Jun Huo

Master

School of Petroleum and Chemical Engineering, Dalian University of Technology No.2 Linggong Road, Ganjingzi District, Dalian City, Liaoning Province, P.R.China

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Education

Sep. 2016 - Present M.S. in Chemical Engineering

School of Petroleum and Chemical Engineering,

Dalian University of Technology

Sep. 2012 - Jul. 2016 B.Eng. in Chemical Engineering and Technology

College of Chemistry, Chemical Engineering and Environmental

Engineering, Liaoning Shihua University

Research Interests

Combining Machine Learning and Molecular Simulation to study polymer structure

Publications

- * indicates co-first author.
 - Jun Huo, Wenxu Qi, Hongda Zhu, et al., Molecular Dynamics Simulation on the Effect of Water Uptake on Hydrogen Bond Network for OH⁻ Conduction in Imidazolium-g-PPO Membrane, *International Journal of Hydrogen Energy*, 2019, 44: 3760–3770. doi:10.1016/j.ijhydene.2018.12.090
 - Ning Zhang, Boyun Yang, Jun Huo, et al., Hydration Structures of Vanadium/Oxovanadium Cations in the Presence of Sulfuric Acid: A Molecular Dynamics Simulation Study, Chemical Engineering Science, 2019, 195: 683-692. doi:10.1016/j.ces.2018.10.014
 - 4. Ning Zhang*, **Jun Huo***, Boyun Yang, et al., Understanding of Imidazolium Group Hydration and Polymer Structure for Hydroxide Anion Conduction in Hydrated Imidazolium-g-PPO Membrane by Molecular Dynamics Simulations, *Chemical Engineering Science*, **2018**, 192: 1167–1176. doi:10.1016/j.ces.2018.08.051
 - 3. Yuechun Song, **Jun Huo**, Ning Zhang, et al., Structural Characteristics of Hydrated Protons in Ion Conductive Channels: Synergistic Effect of the Sulfonate Group and Fluorine Studied by Molecular Dynamics Simulation, *The Journal of Physical Chemistry C*, **2018**, 122(4): 1982-1989. doi:10.1021/acs.jpcc.7b11020
 - Ning Zhang, Shaomin Chen, Boyun Yang, Jun Huo, et al., Effect of Hydrogen-Bonding Interaction on the Arrangement and Dynamics of Water Confined in a Polyamide Membrane: A Molecular Dynamics Simulation, *The Journal of Physical Chemistry B*, 2018, 122(17): 4719-4728. doi:10.1021/acs.jpcb.7b12790

 Ning Zhang, Yuechun Song, Jun Huo, et al., Formation Mechanism of the Spiral-Like Structure of a Hydrogen Bond Network Confined in a Fluorinated Nanochannel: A Molecular Dynamics Simulation, *The Journal of Physical Chemistry C*, 2017, 121(25): 13840-13847. doi:10.1021/acs.jpcc.7b01074

Papers submitted / under review

1. **Jun Huo**, Wenxu Qi, Gaohong He, et al., Structural Characteristics of Hydrated Protons in the Conductive Channels: Effect of Electric Field Studied by Molecular Dynamics Simulation, *Nanoscale*. (Under revison)

Skills

Languages Mandarin Chinese, English.

Programming Python, C/C++, Tcl/Tk, Fortran, LATEX, etc.

Software NAMD, Lammps, VMD, Materials Studio, Gaussian, Office, Photoshop, etc.

Awards

2018 - 2019	The First Prize of Excellent Master Degree Scholarship
2017 - 2018	The First Prize of Excellent Master Degree Scholarship
2016 - 2017	Outstanding Graduate Student Leader
2014 - 2015	Liaoning Provincial Government Scholarship
2013 - 2014	Outstanding Student Leader

Experience

Sep. 2016 - Present	Natural Science Foundation of China Research on the structure and formation mechanism of hydrated proton in the ion transport channel of proton exchange membrane.
Sep. 2016 - July. 2017	Director of Publicity Department, the Graduate Student Union
Sep. 2016 - July. 2017	College Office Assistant
Sep. 2014 - July. 2015	Director of Publicity Department, the Student Union