

Biographical Sketch

James Daniel Whitfield

Contact

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Academic Appointments

06/2016 - Present **Assistant Professor of Physics**, Dartmouth College
Department of Physics and Astronomy

Postdoctoral Fellowships

- 2015-2016 **Postdoctoral Fellow**, University of Ghent (Ghent, Belgium)
Advisor: Frank Verstraete
- 2012-2015 **VCQ Postdoctoral Fellow**, Vienna Center for Quantum Science and Technology (Vienna, Austria)
Advisor: Frank Verstraete
- 2011-2012 **Postdoctoral Fellow**, Columbia University (New York, NY)
Advisors: Boris Altshuler
- 2011-2012 **Postdoctoral Fellow**, NEC Laboratories America (Princeton, NJ)
Advisors: Jérémie Roland

Education

- 2011 **Ph.D. Chemical Physics**, Harvard University (Cambridge, MA)
Thesis: *At the Intersection of Quantum Computing and Quantum Chemistry*
Advisor: Alán Aspuru-Guzik
- 2009 **A.M. Chemistry**, Harvard University (Cambridge, MA)
- 2006 **B.S. Mathematics and Chemistry**, *Magna cum laude*, Morehouse College (Atlanta, GA)

Awards

- 2015 Ford Postdoctoral Fellowship
- 2011 Molecular Physics Young Author Prize, *Journal of Molecular Physics*
For "Simulation of electronic structure Hamiltonians using quantum computers"
- 2006-11 Harvard University Graduate Prize Fellowship, Harvard University
- 2016 Walter and Constance Berke Fellowship

Professional Activities

- 2018 Participant, Summit on Advancing American Leadership in Quantum Information Science, White House
- 2018 Long-term visitor at Simons Institute for workshop “Challenges in Quantum Computing” (UC Berkeley)
- 2018 NSF Young Investigator’s Workshop
- 2017 American Association of Physics Teachers New Faculty Workshop participation
- 2017 Programme Committee for conference *Theory of Quantum Computation, Communication, and Cryptography*
- 2017 Ford Foundation Fellowships Review Panel
- 2015 Co-organized conference *Novel Computing Approaches to Quantum Chemistry*, Telluride, CO
- 2012 Visiting Scholar, Max Planck Institute for the Physics of Complex Systems, Condensed Matter Group
- 2012 Visiting Scientist, Institute for Scientific Interchange, Quantum Science Laboratory

Reviewer for various journals (Quantum Information and Computation, New Journal of Physics, Nature, Nature Communications, Physical Review A, Physical Review Letters, Quantum Science and Technology, Journal of Physics A: Mathematical and Theoretical, International Journal of Quantum Chemistry, NSF Ad-Hoc panels)

Professional memberships

- American Physical Society
- American Chemical Society

College Committees

- Colloquium Committee member, Fall 2018 - Fall 2019
- Digital Outreach Committee chair, since Fall 2016
- Graduate Admissions and Policy committee member, since Fall 2016

Teaching

- Fall 2016 Introduction to Condensed Matter, Phys 73
- Spring 2017 Quantum Physics of Matter, Phys 40
- Fall 2017 Introduction to Condensed Matter, Phys 73
- Winter 2018 Introductory Physics I, Phys 13
- Spring 2018 Quantum Information, Phys 116
- Fall 2018 Introduction to Condensed Matter, Phys 73
- Winter 2019 Introductory Physics I, Phys 13
- Spring 2019 Quantum Physics of Matter, Phys 40

Undergraduate theses supervised

- Samuel Greydanus, Undergraduate thesis 2017
“Approximating Matrix Product States with Machine Learning”
- Erik Weis, Undergraduate thesis 2018
“Benchmarking Quantum Computers Using Electronic Structure Algorithms”
- Shaket Chaudhary, Undergraduate thesis 2019
”Constructing Random Ensembles of Fermionic Systems”

