Tanmoy Sanyal

Scientist, Protein Design || Structural biology

About	Education		
San Francisco Bay Area, CA tanmoy.7989@gmail.com	University of California Santa Barbara Santa Babara, CA, USA	2013 - 2018	
(805) 637-0375	Ph.D Chemical Engineering Graduate emphasis in Computational Science & Engineering (Parallel Computing & Numerical methods for ODEs / PDEs)		
tanmoy7989.github.iotwitter.com/hiddenvariable2	Indian Institute of Technology Kharagpur Kharagpur, WB, India	2008 - 2013	
C tanmoy7989	Masters and Bachelors of Technology (Hons.) integrated dual degree Chemical Engineering		
Protein Design	Research Experience		
Rosetta AlphaFold	Amgen Senior protein design scientist	2023 - Present	
RFDiffusion ProteinMPNN	Novo Nordisk Research Center Seattle Protein design scientist	2022 - 2023	
Molecular modeling openMM LAMMPS GROMACS IMP Modeller UCSF ChimeraX PyMOL ML frameworks	 Biologics design: (i) Design of mini-proteins and antibodies for multiple therap Rosetta and AlphaFold / RFDiffusion / ProteinMPNN pipelines (ii) SAR analysis protein language models. (iii) Optimizing non-canonical modifications to peptid protraction. Scientific software development: Worked closely with AWS to re-tool AlphaF a faster cloud-native in-house version for robust and scalable (~100k sequence prediction; reguarly applied now in variant design within the company. Leadership: (i) Program lead for refining protein interface prediction ML mode collaboration with the Institute for Protein Design at UW, Seattle. (ii) Led and e proof of concept for onboarding virtual reality based structural rendering softw collaborative teams of computational design scientists and synthetic chemists. 	s based on les for half-life Fold and deploy es) structure els in evaulated a	
PyTorch Tensorflow-probability PyMC3	Department of Bioengineering, University of California San Francisco Postdoctoral scholar, Andrej Sali lab	2019 - 2022	
Languages Python (+Cython) (~20000+ lines) Fortran-90 (~1000+ lines) C++ (~4000+ lines) Bash TCL Open Source nbspike Integrative epitope prediction IMP Crosslink guided domain discovery	 Protein structure modeling using chemical crosslinks: Homology models + cl crosslinks + cryo-EM data for protein complex structure modeling; graph samp to extract domain information from crosslinks. Applications include SMC and D protein families. Nanobody biophsyics: Developed integrative docking scores for nanobody do receptors (SARS-CoV-2 and variant spike proteins) using chemical crosslinking mutation data. Developed design principles for increasing nanobody thermosta Whole cell models: Developed Bayesian Network models to combine continuu (pharmacokinetic), particle scale (Brownian dynamics) and network scale (enzy models of the pancreatic beta-cell into build a proof-of-concept multimodal dig Served as software lead within the Pancreatic Beta-Cell consortium (collaborat UCSF, USC, ShanghaiTech and HUJI) 	ling methods DNA helicase cking on target and escape ability. um scale me pathways) gitial twin.	
PDB-Dev deposition Structure of SMC5/6			

LAMMPS Local density potential

Research Interests

Protein language models

Integrative structure modeling

Non-canonical amino acids

Nanobodies

Forcefield development

Chemical Engineering, University of California Santa Barbara

- **Coarse-graining algorithms for protein folding using variational inference techniques**: Developed protein backbone models for (i) template-free folding of 200+ residue coarsegrained 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

2013 - 2018

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 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

IUPAB best oral presentation

la	gdis Bose National Science Talent Search (JBNSTS) fellowship
See	ed money for prototype scale-up of bioreactors for algal biodiesel generation
	chnopreneurship Promotion Programme inistry of Science and Technology, Govt. of India
44	th Indian Biophysical Society Meeting

2009

2009