
- Exploring Chemical Space with Machine Learning and Quantum Mechanics CECAM workshop, ETHZ, Zurich, Switzerland Jun. 1, 2016
- 251th ACS National Meeting, ENFL division session “Application of Computational Chemistry for Fuel and Energy Production”, San Diego, CA Mar. 16, 2016
- 251th ACS National Meeting, CATL division session “Condensed Phase Catalysis Symposium”, San Diego, CA Mar. 15, 2016
- I²CNER International Workshop “CO₂ capture and utilization division”, Kyushu University, Fukuoka Japan Feb. 4, 2016
- I²CNER Annual Symposium “Computational solutions to fundamental problems in carbon-neutral energy research”, Kyushu University, Fukuoka Japan Feb. 1, 2016
- 2016 Electrochemistry Gordon Research Conference, Ventura, California Jan. 11, 2016

10. 250th ACS National Meeting, COMP division session “Calculating pK_as and Redox Potentials”, Boston, MA Aug. 18, 2015
9. LONI Institute/LA-SiGMA seminar, Louisiana State University, Baton Rouge, LA Apr. 1, 2015
8. ACS Central Regional Meeting 2014, Session: “The Science of CO₂ Capture in Energy Production”, Pittsburgh, PA Oct. 31, 2014
7. Pitt Center for Simulation and Modeling symposium: “Advancing Research Through High Performance Computing”, Pittsburgh, PA Oct. 15, 2014
6. 248th ACS National Meeting, ENFL division session “Applications of Theoretical Chemistry for Energy and Fuel Production”, San Francisco, CA Aug. 12, 2014
5. 248th ACS National Meeting, PHYS division session “Renewable Energy Generation at the Interface between Theory and Experiment”, San Francisco, CA Aug. 11, 2014
4. Pittsburgh-Cleveland Catalysis Society, Pittsburgh, PA (presented by Aude Marjolin) Jun. 02, 2014
3. 247th ACS National Meeting, ENFL division session on Innovations in Carbon Dioxide Capture, Storage, Conversion, and Utilization, Dallas, TX Mar. 16, 2014
2. Joint Center for Artificial Photosynthesis (JCAP), Pasadena, CA Oct. 18, 2013
1. 246th ACS National Meeting, PHYS division session on Solar Energy Conversion, Indianapolis, IN Sep. 08, 2013

CONTRIBUTED
PRESENTATIONS
SINCE ARRIVING AT
PITT

‘S’ - denotes student presentations

33. 25th North American Catalysis Society Meeting, Denver, CO; The mechanism of isobutylene polymerization: new insight into proton-catalyzed polymerizations from the growing string method (presented by Minh Nguyen Vo) Jun 5, 2017
S
32. Pittsburgh Cleveland Catalysis Society, Akron, OH; Accurate computational modeling of chemical reactions in polar solvents using cluster-continuum modeling (presented by Yasemin Basdogan) May 25, 2017
S
31. 253rd ACS meeting, San Francisco, CA; Deoptimizing the oxygen reduction reaction on doped amorphous TiO₂ coatings for corrosion inhibition Apr. 6, 2017
30. 16th Annual AIChE, San Francisco, CA; Alloy Catalyst Discovery Using Computational Alchemy (presented by Karthikeyan Saravanan) Nov. 18, 2016
S
29. 16th Annual AIChE, San Francisco, CA; First Principles Quantum Chemistry Calculations to Model CO₂ Electroreduction on SnO₂ Particles (presented by Karthikeyan Saravanan) Nov. 16, 2016
S
28. 16th Annual AIChE, San Francisco, CA; Neural Network and Reaxff Comparison for Au Properties (presented by Jacob Boes, CMU student) Nov. 15, 2016
S
27. 16th Annual AIChE, San Francisco, CA; Comparing the Effect of Counter Ions, Solvent Molecules, and Electron Correlation on Homogeneous Reaction Models (presented by Mitchell C. Groenenboom) Nov. 15, 2016
S

