

# Rasha Atwi

PHD CANDIDATE · STONY BROOK UNIVERSITY

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## Education

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### Stony Brook University / Tufts University

DOCTOR OF PHILOSOPHY, CHEMICAL ENGINEERING

Advisor: Prof. Nav Nidhi Rajput

Stony Brook, NY / Medford, MA

Sep 2018 - May 2024

### Northeastern University

MASTER OF SCIENCE, CHEMICAL ENGINEERING

Advisor: Prof. Richard West

Boston, MA

Sep 2016 - Aug 2018

### American University of Beirut

BACHELOR OF ENGINEERING, CHEMICAL ENGINEERING

With distinction

Beirut, Lebanon

Sep 2012 - May 2016

## Research

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### Stony Brook University / Tufts University

GRADUATE STUDENT RESEARCHER

- Explore structural and dynamical properties of electrolytes and electrode-electrolyte interfaces in batteries using DFT and MD simulations.
- Build computational databases for high-throughput screening of electrolyte materials for Li-S batteries.
- Develop new computational methods for predicting spectroscopic signatures in liquid solutions. The methods address some of the most challenging aspects of computational spectroscopy - equilibrium conformers and solvation effects.
- Write open-source software packages for automating high-throughput multi-scale simulations on supercomputers.

Stony Brook, NY / Medford, MA

2018 - Present

### Northeastern University

GRADUATE STUDENT RESEARCHER

Developed a kinetics model using *ab-initio* thermodynamic simulations for the formation of nitrogen heterocycles during hydrothermal liquefaction of micro-algae.

Boston, MA

2016 - 2018

### Centre for the Theory and Application of Catalysis, Queen's University Belfast

VISITING RESEARCH ASSOCIATE

Synthesized, characterized, and tested heterogeneous catalysts for Fischer Tropsch reaction and operated and controlled chemical rigs.

Belfast, UK

2015

### American University of Beirut

UNDERGRADUATE STUDENT RESEARCHER

Identified pharmaceutical compounds present in the Lebanese water bodies based on physico-chemical properties and prescription data.

Beirut Lebanon

2013 - 2014

## Work Experience

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### Merck & Co.

COMPUTATIONAL & STRUCTURAL CHEMISTRY INTERN

- Investigated the reaction mechanism behind nitrosamine formation in drug products—a carcinogenic and genotoxic compound causing FDA recalls.
- Built and validated a model utilizing physics-based and machine learning approaches to predict nitrosamine yield. The model streamlines risk assessments by identifying secondary amines with high risk of forming nitrosamines.

Rahway, NJ

Jun 2023 - Aug 2023

### Biogen

COMPUTATIONAL CHEMISTRY CO-OP

- Developed a user-friendly 3D molecular shape comparison algorithm for virtual high-throughput screening applications.
- Utilized GPU acceleration for optimizing the performance of the toolkit.
- Implemented the user-interface of the toolkit on Biogen Molecular Informatics Platform.

Cambridge, MA

Jan 2023 - May 2023

### CF Technologies Inc.

OIL EXTRACTION HAZOP TEAM MEMBER

- Reviewed the hazard and operability considerations of a pilot unit that extracts oil from brown grease using supercritical fluids.
- Developed a process flow diagram (PFD) and a piping and instrumentation diagram (P&ID) for the pilot unit.

Hyde Park, MA

Jan 2018 - May 2018

### Power Advocate (Part of Wood Mackenzie)

ENERGY BUSINESS ANALYST CO-OP

- Created cost models for specialty chemicals, equipment, and services used by the energy sector.
- Developed commodity forecasts, performed market analysis, and conducted research for publishing metals and chemicals foresights.

Boston, MA

Jul 2017 - Dec 2017

# Software Development

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## ROSHAMBO

MAIN DEVELOPER & MAINTAINER

Jan 2023 - Present

A Python package for robust Gaussian molecular shape comparison.

GitHub repository: <https://github.com/rashatwi/roshambo>

## MISPR: Materials Informatics for Structure Property Relationships

MAIN DEVELOPER & MAINTAINER

Sep 2018 - Present

An open-source Python package for automating materials science computations (DFT and MD simulations).

GitHub repository: <https://github.com/molmd/mispr>

## MDPropTools

MAIN DEVELOPER & MAINTAINER

Sep 2018 - Present

An open-source Python package for statistical analysis of MD trajectories and output files to compute properties of liquid solutions.