Talks since 2016

Conference Presentations

- Quantum Indeterminacy Workshop (invited talk) — Hanover, NH — July 2019
 - Computation versus Experiment: Quantum technology meets quantum computing. July 13, 2019
- American Mathematics Society Joint Mathematics Meeting (contributed talk) — Baltimore, MD — Jan 2019
 - Quantum Measurement Problem. Session: AMS Special Session on 25 years of Conferences for African-American Researchers in the Mathematical Sciences (CAARMS times 25), II. Jan 16, 2019
- American Physical Society Meeting (contributed talk) — Boston, MA — Mar 2019
 - Quantum simulation and Time-Dependent Density Functional Theory. Session: Applications of Noisy Intermediate Scale Quantum Computers III. Jan 5, 2019
- International Workshop on Quantum Chemical Calculations on Quantum Computers (invited talk) — Osaka City University (OCU), Osaka, Japan — Mar 2018
 - Fermionic Algebras for Quantum Computing. Mar 29, 2018

Seminars/Colloquia

- Stony Brook University — Stony Brook, NY Quantum Simulation of Fermions: Fermion-to-spin mappings and TDDFT. May 16, 2019
- Rigetti Quantum Computing — Berkeley, CA Hydrogen Quantum Simulation. July 18, 2018
- Simons Institute — Berkeley, CA Formalizing Electronic Structure Problems. July 12, 2018
- Sandia National Laboratories — Livermore, California Quantum Simulation of Fermions. July 9, 2018
- Pacific Northwest Laboratories — Richland, Washington. Quantum Simulation of Fermions. August 7, 2018
- Microsoft — Redmond, Washington Quantum Simulation and Applications. August 9, 2018
- Virginia Institute of Technology — Blacksburg, VA Quantum Simulation of Fermions. Sept 10, 2018
- Bates College — Lewiston, Maine The Promise of Quantum Technology. November 15, 2018
- IBM T. J. Watson — Yorktown Heights, NY. Fermionic simulation on quantum computers, June 9, 2018
- Quantum Lunch Seminar at Los Alamos National Laboratory — Los Alamos, NM. Fermionic algebras for qubits, January 26, 2017
- Georgia Tech Quantum Information Seminar at Georgia Institute of Technology — Atlanta, GA Electronic structure on quantum computers with ultra-local qubit operators. January 31, 2017

Funding since 2016

1. National Science Foundation Grant PHYS-1820747, 2018-2021
“Topological Fermionic Quantum Simulation,” PI, \$406,713/3 years.
2. Department of Energy Grant DE-SC0019374, 2018-2021
“Quantum Chemistry for Quantum Computers,” Co-PI, \$536,000/3 years.
3. Department of Energy Grant DE-SC0019374, 2018-2022
Optimization, Verification, and Engineered Reliability of Quantum Computers co-PI, \$727,244/4 years.

Publications since 2016

1. S Gulania, James Daniel Whitfield. Young frames for quantum chemistry. (Preprint) arXiv:1904.10469.
2. J Brown, J Yang, James Daniel Whitfield. Solver for the electronic V-representation problem of time-dependent density functional theory. (Preprint) arXiv:1904.10958.
3. R W Chien, S Xue, T S Hardikar, K Setia, James Daniel Whitfield. Analysis of Superfast Encoding Performance for Electronic Structure Simulations. (Preprint) arXiv:1907.02976. Accepted for publication at Physical Review A.
4. K Setia, S Bravyi, A Mezzacapo, James Daniel Whitfield. Superfast encodings for fermionic quantum simulation. (Preprint) arXiv:1810.05274. Accepted for publication at Physical Review Research.
5. J Brown, James Daniel Whitfield. Basis set convergence of Wilson basis functions for electronic structure. *Journal of Chemical Physics*. 151, 064118, 2019.
6. K Setia, James Daniel Whitfield. Bravyi-Kitaev Superfast simulation of fermions on a quantum computer. *The Journal of Chemical Physics*, 148, 164104, 2018.
7. C Schilling, M Altunbulak, S Knecht, A Lopes, James Daniel Whitfield, M Christandl, D Gross, M Reiher. Generalized Pauli constraints in small atoms. *Physical Review A*, 97, 052503, 2018.
8. G Zhu, Y Subasi, James Daniel Whitfield, M Hafezi. Hardware-efficient fermionic simulation with a cavity-QED system. *New Physics Journal Quantum Information*, 4, 16, 2018.
9. V Havlíček, M Troyer, James Daniel Whitfield. Operator Locality in Quantum Simulation of Fermionic Models. *Physical Review A*, 95, 032332, 2017.
10. James Daniel Whitfield, V Havlíček, M Troyer. Local spin operators for fermion simulations. *Physical Review A*, 94, 030301, 2016.