26. **S** Electrochemical Energy Symposium, Pittsburgh, PA; Deoptimizing the oxygen reduction reaction on doped amorphous TiO₂ surfaces (presented by Mitchell C. Groenenboom) Oct. 21, 2016
25. **S** 252th ACS meeting, Philadelphia, PA; Tailoring materials for electrocatalytic reduction of CO₂ (presented by Karthikeyan Saravanan) Aug. 24, 2016
24. **S** 252th ACS meeting, Philadelphia, PA; Deoptimizing the oxygen reduction reaction on doped amorphous TiO₂ surfaces (presented by Mitchell C. Groenenboom) Aug. 22, 2016
23. **S** PQI2016, Pittsburgh, PA; Traversing the chemical space - Alloy catalysts discovery using Alchemy (presented by Karthikeyan Saravanan) Apr. 20, 2016
22. **S** PQI2016, Pittsburgh, PA; Explicitly unraveling the roles of counter ions and solvent molecules; (Poster presented by Mitchell C. Groenenboom) Apr. 20, 2016
21. 15th Annual AIChE, Salt Lake City, UT; Liquid Mixtures Freezing at Room Temperature: More Insights into Crystallization and Applications of Poly(trimethylene glycol)/Water Mixtures Nov. 11, 2015
20. 15th Annual AIChE, Salt Lake City, UT; New Perspectives on Aqueous Phase Reaction Mechanisms with Ab Initio Molecular dynamics, Nudged-Elastic Band, and Wavefunction Theory-in-DFT Embedding Nov. 9, 2015
19. 15th Annual AIChE, Salt Lake City, UT; Coincidences and Insights into Molecular Heterocycles That Catalyze CO₂ Reduction with Low Overpotentials Nov. 8, 2015
18. **S** 228th Electro Chemical Society meeting, Phoenix, AZ; Exploring the non innocence of inorganic complex ligands in (photo)electrochemical CO₂ reduction (presented by Karthikeyan Saravanan) Oct 11, 2015
17. **S** Science2015 hosted by the Pittsburgh Quantum Institute, Pittsburgh, PA; Aqueous phase CO₂ reduction with sodium borohydride: An ab initio molecular dynamics and nudged-elastic band mechanistic study; (Poster presented by Mitchell C. Groenenboom) Oct. 8, 2015
16. **S** Catalysis in Energy Group Poster Fair, Pittsburgh, PA; Aqueous phase CO₂ reduction with sodium borohydride: An ab initio molecular dynamics and nudged-elastic band mechanistic study; (Poster presented by Mitchell C. Groenenboom) Aug. 11, 2015
15. **S** 24th North American Catalysis Society Meeting, Pittsburgh, PA; Nitrogen Enriched Nanocarbons as a Metal-Free Water Reducing Catalysts (presented by Eric Gottlieb) Jun. 19, 2015
14. **S** 24th North American Catalysis Society Meeting, Pittsburgh, PA; The Mechanism for C-H Borylation By Cu-Fe Heterobimetallic Catalysts (Poster); (presented by Yaqun Zhu) Jun. 17, 2015
13. **S** 24th North American Catalysis Society Meeting, Pittsburgh, PA; Pourbaix Diagrams of Ruthenium Chromophores Under CO₂ Reduction Conditions; (presented by Karthikeyan Saravanan) Jun. 16, 2015
12. **S** 24th North American Catalysis Society Meeting, Pittsburgh, PA; Unraveling the Electrochemical Reactivities of Aromatic N-Heterocycles with Quantum Chemistry; (presented by Mitchell C. Groenenboom) Jun. 16, 2015