GitHub repository: <https://github.com/molmd/mdproptools>

# Skills

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<b>Programming</b>	Python, Cython, MATLAB, NoSQL
<b>Software</b>	Gaussian, LAMMPS, Schrödinger Suites, OpenEye Toolkits, AmberTools, TensorFlow, Keras, Aspen HYSYS, LABVIEW
<b>Other</b>	Git, Unix systems, Shell, Jupyter, LATEX

# Honors and Awards

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<b>Early Career Group Best Poster Award</b> , Battery+Energy Storage Conference, Argonne National Lab	2023
<b>IACS Junior Researcher Award</b> , Stony Brook University	2021 - Present
"Given to continuing PhD graduate students who are recognized as outstanding junior researchers by institute faculty or affiliates."	
<b>Dean's Fellowship</b> , Tufts University	2018 - 2020
"To recognize and recruit exceptional incoming doctoral students."	
<b>College of Engineering Top GPA Award</b> , Northeastern University	2019
<b>Outstanding Seminar Award</b> , Northeastern University	2018
<b>Best Student Poster Award</b> , American University of Beirut	2016
<b>Dean's Honor List</b> , American University of Beirut	2016
"Ranked in the top 10% of the class."	

# Teaching and Research Mentoring

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<b>Workshop Helper</b> , Software Carpentry	2023
Assisted learners one-on-one with software installation, understanding specific lines of code, and other aspects of the learning process related to bash, Git version control, and programming in Python. <a href="#">Link</a> .	
<b>MISPR Workshop Instructor</b> , Stony Brook University	Jul 2022
Organized and ran MISPR hands-on workshop at iACS to introduce high-school students to materials informatics techniques. <a href="#">Link</a> .	
<b>Research Mentor</b> , Stony Brook University / Tufts University	2019 - 2022
<ul style="list-style-type: none"><li>Mentored high-school and undergraduate students on projects related to force field database development and electrolyte property predictions.</li><li>Introduced students to computational chemistry techniques, including DFT, and provided hands-on experience in programming to automate tasks and analyze data effectively.</li></ul>	
<b>Teaching Assistant</b> , Tufts University	2018 - 2020
<ul style="list-style-type: none"><li>Courses: Transport phenomena (CHBE-0022), Advanced thermodynamics (CHBE-0203).</li><li>Held office hours and reviewed and graded homework problems, reports, and exams.</li><li>Led recitation and tutorial sessions to review concepts and work through practice problems.</li></ul>	

# Professional and Community Activities

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<b>Academic Peer Reviewer</b> , WoS ResearcherID: <a href="#">IXN-5252-2023</a> .	2023 - Present
Conferences: ICLR-ml4materials (2), NeurIPS-AI4Mat (3). Journals: J. Open Source Softw. (1)	

Develop a plan to increase membership of students, industry, and international members in the division.

Review travel fellowship applications for the supercomputing conference (SC) held each year.

Helped international graduate students adjust to life in the U.S. and build confidence in their conversational English through weekly one-on-one meetings.

## Publications

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7. **Atwi, R.** & Rajput, N. N. (2023). "Guiding maps of solvent classes for lithium-sulfur batteries via a computational data-driven approach". *Patterns*, 4(9). [Link](#).
6. Li, Z., Rao, H., **Atwi, R.**, Sivakumar, B.M., Gwalani, B., Gray, S., Han, K.S., Everett, T.A., Ajantiwalay, T.A., Murugesan, V., Rajput, N.N., & Pol, V. G. (2023). "Non-polar ether-based electrolyte solutions for stable high-voltage non-aqueous lithium metal batteries". *Nature Communications*, 14, 868. [Link](#).
5. **Atwi, R.**, Bliss, M., Makeev, M., & Rajput, N. N. (2022). "MISPR: An open-source package for high-throughput multiscale molecular simulations". *Scientific Reports*, 12, 15760. [Link](#).
4. **Atwi, R.**, Chen, Y., Han, K. S., Mueller, K. T., Murugesan, V., & Rajput, N. N. (2022). "An automated framework for high-throughput predictions of NMR chemical shifts within liquid solutions". *Nature Computational Science*, 2(2), 112-122. [Link](#).
3. LeClerc, H. O., **Atwi, R.**, Niles, S. F., McKenna, A. M., Timko, M. T., West, R. H., & Teixeira, A. R. (2022). "Elucidating the role of reactive nitrogen intermediates in hetero-cyclization during hydrothermal liquefaction of food waste". *Green Chemistry*, 24(13), 5125-5141. [Link](#).
2. Chen, Y., **Atwi, R.**, Han, K. S., Ryu, J., Washton, N. M., Hu, J. Z., Rajput, N. N., Mueller, K. T., & Murugesan, V. (2021). "Role of a multi-valent ion-solvent interaction on restricted  $Mg^{2+}$  diffusion in dimethoxyethane electrolytes". *The Journal of Physical Chemistry B*, 125(45), 12574-12583. [Link](#).
1. Blanco, D. E., **Atwi, R.**, Sethuraman, S., Lasri, A., Morales, J., Rajput, N. N., & Modestino, M. A. (2020). "Effect of electrolyte cations on organic electrosynthesis: the case of adiponitrile electrochemical production". *Journal of The Electrochemical Society*, 167(15), 155526. [Link](#).

