HUNG Q. PHAM

Computational Chemist

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EXPERIENCE

Research Assistant

• General Chemistry Laboratory

VNUHCM-US

→ Ho Chi Minh, Vietnam (Oct 2011 – Jul 2015)

• General Chemistry Laboratory, Physical Chemistry Laboratory

Teaching Assistant

University of Minnesota ♀ Minneapolis, USA (Sep 2015 – Dec 2016)

• Developing electronic structure theories for periodic systems.

VNUHCM-US ♥ Ho Chi Minh, Vietnam (Oct 2011 – Jul 2015)

• First-principle calculations on framework materials (MOFs, COFs).

PUBLICATIONS

- 1. **H. Q. Pham**, M. R. Hermes, L. Gagliardi; *Periodic Density Matrix Embedding Theory: A Wave Function Approach for Strongly Correlated Materials*,, **2019**, Manuscript under preparation. §
- H. Q. Pham*, D. Q. Le, N.-N. Pham-Tran, Y. Kawazoe, D. Nguyen-Manh; Electron Delocalization in Single-Layer Phthalocyanine-Based Covalent Organic Frameworks: A First Principle Study, RSC Advances, 2019, Manuscript under revision.
- 3. R. Sanner, N. Cherepy, H P. Martinez, H. Q. Pham, V. Young; Highly Efficient Phosphorescence From Cyclometallated Iridium(III) Compounds: Improved Syntheses of Picolinate Complexes and Quantum Chemical Studies of Their Electronic Structures, Inorganica Chim. Acta, 2019, 496, 119040.
- 4. **H. Q. Pham**, R. J. Holmes, E. S. Aydil, L. Gagliardi; Lead-Free Double Perovskites $Cs_2 lnCuCl_6$ and $(CH_3NH_3)_2 lnCuCl_6$: Electronic, Optical, and Electrical Properties, Nanoscale, 2019, 11, 11173. **%**
- H. Q. Pham, V. Bernales, L. Gagliardi; Can Density Matrix Embedding Theory with the Complete Active Space Self-Consistent Field Solver Describe Single and Double Bond Breaking in Molecular Systems?, J. Chem. Theory Comput., 2018, 14, 1960.
- 6. D. Ray, C. Clarke, **H. Q. Pham**, J. Borycz; T. Zhang, E. S. Aydil, R. J. Holmes, L. Gagliardi; A Computational Study of Structural and Electronic Properties of Lead-Free CsMl₃ Perovskites (M = Ge, Sn, Pb, Mg, Ca, Sr, Ba), **J. Phys. Chem. C**, 2018, 122, 7838.
- P.T. K. Nguyen, H. T. D. Nguyen, H. Q. Pham, J. Kim, K. E. Cordova, H. Furukawa; Synthesis and Selective CO₂ Capture Properties of a Series of Hexatopic Linker-Based Metal-Organic Frameworks, Inorg. Chem., 2015, 54 (20), 10065.
- 8. T. L. H. Doan, H. L. Nguyen, **H. Q. Pham**, N.-N. Pham-Tran, T. N. Le, K. E. Cordova; *Tailoring the Optical Absorption of Water-Stable Zr^{IV-} and Hf^{IV-}-Based Metal-Organic Framework Photocatalysts*, **Chem. Asian J.**, 2015, 10, 2660. %
- H. Q. Pham, T. Mai, N.-N. Pham-Tran, Y. Kawazoe, H. Mizuseki, D. Nguyen-Manh; Engineering of Band Gap in Metal-Organic Frameworks by Functionalizing Organic Linker: A Systematic Density Functional Theory Investigation, J. Phys. Chem. C, 2014, 118 (9), 4567.

EDUCATION

Ph.D. in Chemistry
University of Minnesota, USA

B.S. in Chemistry VNUHCM-US, Vietnam

🛗 Sep. 2007 - Oct. 2011

AREA OF INTEREST

- Wave function theories for condensed phases
- Quantum embedding theory
- Density functional theory
- Semiconducting, framework materials

HONORS & AWARDS

Lawrence S. Ting Scholarship
Lawrence Ting Memorial Fund, Vietnam (2008)

Academic Excellent Scholarship VNUHCM-US, Vietnam (2008,2010,2011)

SKILLS

PySCF VASP CRYSTAL Gaussian

Materials Studio Crystal Maker

OpenMP Git Scikit-learn Scipy

Numpy Matplotlib

LANGUAGES

Python C/C++ Fortran Shell script



REVIEWING ACTIVITY

- Journal of Chemical Theory and Computation, ACS %
- Solid State Communications, Elsevier %