11. 249th ACS National Meeting, Denver, CO; First-principles quantum chemical investigations on the selectivity of borohydride for carbon dioxide and bicarbonate reduction in protic conditions Mar. 25, 2015
10. 249th ACS National Meeting, Denver, CO; Aqueous phase CO₂ reduction with sodium borohydride: An ab initio molecular dynamics and nudged-elastic band mechanistic study (Poster); (presented by Mitchell C. Groenenboom) Mar. 24, 2015
9. 14th Annual AIChE, Atlanta, GA; Water-Induced Crystallization of Poly(trimethyleneglycol); (presented by Robert Enick) Nov. 19, 2014
8. 14th Annual AIChE, Atlanta, GA; Thermochemical Descriptors for Unraveling Molecular Promoted CO₂ Conversions Nov. 18, 2014
7. 14th Annual AIChE, Atlanta, GA; First Principles Quantum Chemical Modeling of Radium in Barite for Fracking Wastewater Remediation (Poster); (presented by Gina Wagner) Nov. 17, 2014
6. 14th Annual AIChE, Atlanta, GA; Benchmarking Modern Range Separated DFT Functionals and Ab Initio Wavefunction Theory-in-DFT Embedding for Computational Catalysis Applications Nov. 17, 2014
5. Catalysis in Energy Group meeting, Pittsburgh, PA; A combined AIMD/NEB mechanistic study of aqueous phase CO₂ reduction with sodium borohydride; (presented by Mitchell C. Groenenboom) Nov. 5, 2014
4. 2014 ACS Central Regional Meeting, Pittsburgh, PA; Unraveling mechanistic aspects of heterocycle-promoted CO₂ electroreduction with quantum chemistry Oct. 30, 2014
3. 2014 ACS Central Regional Meeting, Pittsburgh, PA; Investigations of nitrogen doping density in graphene and hydrogen adsorption by DFT; (presented by Eric Gottlieb) Oct. 30, 2014
2. 2014 ACS Central Regional Meeting, Pittsburgh, PA; Preventing corrosion by controlling cathodic reaction kinetics (Poster); (presented by Victor B. Oyeyemi) Oct. 29, 2014
1. 2014 ACS Central Regional Meeting, Pittsburgh, PA; Aqueous phase CO₂ reduction with sodium borohydride: An ab initio molecular dynamics and nudged-elastic band mechanistic study. (Poster); (presented by Mitchell C. Groenenboom) Oct. 29, 2014

