Carlos H. Borca | Curriculum Vitæ

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Summary

- Pragmatic theoretical chemist that enjoys doing collaborative research on basic science, engaging real-life problems, and developing original scientific software solutions.
- Interested in developing theoretical and computational methods and applying them to problems of broad chemical scope, such as non-covalent interactions in organic crystals, pharmaceutical formulations, biomolecular structure, and materials design.
- o 10 years of research and teaching experience in academic institutions and national laboratories in the U.S.
- o 8 articles published in peer-reviewed scientific journals, 2 invited and 14 contributed oral presentations, 15 poster presentations.
- o More than \$25,000 in scholarships, travel grants, and awards.

Education

Postdoctoral Fellowship

Georgia Institute of Technology - School of Chemistry and Biochemistry - Atlanta, GA, USA	0/2011 1 1030110
Ph.D. in Chemistry Purdue University - Department of Chemistry - West Lafayette, IN, USA	8/2012 - 5/2017
Applied Management Principles Mini-MBA Purdue University - Krannert School of Management - West Lafayette, IN, USA	5/2016 - 6/2016
Professional (5-Year) Degree in Chemistry Honors mention for meritorious research thesis Universidad del Valle - School of Natural and Exact Sciences - Cali, Colombia	8/2004 - 11/2009
Experience	
Research	
Development of methods to compute properties of molecular crystals Adviser: Prof. C. David Sherrill Georgia Institute of Technology - Atlanta, GA, USA	8/2017 - Present
Development of theory to model molecular interactions through fragmentation schemes Adviser: Prof. Lyudmila V. Slipchenko Purdue University - West Lafayette, IN, USA	8/2012 - 5/2017
Modular implementation of the Tkatchenko-Scheffler model for van der Waals interactions Advisers: Dr. Alfredo A. Correa and Dr. Xavier I. Andrade Lawrence Livermore National Laboratory - Livermore, CA, USA	6/2015 - 8/2015
Molecular modeling of water absorbent materials at the nano-scale Adviser: Prof. Carlos A. Arango Universidad Icesi - Cali, Colombia	1/2011 - 7/2012
Computational study of interactions of single-wall carbon nanotubes/DNA hybrids and small molecules relevant to chemical sensing applications Adviser: Prof. Julio C. Arce Universidad del Valle - Cali, Colombia	8/2007 - 9/2009
Teaching.	

Graduate Teaching Assistant | Purdue University - West Lafayette, IN, USA

CHM 57900: Computational Chemistry - Spring 2015

CHM 11100: General Chemistry - Fall 2012, Fall 2013, Fall 2014

CHM 11500: General Chemistry - Spring 2013

8/2012 - 5/2015

8/2017 - Present

Physical Chemistry I - 2010-I, 2010-II, 2011-I Physical Chemistry II - 2010-II General Chemistry - 2009-II, 2010-I, 2010-II

Honors and Awards

National Science Foundation Travel Award to attend the CMU-GT Symposium on Machine Learning in Science and Engineering 5/2018 Carnegie Mellon University - Pittsburgh, PA, USA

Materials Computational Center Travel Award to attend the 7th TDDFT School and Workshop
University of Illinois - Urbana-Champaign, IL, USA

9/2016

Colombian Students Association at Purdue University (CSAP) Travel Grant Award

Purdue University - West Lafayette, IN, USA

8/2015

LLNL Students Poster Symposium Outstanding Accomplishment Award
Lawrence Livermore National Laboratory - Livermore, CA, USA

7/2015

Eli Lily Scholarship 6/2014

Purdue University - West Lafayette, IN, USA

Young Scientist and Innovator Scholarship of 2011 1/2011

Administrative Department of Science, Technology, and Innovation of the Colombian Government (Colciencias) and Universidad Icesi - Cali, Colombia

Peer-reviewed Articles

Published.

- Martín A. Mosquera, Carlos H. Borca, Mark A. Ratner, and George C. Schatz. Connection Between Hybrid Functionals and Importance of the Local Density Approximation. *The Journal of Physical Chemistry A*, 2016, 120 (9), pp 1605–1612. DOI: 10.1021/acs.jpca.5b10864
- Carlos H. Borca and Carlos A. Arango. Molecular Dynamics of a Water-absorbent Nano-scale Material Based on Chitosan. *The Journal of Physical Chemistry B*, **2016**, *120* (15), pp 3754–3765. DOI: 10.1021/acs.jpcb.5b11230 Featured in the cover art.