## Posters and Presentations

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17. **Atwi, R.** & Rajput, N. N. (Fall 2023). "An automated multi-scale NMR framework helps understand solvent exchange dynamics in multivalent batteries". MRS Meeting. Oral Presentation.
16. **Atwi, R.** & Rajput, N. N. (Fall 2023). "Guiding maps of solvents for lithium-sulfur batteries via a computational data-driven approach". 5th Battery and Energy Storage Conference (Argonne National Laboratory). Poster Presentation.
15. **Atwi, R.** & Rajput, N. N. (Fall 2022). "Data-driven prediction of structure and dynamic features of Li-S electrolytes with an automated multi-scale computational infrastructure". MRS Meeting. Oral Presentation.
14. **Atwi, R.** & Rajput, N. N. (Fall 2022). "Data-driven investigation of solvents for lithium-sulfur battery liquid electrolytes". ACS Meeting. Oral Presentation. [Link](#).
13. **Atwi, R.** & Rajput, N. N. (Fall 2022). "Towards accurate predictions of NMR chemical shifts in liquid solutions". ACS Meeting. Oral Presentation. [Link](#).
12. **Atwi, R.**, Bliss, M. M., & Rajput N. N. (Summer 2022). "MISPR: A novel framework for high-throughput multi-scale modelling of complex liquid solutions". FOMMS: Molecular Modeling and the Data Revolution Conference. Poster Presentation.
11. **Atwi, R.** & Rajput, N. N. (Spring 2022). "Tailoring atomistic interactions in Li-S battery via a computational multi-scale data-driven approach". 241st ECS Meeting. Oral Presentation. [Link](#).
10. **Atwi, R.** (Spring 2022). "Towards next generation batteries via high-throughput modelling of electrolytes". IACS Brown Bag Lunch. Oral Presentation. [Link](#).
9. **Atwi, R.** (Fall 2022). "Tailoring atomistic interactions in Li-S battery via a computational multi-scale data-driven approach". IACS Student Seminar. Oral Presentation. [Link](#).
8. Rajput, N. N. & **Atwi, R.** (Spring 2021). "Designing optimal electrolytes and interfaces in Li-S batteries". 239th ECS Meeting. Invited Oral Presentation. [Link](#).
7. Rajput, N. N. **Atwi, R.**, & Bliss, M. (Spring 2021). "A multi-scale infrastructure for automating materials science computations".

239th ECS Meeting. Invited Oral Presentation. [Link](#).

6. **Atwi, R.** & Rajput, N. N. (Spring 2021). “Effects of functionalized cathode and electrolyte composition on structure and dissolution of polysulfide species in Li-S batteries”. [MRS Meeting](#). Oral Presentation.
5. **Atwi, R.** & Rajput, N. N. (Fall 2020). “Tailoring atomistic interactions in Li-S battery via a computational multi-scale-data-driven approach”. [MRS Meeting](#). Oral Presentation.
4. **Atwi, R.**, Bliss, M. M., & Rajput N. N. (Fall 2020). “A high-throughput multi-scale infrastructure for automating materials science computations”. [MRS Meeting](#). Oral Presentation.
3. Timko, M. T., West, R. H., & **Atwi, R.** (2019). “Understanding formation of nitrogen heterocycles during catalytic hydrothermal liquefaction”. [AIChE Annual Meeting](#). Oral Presentation. [Link](#).
2. **Atwi, R.**, Timko, M. T., & West, R. H. (Fall 2018). “Investigating reaction pathways for the formation of nitrogen heterocycles during hydrothermal liquefaction of microalgae”. [ACS Meeting](#).
1. **Atwi, R.** & West, R. H. (Summer 2017). “Understanding the source of undesired nitrogen heterocycles in biofuel made by hydrothermal liquefaction of micro-algae”. [New England Energy Research Forum](#). Poster Presentation.