Computational Approaches to the Analysis of Biomolecular Sequences, Structures and Their Functions and Applications to Biotechnology and Clinical Data Studies (26- 27 Mar 2020)

Organizing Commettiee

Co-chairs

Igor N. Berezovsky (Bioinformatics Institute, A*STAR) Lars Nordenskiöld (Nanyan Technological University) Frank Eisenhaber (Bioinformatics Institute, A*STAR)

Members

Zhang Louxin (National University of Singapore) Choi Kwok Pui (National University of Singapore) Day 4. Thursday, March 26, 2020 Coarse-Graining in Molecular Simulations and Reconstruction of Biomolecules and their Assemblies

Morning Session 9:30 – 12:00 Chair: Lars Nordenskiöld

9:30 - 10:30

Lanyuan Lu, Nanyang Technlogical University, Singapore Talk "Constructing protein structure ensemble using solution experimental data"

ABSTRACT

Developing algorithms for identifying structure ensembles of protein systems on the basis of small angle X-ray scattering or nuclear magnetic resonance is an emerging research field due to the recent experimental technological advances. The determination of protein structure ensemble from those solution experimental data is usually a highly ill-posed problem and suffers from the over-fitting issue. We systematically investigated the problem using combined molecular simulation and experimental approach, and it was discovered that a well-defined structure ensemble based on low-resolution scattering data can only be obtained from a few low energy structures. A similar conclusion was also achieved using paramagnetic relaxation enhancement data and computer simulation. Results on proteins containing multiple domains and flexible links, such as the non-structural proteins of dengue virus, will be discussed.

11:00 - 12:00

Peter J. Bond, Bioinformatics Institute, A*STAR, Singapore Talk "Multiscale modelling approaches to large systems in biology: understanding infectious diseases and the host response"

ABSTRACT

Infectious diseases due to bacterial and viral pathogens are a significant threat to human health. The interaction and assembly of such pathogens with host cells is critical to infection, the immunological response, and therapeutic interventions. We use multiscale simulations and integrative modelling approaches to investigate these processes, from atomic resolution to highly simplified coarse-grained levels. Here, I will give an overview of this research, with a particular focus on: (1) flavivirus dynamics during its infective life cycle; and (2) mammalian immune receptor responses upon exposure to bacterial pathogens.

For the former, we especially study dengue virus, responsible for more than 400 million infections per year. In serious cases, the virus can cause dengue hemorrhagic fever and shock syndrome, often facilitated by the host via antibody-dependent enhancement. In order to fill existing gaps in our molecular understanding of the viral life cycle, we have been integrating structural, biophysical, and genomic experiments with multiscale modelling, towards the "virtual dengue virus". We have reconstructed and refined the complete dengue envelope in near-atomic resolution, under different states of maturation. These virtual viruses serve as a platform for following mechanistic stages along the life cycle, such as endosomal membrane remodeling and fusion, and enabled us to identify how virus particles interact with antibodies to modulate or even facilitate infectivity, with important consequences for vaccine development.

For the latter, we are interested in the mechanisms by which endotoxic molecules released from bacterial pathogens over-stimulate immune receptors of the innate host defense system, which can lead to sepsis, a condition that kills millions of people each year. To better understand this process, we have developed computational models to trace in atomic and molecular detail the cascades and intermediates associated with transfer of endotoxins from bacterial surfaces to host immune receptors. We have also leveraged these models, to establish previously undisclosed modes of action of anti-endotoxic peptides that occur naturally during wound healing. Collectively, this work has helped to unravel key determinants governing interactions between bacterial pathogens and the host, and should help towards the search for novel therapeutics to tackle ongoing issues of antimicrobial resistance.

Afternoon Session 13:30 – 16:15 Chair: Lars Nordenskiöld

13:30 - 14:30

Alexander Lyubartsev, Stockholm University, Sweden *Talk "Multiscale modeling of macromolecular systems by structure-based coarse-graining"*

ABSTRACT

Molecular simulations of many phenomena related to biomolecular systems, soft matter and nanomaterials requires consideration of length scales above 10 nm and time scales longer than 1 mks, which necessitates the use of coarse-grained (low resolution) models, when each site of the model represents a group of atoms, and the solvent is often omitted. While many of coarse-grained models used in different studies in recent years relay on empirically parametrized interaction potentials, the systematic structure-based coarse-graining approach is based on determination of coarse-grained potentials from atomistic (high resolution) simulations.

Here a multiscale modeling approach based on the inverse Monte Carlo method is presented, in which radial distribution functions (RDF) and distributions of internal degrees of freedom of molecular structure, obtained in high-resolution atomistic simulations, are used to reconstruct effective potentials which reproduce the same structural properties within low-resolution coarse-grained model. The statistical-mechanical equations expressing canonical properties such as RDFs in terms of potential parameters can be inverted and solved numerically according to the iterative Newton scheme. The approach is illustrated on several examples of varying complexity: ionic solution; ionic liquids, coarse-grained lipid model, coarse-grained DNA model and nucleosome core particles. It is demonstrated further how effective potentials, derived exclusively from atomistic simulations, can be used to model such phenomena as lipid self-assembly, formations of vesicles and other ordered structures at varying lipid composition and concentration of different components, modeling of long DNA fragments at varying ionic conditions. A software package MagiC implementing the inverse Monte Carlo method for computation of effective potentials for coarse grained models of arbitrary structure from atomistic trajectories is presented.

