





## SUMAN SAMANTRAY

IBI-7: Structural Biochemistry,  
Forschungszentrum Jülich GmbH,  
Wilhelm-Johnen-Straße, 52425 Jülich, Germany

+49(0)176 – 20986662   [s.samantray@fz-juelich.de](mailto:s.samantray@fz-juelich.de)   [suman.rishis@gmail.com](mailto:suman.rishis@gmail.com)     
[@samantray\\_1990](https://twitter.com/samantray_1990)   [suman – samantray](https://github.com/suman-samantray)   [suman – samantray.github.io](https://suman-samantray.github.io)   

---

### RESEARCH INTERESTS

Amyloid aggregation, Interfacial phenomena, Molecular simulation, Computational Chemistry, Machine Learning, Drug discovery, Graph convolutional neural networks

### EDUCATION

- **IBI-7: Structural Biochemistry, FZ Jülich GmbH, Jülich, Germany / AICES fellow, RWTH Aachen University, Aachen, Germany** *Aug 2018 – present*  
*Doctor of Philosophy (Dr. rer. nat.) in Computational Biochemistry*

- Dissertation : Simulation of Amyloid Aggregation under *in vivo* Conditions  
Supervisor : Prof. (Dr.) Birgit Strodel

- **State University of New York, Buffalo, USA** *Aug 2013 – Aug 2015*  
*Master of Engineering (M.Eng) in Chemical Engineering*

- Dissertation : Calculation of Saturation and Interfacial properties of model Carbon Dioxide-Water system using Monte Carlo Simulation.  
Supervisor : Prof. (Dr.) Jeffrey R. Errington

- **Indira Gandhi Institute of Technology, Odisha, India** *Aug 2009 – Jun 2013*  
*Bachelor of Technology (B.Tech) in Chemical Engineering*

- Dissertation : Synthesis of activated carbon from Jackfruit peel waste and Coconut husk for Purification of water.  
Supervisor : Prof. (Dr.) Satyabrata Mohanta

### RESEARCH EXPERIENCE

- **Ph.D. Researcher** *Aug 2018–Present*  
*IBI-7: Structural Biochemistry, FZ Jülich GmbH, Germany* Prof. (Dr.) Birgit Strodel

- Determination of molecular mechanics parameters and building kinetic transition models to elucidate the amyloid- $\beta$  aggregation pathways.
- Identification of bio-mimetic molecules inhibiting amyloid- $\beta$  aggregation.
- Development of simulation methods for studying amyloid aggregation under the influence of glycosaminoglycans.
- Co-organiser of hands-on workshop on **Molecular Dynamics Simulations of Proteins** at [IHRS BioSoft](#).
- Co-supervision of Strodel group online code databases on [GitHub](#).
- Maintenance of Strodel group computing clusters and cloud storage services.
- Co-mentoring of HiWi and M.Sc. students in the Strodel group.

■ **College of Science postgraduate fellow**

*School of Chemistry, NUI Galway, Ireland*

*Jan 2017–Jul 2018*  
Prof. (Dr.) David L. Cheung

- Using molecular simulation to understand the behaviour of Intrinsically disordered/ Amyloidogenic proteins at air-water interface (AWI).
- Using the replica exchange and metadynamics simulations to investigate protein structures at liquid interfaces.

■ **Research Associate**

*Dept. of Industrial Design, NIT Rourkela, India*

*Oct 2016–Dec 2016*  
Prof. (Dr.) Dibya Prakash Jena

- Worked in the Industrial Acoustics lab to identify a benchmark acoustic cloaking device.
- Built an impedance tube with an attached cylindrical Helmholtz resonator to evaluate net acoustic transmission loss using transfer matrix method.

■ **Graduate Research Student**

*SUNY Buffalo, NY, USA*

*Sep 2013–Sept 2015*  
Prof. (Dr.) Jeffrey R. Errington

- Applied Grand Canonical Monte Carlo simulation method to compute vapor-liquid coexistence properties of carbon dioxide and water fluid mixture.
- Used free energy-based approach to determine interfacial properties of the binary fluid mixture including activity fraction expanded ensemble technique on atomistic silica-like surface.
- Developed algorithms in python to analyze and interpret data from GCMC simulation.