- Fulizi Xiong, **Carlos H. Borca**, Lyudmila V. Slipchenko, and Paul B. Shepson. Photochemical Degradation of Isoprene-derived 4,1-Nitrooxy Enal. *Atmospheric Chemistry and Physics*, **2016**, *16*, pp 5595–5610. DOI: 10.5194/acp-16-5595-2016
- Laura I. Mosquera-Giraldo, Carlos H. Borca, Xiangtao Meng, Kevin J. Edgar, Lyudmilla V. Slipchenko, and Lynne S. Taylor. Mechanistic Design of Chemically Diverse Polymers with Applications in Oral Drug Delivery. *Biomacromolecules*, 2016, 17 (11), pp 3659–3671. DOI: 10.1021/acs.biomac.6b01156
- Carlos H. Borca, Lyudmila V. Slipchenko, and Adam Wasserman. Ground-State Charge Transfer: Lithium-Benzene and the Role of Hartree-Fock Exchange. *The Journal of Physical Chemistry A*, **2016**, *120* (41) pp 8190–8198. DOI: 10.1021/acs.jpca.6b09014
- Na Li, Laura I. Mosquera-Giraldo, Carlos H. Borca, James D. Ormes, Michael Lowinger, John D. Higgins, Lyudmilla V. Slipchenko, and Lynne S. Taylor. A Comparison of the Crystallization Inhibition Properties of Bile Salts. Crystal Growth & Design, 2016, 16 (12) pp 7286–7300. DOI: 10.1021/acs.cgd.6b01470
- o Joel D. Rindelaub, **Carlos H. Borca**, Matt A. Hostetler, Mark A. Lipton, Lyudmila V. Slipchenko, and Paul B. Shepson. The Acid-Catalyzed Hydrolysis of an α -Pinene-Derived Organic Nitrate: Kinetics, Products, Reaction Mechanisms, and Atmospheric Impact. *Atmospheric Chemistry and Physics*, **2016**, 16, pp 15425–15432. DOI: 10.5194/acp-2016-726
- Sarah F. Tyler, Eileen C. Judkins, Dmitry Morozov, Carlos H. Borca, Lyudmila V. Slipchenko, and David R. McMillin. To Be or Not to Be Symmetric: That is the Question for Potentially Active Vibronic Modes. The Journal of Chemical Education, 2017, 94 (9), pp 1232–1237. DOI: 10.1021/acs.jchemed.7b00289

Under Review

• Naila A. Mugheirbi, Laura I. Mosquera-Giraldo, **Carlos H. Borca**, Lyudmila V. Slipchenko, and Lynne S. Taylor. Phase Behavior of Solid Dispersions Produced using Various Solvent Systems: Mechanistic Understanding of the Role of Polymer using Experimental and Theoretical Methods. *Under review*, **2018**.

In Preparation (Draft available)

- Laura I. Mosquera-Giraldo, **Carlos H. Borca**, Andrew S. Parker, Yifan Dong, Kevin J. Edgar, Stephen P. Beaudoin, Lyudmila V. Slipchenko, and Lynne S. Taylor. Crystallization Inhibition Properties of Cellulose Ester and Ethers for a Group of Chemically Diverse Drugs Experimental and Computational Insight. *Manuscript in Preparation*, **2018**.
- **Carlos H. Borca** and Lyudmila V. Slipchenko. Exploiting the Timescale Separation between Energy Contributions to Accelerate Molecular Dynamics in the Effective Fragment Potential. *Manuscript in Preparation*, **2018**.
- Carlos H. Borca, Brandon W. Bakr, Lori A. Burns, and C. David Sherrill. CrystaLattE: Automated Computation of Benchmark-level Lattice Energy of Molecular Crystals. *Manuscript in Preparation*, **2018**.