PUBLICATIONS DONE AT PITT

- [1] Keith, J. A.; Carter, E. A. Theoretical Insights into Electrochemical CO₂ Reduction Mechanisms Catalyzed by Surface-Bound Nitrogen Heterocycles. *J. Phys. Chem. Lett.* **2013**, *4*, 4058–4063, DOI: 10.1021/jz4021519. **Correction** - DOI: 10.1021/acs.jpcllett.5b00170.
- [2] Muñoz-García, A. B.; Ritzmann, A. M.; Pavone, M.; Keith, J. A.; Carter, E. A. Oxygen Transport in Perovskite-Type Solid Oxide Fuel Cell Materials: Insights from Quantum Mechanics. *Acc. Chem. Res.* **2014**, *47*, 3340–3348, DOI: 10.1021/ar4003174.
- [3] Li, P.; Henkelman, G.; Keith, J. A.; Johnson, J. K. Elucidation of Aqueous Solvent-Mediated Hydrogen-Transfer Reactions by ab initio Molecular Dynamics and Nudged Elastic-Band Studies of NaBH₄ Hydrolysis. *J. Phys. Chem. C* **2014**, *118*, 21385–21399, DOI: 10.1021/jp507872d.
- [4] Marjolin, A.; Keith, J. A. Thermodynamic Descriptors for Molecules That Catalyze Efficient CO₂ Electroreductions. *ACS Catal.* **2015**, *150*, 1123–1130, DOI: 10.1021/cs501406j.
- [5] Grice, K. A.; Groenenboom, M. C.; Manuel, J. D. A.; Sovereign, M. A.; Keith, J. A. Examining the Selectivity of Borohydride for Carbon Dioxide and Bicarbonate Reduction in Protic Conditions. *Fuel* **2015**, *150*, 139–145, DOI: 10.1016/j.fuel.2015.02.007.
- [6] Parmelee, S. R.; Mazzacano, T. J.; Zhu, Y.; Mankad, N. P.; Keith, J. A. A Heterobimetallic Mechanism for C-H Borylation Elucidated from Experimental and Computational Data. *ACS Catal.* **2015**, *150*, 3689–3699, DOI: 10.1021/acscatal.5b00275.
- [7] Boes, J. R.; Groenenboom, M. C.; Keith, J. A.; Kitchin, J. R. Neural network and ReaxFF comparison for Au properties. *Int. J. Quantum Chem.* **2016**, *116*, 979–987, DOI: 10.1002/qua.25115.
- [8] Saravanan, K.; Keith, J. A. Standard redox potentials, pK_as, and hydricities of inorganic complexes under electrochemical conditions and implications for CO₂ reduction. *Dalton Trans.* **2016**, *45*, 15336–15341, DOI: 10.1039/C6DT02371A.
- [9] Groenenboom, M. C.; Saravanan, K.; Zhu, Y.; Carr, J. M.; Marjolin, A.; Faura, G. G.; Yu, E. C.; Dominey, R. N.; Keith, J. A. Structural and Substituent Group Effects on Multielectron Standard Reduction Potentials of Aromatic N-Heterocycles. *J. Phys. Chem. A* **2016**, *120*, 6888–6894, DOI: 10.1021/acs.jpca.6b07291.
- [11] Groenenboom, M. C.; Keith, J. A. Explicitly Unraveling the Roles of Counterions, Solvent Molecules, and Electron Correlation in Solution Phase Reaction Pathways. *J. Phys. Chem. B* **2016**, *120*, 10797–10807, DOI: 10.1021/acs.jpccb.6b07606.
- [10] Saravanan, K.; Gottlieb, E.; Keith, J. A. Nitrogen-doped Nanocarbon Materials Under Electroreduction Operating Conditions and Implications for Electrocatalysis. *Carbon* **2017**, *111*, 859–866, DOI: 10.1016/j.carbon.2016.10.084.
- [12] Gray, C. M.; Saravanan, K.; Wang, G.; Keith, J. A. Quantifying solvation energies at solid/liquid interfaces using continuum solvation methods. *Mol. Sim.* **2017**, *43*, 420–427, DOI: 10.1080/08927022.2016.1273525.
- [13] Saravanan, K.; Basdogan, Y.; Dean, J. R.; Keith, J. A. Computational investigation of CO₂ electroreduction on tin oxide and predictions of Ti, V, Nb and Zr dopants for improved catalysis. *J. Mater. Chem. A* **2017**, *5*, 11756–11763, DOI: 10.1039/C7TA00405B.