## PROFESSIONAL EXPERIENCE

■ **Reviewer**

*Jan 2021–present*

- *Molecular Pharmaceutics, International Journal of Molecular Science*

■ **Computing Assistant**

*Information Solutions and Services, NUI Galway, Ireland*

*Aug 2017–Jul 2018*  
Mr. Peter Crampton

- Responsible for the management, development, physical upkeep and maintenance of the ISS and departmental PC suites across campus.
- Assist the Desktop Services, Provisioning and Support Manager, ensuring efficient operationally of all PC suites.

■ **Senior Application Developer**

*Digital products and interactive media (DPIM) III,  
NBC Universal, NY, USA*

*Oct 2015–Sept 2016*  
Mr. Wen Qu, Mrs. Dana Fleur

- Lead a team of 3 Dev's and 2 QA's to develop MPS mobile SDK and built a Test App to displays ads fetch SDK users (NBC native apps) and configure it for vendors purposes during **Rio Olympics 2016**.
- Developed the [NBCUView](#) and recently implemented Apple Push Notification Service. Documented the app architecture including identification of the service end points.

## TEACHING EXPERIENCE

■ **Teaching Instructor**


*School of Chemistry, NUI Galway, Ireland*

*Oct 2017–Apr 2018*  
Prof. (Dr.) David L. Cheung

- Teaching Assistant for Computational Drug Design and Drug Discovery laboratory, Spring 2018
- Teaching Assistant for Physical Chemistry laboratory, Fall 2017

- Demonstration and lecturing on chemistry experiments to primary and secondary school students.

## PUBLICATIONS

Peer Reviewed Publications (Total Publications: 6, Total Citations: 22, h-index: 3) 

\* - equal authorship

- [8] Elucidating the structure of glycosaminoglycans under different physiological conditions, S. Samantray, O.O. Olubiyi, B. Strodel, (*In preparation*), *Biomolecules*
- [7] The Effect of Different Glycosaminoglycans on the Amyloid Aggregation Pathways in Molecular Dynamics Simulations, S. Samantray, B. Strodel, (*Submitted*), *Journal of Physical Chemistry B*
- [6] Thermodynamics and kinetics of the amyloid- $\beta$  peptide revealed by Markov state models based on MD data in agreement with experiment, A. Paul\*, S. Samantray\*, M. Anteghini, B. Strodel, (*In Review*), *Chemical Science*
- [5] Molecular dynamics simulations of protein aggregation: protocols for simulation setup and analysis with Markov state models and transition networks, S. Samantray, W. Schumann, A.-M. Illig, M.-C. Pacheco, A. Paul, B. Barz, B. Strodel, (*In Press*), *Methods in Molecular Biology (Springer)*
- [4] Effect of the Air-Water Interface on the Conformation of Amyloid Beta, S. Samantray, D.L. Cheung, *Biointerphases*, 15(6), 061011 (2020) (*Selected as Featured Article and mentioned in AIP Scilight.*)
- [3] Different force fields give rise to different amyloid aggregation pathways in molecular dynamics simulations, S. Samantray, F. Yin, B. Kav, B. Strodel, *Journal of Chemical Information and modelling*, 60(12), 6462–6475 (2020)
- [2]  $\beta$ -Turn mimetic synthetic peptides as amyloid- $\beta$  aggregation inhibitors, S. Deike, S. Rothemund, B. Voigt, S. Samantray, B. Strodel, W.H. Binder, *Bioorganic Chemistry*, 101, 104012 (2020)
- [1] Molecular dynamics simulation of protein biosurfactants, D.L. Cheung, S. Samantray, *Colloids Interfaces*, 2 (3), 39 (2018)

## CONFERENCE PRESENTATIONS

- [1] “Behaviour of intrinsically disordered proteins at liquid interfaces: insights from molecular simulations”, Nanoscale Simulators Meeting of Ireland, University of Limerick, Ireland (May 2018).