Scientific Events

Events Organized	
Second Annual Academic Event of the Colombian Student Association at Purdue University - West Lafayette, IN, USA Head of the Academic Event Organization Committee	10/2016
Invited Talks	
Academic Presentations of the Colombian Student Association at Purdue University - West Lafayette, IN, USA Developing Software to Model van der Waals Interactions in Materials	9/2015
Special Guest Talk at the Graduate Physical Chemistry Seminar at Universidad del Valle - Cali, Valle del Cauca, Colombia Charge Distribution in Carbon Nanopores Via Density Functional Theory	5/2014
Contributed Talks	
Meeting of the Southeastern Theoretical Chemistry Association (SETCA 2017) - Oxford, MS, USA CAM-LDAO: Reincarnating the Local Density Approximation	5/2017
Graduate Physical Chemistry Seminar - West Lafayette, IN, USA Molecular Dynamics with the Effective Fragment Potential Method	11/2016
252th National Meeting & Exposition of the American Chemical Society (ACS 2016) - Philadelphia, PA, USA • <i>CAM-LDAO: The Reincarnation of the Local Density Approximation</i>	8/2016
• Timescale Separation between Energy Contributions in the Effective Fragment Potential	
 Molecular Dynamics of Water-Absorbent Nanoscale Materials Based on Chitosan 	
48th Midwest Theoretical Chemistry Conference (MWTCC 2016) - Pittsburgh, PA, USA Exploiting the Timescale Separation between Energy Contributions to Accelerate Molecular Dynamics in the Effective Fragment Potential	6/2016
Academic Presentations of the Colombian Student Association at Purdue University - West Lafayette, IN, USA Molecular Dynamics of a Water-Absorbent Nanoscale Material Based on Chitosan	6/2016
250th National Meeting & Exposition of the American Chemical Society (ACS 2015) - Boston, MA, USA • Charge Transfer in the Lithium-Benzene Complex via Density Functional Theory	8/2015
• Developing materials-modeling software for electron dynamics with van der Waals interactions	
O Determining the Melting Point of Ice with the Effective Fragment Potential	
V National Meeting of Theoretical and Computational Chemists (V ENQTC) - Guatapé, Antioquia, Colombia Charge Distribution in Carbon Nanopores Via Density Functional Theory	5/2014
IX National Congress of Pure and Applied Chemistry Students (IX CONEQ) - Cali, Valle del Cauca, Colombia Computational Study of the Interactions between Carbon Nanotube/DNA Hybrids and Simple Molecules Relevant in Chemical Sensing	10/2009
III National Symposium of Nanotechnology (NANOCOLOMBIA 2009) - Bogotá, D.C., Colombia Electronic Properties of Chemical Transducers Based on Carbon Nanotubes Functionalized with Homo-DNA polynucleotides	4/2009
Posters	
49th Midwest Theoretical Chemistry Conference (MWTCC 2017) - East Lansing, MI, USA CAM-LDAO: Reincarnating the Local Density Approximation	6/2017
7th Time-Dependent Density-Functional Theory: Prospects and Applications (7th TDDFT) - Benasque, Aragón, Spain CAM-LDAO: The Reincarnation of the Local Density Approximation	9/2016
IX Congress of the International Society for Theoretical Chemical Physics (IX ISTCP 2016) - Grand Forks, ND, USA Exploring the Temporal Evolution of the Energy Components in the Effective Fragment Potential Molecular Dynamics	7/2016

