Tanmoy Sanyal

Scientist, Protein Design | Structural biology

About

Greater Seattle area, WA

tanmoy.7989@gmail.com

(805) 637-0375

tanmoy7989.github.io

twitter.com/hiddenvariable2

tanmoy7989

Protein Design

Rosetta AlphaFold RFDiffusion ProteinMPNN

Molecular modeling

openMM LAMMPS **GROMACS** IMP Modeller UCSF ChimeraX **PyMOL**

MI frameworks

PvTorch Tensorflow-probability PyMC3

Languages

Python (+Cython) (~20000+ lines) Fortran-90 (~1000+ lines) C++ (~4000+ lines) Bash TCI

Open Source

nbspike Integrative epitope prediction

IMP Crosslink guided domain discovery

PDB-Dev deposition Structure of SMC5/6

LAMMPS Local density potential

Education

University of California Santa Barbara

Santa Babara, CA, USA

Ph.D Chemical Engineering

Graduate emphasis in Computational Science & Engineering (Parallel Computing & Numerical methods for ODEs / PDEs)

Indian Institute of Technology Kharagpur

Kharagpur, WB, India

Masters and Bachelors of Technology (Hons.) integrated dual degree Chemical Engineering

Research Experience

2023 - Present Amgen

Senior protein design scientist

protein families.

Novo Nordisk Research Center Seattle

2022 - 2023 Protein design scientist

- Biologics design: (i) Design of mini-proteins and antibodies for multiple therapy areas using Rosetta and AlphaFold / RFDiffusion / ProteinMPNN pipelines (ii) SAR analysis based on protein language models. (iii) Optimizing non-canonical modifications to peptides for half-life protraction.
- Scientific software development: Worked closely with AWS to re-tool AlphaFold and deploy a faster cloud-native in-house version for robust and scalable (~100k sequences) structure prediction; reguarly applied now in variant design within the company.
- **Leadership**: (i) Program lead for refining protein interface prediction ML models in collaboration with the Institute for Protein Design at UW. Seattle. (ii) Led and evaulated a proof of concept for onboarding virtual reality based structural rendering software among collaborative teams of computational design scientists and synthetic chemists.

Department of Bioengineering, University of California San Francisco Postdoctoral scholar, Andrej Sali lab

2019 - 2022

2013 - 2018

2008 - 2013

- Protein structure modeling using chemical crosslinks: Homology models + chemicalcrosslinks + cryo-EM data for protein complex structure modeling; graph sampling methods to extract domain information from crosslinks. Applications include SMC and DNA helicase
- Nanobody biophsyics: Developed integrative docking scores for nanobody docking on target receptors (SARS-CoV-2 and variant spike proteins) using chemical crosslinking and escape mutation data. Developed design principles for increasing nanobody thermostability.
- Whole cell models: Developed Bayesian Network models to combine continuum scale (pharmacokinetic), particle scale (Brownian dynamics) and network scale (enzyme pathways) models of the pancreatic beta-cell into build a proof-of-concept multimodal digitial twin. Served as software lead within the Pancreatic Beta-Cell consortium (collaborating across UCSF, USC, ShanghaiTech and HUJI)

Research Interests

Protein language models

Integrative structure modeling

Non-canonical amino acids

Nanobodies

Forcefield development

Chemical Engineering, University of California Santa Barbara

PhD candidate, Shell lab

• Coarse-graining algorithms for bulk liquids: Introduced the local density potential for structurally and thermodynamically accurate coarse-grained MD simulations of liquid-liquid phase separation, which are relatively rare in the chemical engineering literature

- Coarse-graining algorithms for protein folding using variational inference techniques: Developed protein backbone models for (i) template-free folding of 200+ residue coarse-grained protein domains using MD simulations, and (ii) studying self-assembly in amyloidogenic peptides
- **High Performance Computing**: As system administrator, set up 144 core Linux (Rocks 6.2) cluster and wrote utilities for automating (i) network-attached-storage management, and (ii) conflict-free compute resource sharing between lab members.

Volunteer

Peer-review 2019 - Present

Structure, Journal of Physical Chemistry, Proteins: Structure, Function and Bioinformatics, Review of Scientific Instruments, Rapid Reviews Covid-19, Life

Graduate Simulation Seminar Series, University of California Santa Barbara

2014 - 2016

2013 - 2018

Co-founder

Conception, organization, public outreach and long-term funding support from ChE, Materials Science and Computer Science departments.

Selected Publications

Highly synergistic combinations of nanobodies that target SARS-CoV-2 and are resistant to escape F.D. Mast, P.C. Fridy, N.E. Karen, J. Wang, E.Y. Jacobs, J.P. Olivier, T. Sanyal, et al, *eLife*, 2021

Bayesian metamodeling of complex biological systems across varying representations B. Raveh, L. Sun, K.L. White, T. Sanyal, et al, *PNAS*, 2021

Integrative analysis reveals unique structural and functional features of the Smc5/6 complex F.D. Mast, P.C. Fridy, N.E. Karen, J. Wang, E.Y. Jacobs, J.P. Olivier, T. Sanyal, et al, PNAS, 2021

A hybrid, bottom up, structurally-accurate, Go-like coarse-grained protein model T. Sanyal, J. Mittal & M. Scott Shell, *JCP*, 2019

Transferable coarse-grained models of liquid-liquid equilibrium using local density potentials optimized with the relative entropy T. Sanyal & M. Scott Shell, *JPC-B*, 2018

Coarse-grained models using local-density potentials optimized with the relative entropy: Application to implicit solvation T. Sanyal & M. Scott Shell, *JCP*, 2016

Recent conference presentations

7th Annual Computational Drug Development for Biologics Summit, Boston, MA, USA (Talk): Unpacking a workflow for automating protractor placement on peptides for enhanced half life, Oct-2022

Indian Biophysical Society Meeting, Navi Mumbai, India (Talk): Integrative modeling of higher order complexes from binary binding modes, Apr-2022

Biophysical Society Annual Meeting, San Francisco, CA, USA (Talk): Rigid body assignment in integrative determination of protein complex structures, Feb-2022

PDB50: A special symposium celebrating the 50th anniversary of the Protein Data Bank, (virtiual) (Poster): Integrative modeling of the SMC5/6 complex, May-2021

USC Brige Institute - Pancreatic Beta Cell Consortium Retreat, Catalina Island, CA, USA (Workshop): Meta-modeling: Probabilistic graphical models and your data, Nov-2019

Berkeley Mini Stat Mech Meeting, Berkeley, CA, USA

(Poster): Coarse-grained models for protein folding and self-assembly with the relative entropy, Jan-2019

Awards

44th Indian Biophysical Society Meeting	
Technopreneurship Promotion Programme Ministry of Science and Technology, Govt. of India	2009
Seed money for prototype scale-up of bioreactors for algal biodiesel generation	

2009

Jagdis Bose National Science Talent Search (JBNSTS) fellowship