Previous Publications

11. S Barz, B Dakic, Y O Lipp, F Verstraete, James Daniel Whitfield, P Walther. Linear-optical generation of eigenstates of the two-site XY model. *Physical Review X*, 5(2):021010, 2015.
12. Y Wang, F Dolde, J Biamonte, R Babbush, V Bergholm, S Yang, I Jakobi, P Neumann, A Aspuru-Guzik, James Daniel Whitfield, and J Wrachtrup. Quantum Simulation of Helium Hydride Cation in a Solid-State Spin Register. *ACS Nano*, 9(8):7769-7774, 2015.
13. James Daniel Whitfield, M-H Yung, D G Tempel, S Boixo, A Aspuru-Guzik. Computational complexity of time-dependent density functional theory. *New Journal of Physics*, 16(8):083035, 2014.
14. James Daniel Whitfield, Z Zimborás. On the NP-completeness of the Hartree-Fock method for translationally invariant systems. *The Journal of Chemical Physics*, 141(23):234103, 2014.
15. James Daniel Whitfield. Communication: Spin-free quantum computational simulations and symmetry adapted states. *The Journal of Chemical Physics*, 139:021105, 2013.
16. James Daniel Whitfield, P J Love, and A Aspuru-Guzik. Computational complexity in electronic structure. *Physical Chemistry Chemical Physics*, 15(2):397-411, 2013.
17. Z Zimboras, M Faccin, Z Kadar, James Daniel Whitfield, B Lanyon, and J Biamonte. Quantum transport enhancement by time-reversal symmetry breaking. *Scientific Reports*, 3:2361, 2013.

18. N C Jones, James Daniel Whitfield, P L McMahon, M-H Yung, R Van Meter, A Aspuru-Guzik, and Y Yamamoto. Faster quantum chemistry simulation on fault-tolerant quantum computers. *New Journal of Physics*, 14(11):115023, 2012.
19. James Daniel Whitfield, M Faccin, and J D Biamonte. Ground-state spin logic. *EPL (Europhysics Letters)*, 99(5):57004, 2012.
20. James Daniel Whitfield. Chapter 7 - Electronic Structure in “Mathematical Modeling II: Quantum Mechanics and Spectroscopy” by Troy L. Story. Zip Publishing, 2012.
21. J D Biamonte, V Bergholm, James Daniel Whitfield, J Fitzsimons, and A Aspuru-Guzik. Adiabatic quantum simulators. *AIP Advances*, 1(2):022126-022126, 2011.
22. I Kassal, James Daniel Whitfield, A Perdomo-Ortiz, M-H Yung, and A Aspuru-Guzik. Simulating chemistry using quantum computers. *Annual Review of Physical Chemistry*, 62:185-207, 2011.
23. Z Li, M-H Yung, H Chen, D Lu, James Daniel Whitfield, X Peng, A Aspuru-Guzik, and J Du. Solving quantum ground-state problems with nuclear magnetic resonance. *Scientific Reports*, 1, 2011.
24. James Daniel Whitfield, J. Biamonte, and A. Aspuru-Guzik. Simulation of electronic structure Hamiltonians using quantum computers. *Molecular Physics*, 109(5):735-750, 2011.
25. B P Lanyon, James Daniel Whitfield, G G Gillett, M E Goggin, M P Almeida, I Kassal, J D Biamonte, M Mohseni, B J Powell, M Barbieri, et al. Towards quantum chemistry on a quantum computer. *Nature Chemistry*, 2(2):106-111, 2010.
26. James Daniel Whitfield, C A Rodriguez-Rosario, and A Aspuru-Guzik. Quantum stochastic walks: A generalization of classical random walks and quantum walks. *Physical Review A*, 81(2):022323, 2010.
27. M-H Yung, D Nagaj, James Daniel Whitfield, and A Aspuru-Guzik. Simulation of classical thermal states on a quantum computer: A transfer-matrix approach. *Physical Review A*, 82(6):060302, 2010.