Ganesh Srinivasan Anand, Department of Biological Sciences, National University of Singapore, Singapore *Talk "Whole Viral Particle Metastability and Dynamics by HDX Mass Spectrometry"*

ABSTRACT

Viruses are metastable macromolecular assemblies that undergo reversible conformational rearrangements in solution referred to as 'breathing' that are critical for sensing host-specific environmental cues, receptor interactions and for initiating host entry. Host specific environmental cues including temperature, osmolyte and pH trigger large changes in viral assembly leading to formation of disassembly intermediates and eventual disassembly with release of genomic material. Amide hydrogen deuterium exchange mass spectrometry (HDXMS) is a powerful tool for measuring hydrogen bonding propensities and solvent accessibility of proteins and protein complexes. This makes it especially suitable for probing the breathing dynamics, measurement of quaternary contacts of viral particles in solution as well as to probe the effects of hostspecific conditions and for mapping epitopes and paratopes of neutralizing antibodies. In this talk, I will present a short overview of mass spectrometry applications in structural biology for mapping protein assemblies and interfaces with lipids and nucleic acids with dengue and zika viral dynamics as the focus. Our HDXMS results show that each dengue/zika serotype/strain display a distinct breathing dynamics profile with an entirely different expansion response to temperature. At 37 °C, DENV serotype 2 strains show temperature-specific changes with the biggest change at the Envelope E- intradimeric interface. However, at 40 °C, a different set of temperature-specific loci were detected in strains of DENV serotype 1. These unique serotype dynamic profiles underscore the importance of protein guaternary contacts, packing of lipid bilayer and RNA genome in each viral particle. HDXMS has also enabled detailed maps of epitope and paratopes for 2 (1 stabilizing and 1 destabilizing) antibody-dengue virion complexes. I will conclude by describing how HDXMS can reveal the role of genomic RNA in viral stability and dynamics in a model plant RNA virus, Turnip Crinkle Virus (TCV).

Firdaus Samsudin, Bioinformatics Institute, A*STAR, Singapore Selected short talk, "To bind or not to bind: IgM versus IgG in Pertuzumab and Trastuzumab"

Firdaus Samsudin^a, Joshua Yi Yeo^a, Samuel Ken-En Gan^a, and Peter J. Bond^{a,b} ^aBioinformatics Institute, A*STAR, Singapore ^bNational University of Singapore, Singapore

ABSTRACT

Antibodies are classified into five major immunoglobulin (Ig) isotypes and they can have different numbers of antigen binding sites. For example, IgG has two antigen binding sites, whereas IgM has up to twelve. While the former is the isotype of choice for most currently available therapeutic antibodies, the latter has superior complement-activating and cell-agglutinating abilities. Utilizing IgM is therefore an attractive option for future immunotherapy. But is having more antigen binding sites always better? In this study, we performed an integrative multiscale modelling and simulation of two well-known monoclonal antibodies used to treat human epidermal growth factor receptor 2 (HER2)positive breast cancer: Pertuzumab and Trastuzumab. We found that only Pertuzumab IgM can utilize all of its twelve antigen binding sites to bind to HER2 extracellular domain, while similar binding in Trastuzumab IgM is hindered by steric clashes. This is subsequently validated by cell count assay showing that Pertuzumab IgM is more efficient at inhibiting the proliferation of HER2-overexpressing cells compared to its IgG counterpart and to Trastuzumab IgM. Our study highlights the importance of understanding the molecular details of antigen-antibody interaction for the design and isotype selection of therapeutic antibodies.

Day 5. Friday, March 27, 2020 Biomedical Modelling, Informatics, and Clinical Data Studies

Morning Session 9:30 – 12:00 Chair: Frank Eisenhaber

9:30 - 10:30

Gerhard Grüber, School of Biological Sciences, Nanyang Technological University, Singapore *Talk "Amino acid sequence insertions or deletions inside Mycobacterium tuberculosis's respiratory complexes provide fitness for the pathogen and new pathways for TB-drug development"*

ABSTRACT

The causative agent of Tuberculosis (TB) *Mycobacterium tuberculosis* (*Mtb*) encounters unfavourable environmental conditions in the lungs, including nutrient limitation, low oxygen tensions and/or low/high pH values. These harsh conditions in the host triggers *Mtb* to enter a dormant state in which the pathogen does not replicate and uses host-derived fatty acids instead of carbohydrates as an energy source. Independent to the energy source, the bacterium's energy currency ATP is generated by oxidative phosphorylation, in which the F₁F₀-ATP synthase uses the proton motive force generated by the respiratory complexes, including two terminal oxidases: a proton-pumping cytochrome *bc*₁:*aa*₃ oxidase supercomplex and a non-proton pumping, less energetically efficient, cytochrome *bd* oxidase. The later reveals regions and amino acid residues unique to mycobacteria, opening a door for TB-specific compound design.