PUBLICATIONS DONE BEFORE ARRIVING AT PITT

- [1] Keith, J. A.; Oxgaard, J.; Goddard III, W. A. Inaccessibility of β -hydride Elimination From -OH Functional Groups in Wacker-type Oxidation. *J. Am. Chem. Soc.* **2006**, *128*, 3132–3133, DOI: 10.1021/ja0533139.
- [2] Keith, J. A.; Behenna, D. C.; Mohr, J. T.; Ma, S.; Marinescu, S. C.; Oxgaard, J.; Stoltz, B. M.; Goddard III, W. A. The Inner-sphere Process in the Enantioselective Tsuji Allylation Reaction with (*S*)-*t*-Bu-phosphinooxazoline Ligands. *J. Am. Chem. Soc.* **2007**, *129*, 11876–11877, DOI: 10.1021/ja070516j.
- [3] Keith, J. A.; Nielsen, R. J.; Oxgaard, J.; Goddard III, W. A. Unraveling the Wacker Oxidation Mechanisms. *J. Am. Chem. Soc.* **2007**, *129*, 12342–12343, DOI: 10.1021/ja072400t.
- [4] Lu, J.-Y.; Keith, J. A.; Shen, W.-Z.; Schürmann, M.; Preut, H.; Jacob, T.; Arndt, H.-D. Regioselective De Novo Synthesis of Cyanohydroxypyridines with a Concerted Cycloaddition Mechanism. *J. Am. Chem. Soc.* **2008**, *130*, 13219–13221, DOI: 10.1021/ja804078v.
- [5] Keith, J. A.; Henry, P. M. The mechanism of the Wacker reaction: a tale of two hydroxypalladations. *Angew. Chem. Int. Ed.* **2009**, *48*, 9038–9049, DOI: 10.1002/anie.200902194.
- [6] Keith, J. A.; Nielsen, R. J.; Oxgaard, J.; Goddard III, W. A.; Henry, P. M. Comment on "Mechanism and Kinetics of the Wacker Process: A Quantum Mechanical Approach". *Organometallics* **2009**, *28*, 1618–1619, DOI: 10.1021/om800013p.
- [7] Keith, J. A.; Jerkiewicz, G.; Jacob, T. Theoretical Investigations of the Oxygen Reduction Reaction on Pt(111). *ChemPhysChem* **2010**, *11*, 2779–2794, DOI: 10.1002/cphc.201000286.
- [8] Gao, W.; Keith, J. A.; Anton, J.; Jacob, T. Theoretical Elucidation of the Competitive Electrooxidation Mechanisms of Formic Acid on Pt (111). *J. Am. Chem. Soc.* **2010**, *132*, 18377–18385, DOI: 10.1021/ja1083317.
- [9] Anderson, B. J.; Keith, J. A.; Sigman, M. S. Experimental and Computational Study of a Direct O₂-coupled Wacker Oxidation: Water Dependence in the Absence of Cu Salts. *J. Am. Chem. Soc.* **2010**, *132*, DOI: 10.1021/ja1057218.
- [10] Keith, J. A.; Fantauzzi, D.; Jacob, T.; van Duin, A. C. Reactive Forcefield for Simulating Gold Surfaces and Nanoparticles. *Phys. Rev. B* **2010**, *81*, 235404, DOI: 10.1103/PhysRevB.81.235404.
- [11] Gao, W.; Keith, J. A.; Anton, J.; Jacob, T. Oxidation of Formic Acid on the Pt(111) Surface in the Gas Phase. *Dalton Trans.* **2010**, *39*, 8450–8456, DOI: 10.1039/C0DT00404A.
- [12] Keith, J. A.; Jacob, T. *Theory and Experiment in Electrocatalysis*; Springer New York, pp 89–132.
- [13] Keith, J. A.; Jacob, T. Atomic-level Elucidation of the Initial Stages of Self-assembled Monolayer Metallization and Nanoparticle Formation. *Chem. Eur. J.* **2010**, *16*, 12381–12386, DOI: 10.1002/chem.201001396.
- [14] Keith, J. A.; Anton, J.; Kaghazchi, P.; Jacob, T. Modeling Catalytic Reactions on Surfaces with Density Functional Theory. *Modeling and Simulation of Heterogeneous Catalytic Reactions* 1–38.
- [15] Kleiner, K.; Comas-Vives, A.; Naderian, M.; Mueller, J. E.; Fantauzzi, D.; Mesgar, M.; Keith, J. A.; Anton, J.; Jacob, T. Multiscale Modeling of Au-Island Ripening on Au(100). *Adv. Phys. Chem.* **2012**, *2011*, DOI: 10.1155/2011/252591.
- [16] Oyeyemi, V. B.; Keith, J. A.; Pavone, M.; Carter, E. A. Insufficient Hartree-Fock Exchange in Hybrid DFT Functionals Produces Bent Alkynyl Radical Structures. *J. Phys. Chem. Lett.* **2012**, DOI: 10.1021/jz201564g.

- [17] Keith, J. A.; Carter, E. A. Theoretical insights into pyridinium-based photoelectrocatalytic reduction of CO₂. *J. Am. Chem. Soc.* **2012**, *134*, 7580–7583, DOI: 10.1021/ja300128e.
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