2016 Conference on Excited State Processes (ESP 2016) - Santa Fe, NM, USA Charge Transfer in the Lithium-Benzene Complex: Understanding the Role of the Hartree-Fock Exchange	6/2016
First Annual Academic Event of the Colombian Student Association at Purdue University - West Lafayette, IN, USA Developing Materials-modeling Software for Electron Dynamics with van der Waals Interactions	11/2015
Lawrence Livermore National Laboratory Student Poster Symposium - Livermore, CA, USA Developing Materials-modeling Software for Electron Dynamics with van der Waals Interactions	7/2015
Meeting of the Southeastern Theoretical Chemistry Association (SETCA 2015) - Orlando, FL, USA Charge Transfer in Lithium-Benzene via Density Functional Theory	5/2015
248th National Meeting & Exposition of the American Chemical Society (ACS 2014) - San Francisco, CA, USA Charge Transfer in Lithium-Benzene via Density Functional Theory	8/2014
46th Midwest Theoretical Chemistry Conference (MWTCC 2014) - Evanston, IL, USA Charge Transfer in Lithium-Benzene via Density Functional Theory	7/2014
VIII National Meeting of Neuroscience - Bogotá, D.C., Colombia Computational Study of Glycosylation and Phosphorylation of Proteins Involved in Neurodegeneration	6/2012
IV National Meeting of Theoretical and Computational Chemists (IV ENQTC) - Cali, Valle del Cauca, Colombia Molecular Modelling of Water Absorbent Nanoscale Materials	5/2012
Fourth Research Socialization Day at Universidad Icesi 2011 - Cali, Valle del Cauca, Colombia Molecular Mechanics Study of Hydrogel-type Biopolymers at the Nanoscale	3/2011
III National Meeting of Theoretical and Computational Chemists (III ENQTC) - San Gil, Santander, Colombia Electronic Response of Chemical Transducers Constituted by Carbon Nanotubes Functionalized with DNA Homopolynucle	4/2010
XXXV Congress of Theoretical Chemists of Latin Expression (QUITEL 2009) - San Andrés Islas, Colombia Electronic Response of Chemical Transducers Constituted by Carbon Nanotubes Functionalized with DNA Homopolynucleotides	9/2009
II National Meeting of Theoretical and Computational Chemists (II ENQTC) - Calarcá, Quindio, Colombia Computational Study of the Interactions between Carbon Nanotube/DNA Hybrids and Simple Molecules Relevant in Chemical Sensing	4/2010
Participations	
PSI4 World Wide Developers Conference 2017 (PSI4 WWDC 2017) Blacksburg, Virginia, USA	11/2017
7th Time-Dependent Density-Functional Theory: Prospects and Applications School and Workshop (7th TDDFT) Benasque, Aragón, Spain	9/2016
2015 Computational Chemistry and Materials Science Summer Institute (CCMS 2015) Livermore, CA, USA	6/2015
Sustainable Software Innovation Institute for Computational Chemistry and Materials Modeling ((SICM) ²) Stony Brook, NY, USA	7/2014
II Colombian School on Theory and Computation in Molecular Sciences (II ECTCCM) Guatapé, Antioquia, Colombia	5/2014
246th National Meeting & Exposition of the American Chemical Society (ACS 2013) Indianapolis, IN, USA	8/2013
45th Midwest Theoretical Chemistry Conference (MWTCC 2013) Urbana-Champaign, IL, USA	7/2013
IX International Seminar of Neuroscience Bogotá, D.C., Colombia	6/2012
I Colombian School on Theory and Computation in Molecular Sciences (I ECTCCM) Cali, Valle del Cauca, Colombia	5/2012
Outreach Programs and Initiatives	
Interchange Program 2016 - West Lafayette, IN, USA Theoretical and Computational Chemistry Science Club	7/2016
Clubes de Ciencia - Colombia 2015 - Puerto Triunfo, Antioquia, Colombia Computational Chemistry Science Club Designer and Instructor	10/2015
Interchange Program 2015 - Medellín, Antioquia, Colombia Chemistry and Biology Instructor	9/2015

Professional Affiliations

Member of the American Chemical Society (ACS) 7/2013 - Present

Member of the Colombian Student Association at Purdue University (CSAP) 7/2012 - Present

Member of the Professional Chemists Council of Colombia (CPQ)

2/2010 - Present

Professional Card No.: PQ-3170

Jesuit Alumni Association of Cali, Colombia (A.S.I.A. Santiago de Cali) 5/2004 - Present

Member of the Board (2008-2012)

Languages

Spanish: Native speaker.

English: Fluently spoken and written. Live and work in the United States since 2012.

Portuguese: Intermediate proficiency.

Computer Skills

Chemistry: LibEFP/EFPMD, GROMACS, NAMD, PSI4, Q-Chem, Octopus, GAMESS, HyperChem, Gaussian,

MOPAC, NWChem, VMD, IQmol, GaussView, Gabedit, Avogadro, Molden, ChemBioOffice.

Basic programing: Python, HTML, C/C++.

Others: LibreOffice, Jupyter.

References

Prof. C. David Sherrill | Postdoctoral Adviser 8/2017 - Present

Professor - Department of Chemistry - Georgia Institute of Technology

Email: sherrill@gatech.edu | Office Phone: +1 (404) 894-7452

Prof. Lyudmila V. Slipchenko | Doctoral Adviser 8/2012 - Present

Associate Professor - Department of Chemistry - Purdue University Email: lslipchenko@purdue.edu | Office Phone: +1 (765) 494-5255

Prof. Lynne S. Taylor | Research Collaborator 8/2014 - 5/2018

Retter Professor of Pharmacy - Department of Industrial and Physical Pharmacy - Purdue University

Email: lstaylor@purdue.edu | Office Phone: +1 (765) 496-6614

Prof. Adam Wasserman | Committee Member 8/2012 - 5/2017

Associate Professor - Department of Chemistry - Purdue University Email: awasser@purdue.edu | Office Phone: +1 (765) 494-2348

Interests

• Computer repair and upgrade. • Aircraft modeling and simulation.

LEGO(R).
 Automobile mechanics.