A special feature of the mycobacterial F-ATP synthase is its inability to establish a significant proton gradient during ATP hydrolysis, and its latent ATPase activity, to prevent energy waste and to control the membrane potential. Recently, we have identified unique epitopes of mycobacterial F_1F_0 -ATP synthase subunits absent in their prokaryotic or mitochondrial counterparts which contribute to the regulation of the low ATPase activity. Most recent structural insights into individual subunits, the F_1 domain or the entire mycobacterial enzyme added to the understanding of mechanisms, regulation and differences of the mycobacterial F_1F_0 -ATP synthase compared to other bacterial and eukaryotic engines. Novel insights providing the basis for the design of new compounds targeting this engine and even novel regimens for multidrug resistant TB will be presented.

11:00 - 12:00

Lim Soon Wong, National University of Singapore, Singapore Talk "A genome assembly quality assessment measure that integrates contiguity, completeness, and correctness"

ABSTRACT

The quality of a genome assembly is assessed by a variety of measures (e.g. NGA50, number of known core genes mapped, and number of translocation). These typically consider the contiguity, completeness, and correctness of the assembly separately and independently. It is often the case that one assembly is slightly superior to another assembly in some measures and is slightly worse in other measures. Thus one cannot easily tell which assembly is better. Indeed, even when one assembly is vastly superior to the other, these measures often suggest very small differences (e.g. 1-2%) between the assemblies. In this talk, I will describe a genome assembly quality assessment measure called PDR (pairwise distance reconstruction). The PDR is based on the average difference between the observed distance of two loci on an assembly and the actual distance on the reference genome. In other words, PDR determines how well the assembly can be used to inform on the distance between two arbitrary loci in the actually genome. It is easy to see that the more contiguous, complete, and correct an assembly is, the better it can inform on the distance between two arbitrary loci on the actual genome. Thus the PDR naturally generalizes and integrates these properties. As the PDR is averaged over all possible pairs of positions on a genome, it is impractical to compute it in a naïve manner for large genomes. Fortunately, an approximation to the PDR that is fast to compute accurately (error is less than E-10) is possible using piecewise integrals. I will show in this talk that this makes the PDR a very practical informative genome assembly quality assessment measure. This talk is based on the PhD thesis of my student Xie Luyu.

Afternoon Session 13:30 – 16:15 Chair: Frank Eisenhaber

13:30 - 14:30

Mahsa Paknezhad, Bioinformatics Institute, A*STAR, Singapore *Tutorial "*Reinforcement learning and its biomedical applications"

ABSTRACT

Reinforcement learning is a machine learning method that is gaining growing popularity in different areas of research. This talk will present an overview of multiple reinforcement learning algorithms. We will introduce Q-learning and policy gradient as two most widely known reinforcement learning methods. We will explain how these techniques are effectively utilized to address problems in biomedical data and image domain.

Liang Kaicheng, Bioinformatics Institute, A*STAR, Singapore Tutorial "Computational imaging with deep learning: biomedical focus"

ABSTRACT

Modern approaches to artificial intelligence using deep learning have achieved significant milestones in computer vision. Beyond image classification and analysis, AI techniques for imaging and image generation have also shown great promise. This talk will review the state of the art in generative AI for computational challenges such as super-resolution and style transfer, with a focus on biomedical applications and microscopy. Future opportunities and potential pitfalls will also be discussed.

Rohan Williams, Singapore Centre for Environmental Life Sciences Engineering (SCELSE), National University of Singapore, Singapore Selected short talk "Sequence similarity networks: a working model for defining the 'known-unknowns' of gene function in microbiomes"

ABSTRACT

It is well understood that a substantial proportion of predicted genes are classifiable as having no known function. Current approaches to defining gene function are highly dependent on comparative genomic methods and the use of reference databases such as KEGG. However this kind of approach will by nature eliminate genes of unknown function from consideration, and this becomes particularly limiting when considering the functional repertoire of member species of complex microbial communities (microbiomes). Here I discuss some recent work that uses sequence similarity networks to provide a working model for organising functional genes irrespective of whether their biological function is currently understood or not. Briefly protein sequence from genes predicted within metagenome assemblies are subjected to all-versus-all protein BLAST, and analysed as a network: I will discuss the validation of these networks against known functional databases (KEGG) and how we have applied this concept to the interpretation of gene catalogues from metagenome assemblies. I will illustrate this approach using metagenome assemblies obtained from bioreactor communities enriched for anaerobic ammonium oxidising bacteria (AnAOB), directed toward the ongoing search for the key genes underpinning the biochemistry of the anaerobic ammonium oxidisation reaction (ANAMMOX).