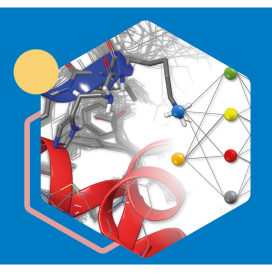


The 2nd Nordic Conference on Computational Chemistry

18-19 March 2025



Preliminary program

Tuesday 18 th March	
09.15 - 10.00	Registration
10.00 - 10.10	Welcome
10.10 – 12.10 10.10 – 10.50 10.50 – 11.30 11.30 – 12.10	 Session 1: Chair: Simone Fulle, Novo Nordisk Title to be announced, Charlotte Deane, Oxford University Computational biophysics at the proteome scale, Kresten Lindorff-Larsen, Copenhagen University An integrative approach to study intrinsically disordered proteins: From solution behavior to interaction with surfaces, Marie Skepö, Lund University
12.10 - 12.40	Poster Flash presentations
12.40 - 13.40	Lunch
13.40 - 15.00 13.40 - 14.20 14.20 - 15.00	 Session 2: Chair: Jens Carlsson, Uppsala University Current Developments in Chemical Space Exploration and Molecular Design, Mathias Rarey, University of Hamburg Trends in GPCR drug discovery, David Gloriam, Copenhagen University
15.00 - 15.20	Selected talk: Discovery of small molecule modulators of FZD ₇ using in silico docking screens, Magdalena M. Scharf, Karolinska Institutet
15.20 - 16.20	Break & poster session
16.20 – 17.20 16.20 – 16.40	Selected talk: The ribosome lowers the entropic penalty of protein folding, Julian O. Streit, University College London
16.40 - 17.00	Innovative computational drug design strategies at the intersection of automated molecule generation, machine learning, and free energy calculations, Aniket Magarkar, Boehringer Ingelheim Pharma GmbH & Co
17.00-17.20	Finding better drugs: advice for optimizing multi-parameter optimization scores, Jenny Viklund, AstraZeneca
19.00 –	Dinner

Wednesday 19 th March	
08.30 - 09.00	Entrance to conference
09.00 - 10.20 09.00 - 09.40 09.40 - 10.20	 Session 3: Chair: Ruth Brenk, University of Bergen Engineering Molecules to Specification with Generative AI, Rocio Mercado, Chalmers Novel Technologies Impacting Drug Discovery across Modalities at Lundbeck, Henrik Keränen, Lundbeck
10.20 - 11.10	Break & poster session
11.10 - 11.50	• The holy grail of generative AI – from hit to candidate drug, Eva Nittinger, AstraZeneca
11.50 - 12.10	Selected talk: Bimodal Data Fusion of Compound Structures and Cell Morphology Features for Al-driven Mechanism of Action Prediction in Cancer Drug Discovery, Osheen Sharma, Karolinska Institutet
12.10 - 13.15	Lunch
13.15 – 14.35 13.15 – 13.55 13.55 – 14.35	 Session 4: Chair: Ola Engkvist, AstraZeneca Exploring Chemical Space with Assembly Theory & Chemputation, Lee Cronin, University of Glasgow Computational studies of permeation enhancers for orally administered peptide drugs, Per Larsson, Uppsala University
14.35 - 15.00	Poster Award and closing
15.10 -	Tour of AstraZeneca

Sponsors