## POSTER PRESENTATIONS

- [8] “Simulation Studies of Amyloid- $\beta$  Peptide and its Interactions with Membranes and Glycosaminoglycans”, 5<sup>th</sup> Ulm Meeting on “Biophysics of Amyloid Formation”, Ulm University, Germany (Feb 2020)
- [7] “Role of physiological environments in the folding of Amyloid- $\beta$ : Insights from molecular simulations”, 3<sup>rd</sup> Düsseldorf-Juelich Symposium on Neurodegenerative Diseases, Düsseldorf, Germany (Nov 2019).
- [6] “Structure and Assembly Dynamics of Amyloidogenic Peptides in Aqueous Solution and at Liquid Interfaces”, Computer Simulation and Theory of Macromolecules, Huenfeld, Germany (Mar 2019).
- [5] “Role of physiological environments in the folding mechanism of intrinsically disordered proteins”, Biennial Meeting of the German Biophysical Society, Düsseldorf, Germany (Sep 2018).
- [4] “Behaviour of intrinsically disordered proteins at liquid interfaces: Insights from molecular simulations”, 70<sup>th</sup> Irish Universities Chemistry Research Colloquium, Queen’s University Belfast, UK (Jun 2018).
- [3] “Behaviour of Amyloidogenic Peptides at Liquid Interfaces: Insights from Molecular Dynamics Simulation”, 7<sup>th</sup> NUIG-UL conference, NUI Galway, Ireland (Apr 2017).

- [2] “Grand Canonical Transition Matrix Monte Carlo simulations for prediction of Vapour- Liquid Equilibria and Interfacial Properties of TraPPE CO2-Tip4p/2005 Water systems on Atomistically Charged Surfaces”, SUNY Buffalo, 17<sup>th</sup> CBE Graduate Research Symposium, NY, USA (Oct 2014).
- [1] “Effect of Oil to Methanol ratio on separation of fatty acids during Transesterification of Rice bran oil”, ICACE-2013, NIT Raipur, India (Apr 2013).

## WORKSHOPS

- “Computer Tutorial in Markov Modeling (PyEMMA)”, Freie Universitaet Berlin, Germany (Feb 2019).
- “CHARMM-GUI CECAM school”, EPFL campus, Lausanne, Switzerland (Oct 2018).
- “CCP5 summer school”, Lancaster University, UK (Jul 2018).
- “Physics of Life”, 49<sup>th</sup> IFF Spring School, FZ Jülich, Germany (Feb 2018).
- “Mapping 3D Objects using a single camera”, Stokes Modelling Workshop, NUI Galway, Ireland (Jun 2017).
- “State of the art in mesoscale and multiscale modelling”, CECAM-IRL, University College Dublin, Ireland (May 2017).

## SCHOLASTIC ACHIEVEMENTS

- Awarded **Aachen Institute of computational engineering science (AICES) fellowship**, RWTH Aachen University, Germany (2018).
- Awarded 120k Class C project CPU hours from Irish High End Computing Centre (ICHEC), NUI Galway, Ireland (2017-18).
- Awarded **College of Science (CoS) postgraduate research scholarship**, NUI Galway, Ireland (2017).
- Selected for **Invitational Internship Program (DAE)** at Variable Energy Cyclotron Centre, Kolkata, India (2012).
- Selected for **Summer Internship Scholarship Program**, NIT Rourkela, India (2011).
- Selected for the 2nd level of Indian National Astronomy Olympiad, India (2005).

## SKILLS

- **Scripting Languages**
  - \* Python, Bash, Objective C, Swift, Xcode IDE ●●●●●
  - \* MATLAB, C++, Fortran 2003, Aspen HYSYS, OpenMP, MPI, R, PyTorch ●●●●●
- **Visualisation and Molecular modelling tools**
  - \* VMD, QTGrace, PyMOL, GROMACS, PLUMED v2.2, CHARMM-GUI, Maestro ●●●●●
  - \* Gaussian, LAMMPS ●●●●●
- **Document Preparation and Operating Systems**
  - \* L<sup>A</sup>T<sub>E</sub>X, MS Office, Linux (Ubuntu), MacOS ●●●●●
  - \* Windows ●●●●●
- **Laboratory Equipment and Techniques**
  - \* SEM, XRD, FTIR Spectroscopy, Particle Size Analyzer, Thermogravimetric analysis ●●●●●

## REFERENCES

Name:	Prof. (Dr.) Birgit Strodel	Dr. Bogdan Barz	Prof. (Dr.) Gunnar Schroeder
Designation:	Professor	Assistant Professor	Professor
Email:	b.strodel@fz-juelich.de	b.barz@fz-juelich.de	gu.schroeder@fz-juelich.de