

imaging across 5 orders of magnitude in length, we are able to both find these rare fusion events, and study a heterogeneous population of osteoclasts for a contextualized understanding of this important cell type. The insights gained in this work explore fundamental features of cell fusion required for development of successful next generation biomaterials that cater to basic osteoclast biology requirements and ensure better replication of the *in vivo* environment.

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Optimizing Excitation Polarization to Probe Fusion Pore Dynamics using TIRF Microscopy

Kasey Hancock^{1,2}, Joerg Nikolaus³, Erdem Karatekin³, David Baddeley⁴.

¹Yale University, New Haven, CT, USA, ²Truman State University, Kirksville, MO, USA, ³School of Medicine, Cellular and Molecular Physiology, Molecular Biophysics and Biochemistry, Yale University, West Haven, CT, USA, ⁴Department of Cell Biology, School of Medicine, Yale University, New Haven, CT, USA.

During hormone or neurotransmitter release via exocytosis, fusion pores may open and close (flicker) repeatedly before resealing (“kiss & run” fusion) or dilating (full fusion) irreversibly. Pore dynamics regulate the amount and kinetics of cargo release, and determine the mode of recycling, but mechanisms that govern pore dynamics are not understood. This is in large part due to a lack of reconstituted assays with single-pore sensitivity and millisecond time resolution. We recently described a polarized total internal reflection (pTIRF) microscopy assay to monitor fusion of proteoliposomes to planar lipid bilayers supported on a soft polymer cushion with single molecule sensitivity and ~15 ms temporal resolution. Fluorescently labeled small unilamellar vesicles, reconstituted with exocytotic/neuronal v-SNAREs (v-SUVs), fuse with a supported bilayer containing the cognate t-SNAREs (t-SBL). Each fusion event is accompanied by changes in the total fluorescence intensity surrounding the fusion site, as the lipid-linked labels diffuse from the liposome into the supported bilayer through the fusion pore. Analysis of the intensity changes, combined with a mathematical model, provides information on pore dynamics (Stratton et al. *Biophys. J.* 2016). In principle, three factors can contribute to intensity changes upon fusion: 1) dequenching of fluorophores, 2) evanescent field decay and 3) a change in the average orientation of the fluorophore dipole moments with respect to the excitation polarization field, as the fluorophores move from the round liposome into the flat bilayer. Here, we systematically varied the polarization of the excitation field and quantified its contribution to intensity changes for different lipid-linked fluorophores. Large increases facilitate detection of fusion events and quantification of lipid release kinetics.

Membrane Structure I

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Analysis of Lipid Domains in Bilayer Simulations using Observables for Lipid Packing

Soohyung Park, Wonpil Im.

Biological Sciences, Lehigh University, Bethlehem, PA, USA.

Lipid rafts are micro- or nano-sized dynamic domains of membranes enriched in cholesterol, glycosphingolipids (and other saturated lipids), and specific membrane proteins, which are involved in many important biological processes such as membrane trafficking, signaling, protein sequestration, and so on. Though the existence of lipid rafts has been widely accepted, the study of their dynamics (size and lifetime) is still challenging due to the limitations in experimental techniques. In this context, computational studies can be a useful approach that complements experiments and provides insight into the properties and dynamics of lipid rafts by monitoring and analyzing membrane structures at the atomic resolution. So far, the analysis of lipid domains in computational studies has been mostly based on the order parameter and neighbor composition of lipids, which could be complicated in the analysis of multicomponent bilayers. To address this issue, we propose a simple yet robust method for the analysis of lipid domains in bilayer simulations using physically transparent observables for lipid packing, area and thickness. In our method, the ordered states of lipids are inferred by a hidden Markov model analysis. Then, the ordered state map onto the Voronoi tessellation of lipids is analyzed using the Getis-Ord local spatial autocorrelation statistics to obtain clusters of ordered lipids (lipid domains). The method is general and can be applied to various situations in that 1) it does not require information other than the mechanical properties of lipids to assign their ordered states, and 2) the spatial autocorrelation statistics allows a robust assignment of the clusters of ordered and

disordered regions. The usefulness of the method is illustrated by analyzing the concentration effects on the membrane order for the bilayers consisting of 1) ganglioside GM1 and 1-palmitoyl-2-oleoyl-*sn*-glycero-3-phosphocholine (POPC), and of 2) cholesterol, 1,2-dimyristoyl-*sn*-glycero-3-phosphocholine (DMPC), POPC, 1,2-dimyristoyl-*sn*-glycero-3-phosphoethanolamine (DMPE), and 1-palmitoyl-2-oleoyl-*sn*-glycero-3-phosphoethanolamine (POPE).

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Software for Direct Comparison of Membrane Scattering Experiments Data to Molecular Dynamics Simulations

Yevhen Cherniavskiy¹, Svetlana Baoukina¹, Bryan W. Holland^{1,2}, Norbert Kučerka^{3,4}, Peter Tieleman¹.

¹Biological Sciences, University of Calgary, Calgary, AB, Canada, ²Simulations Plus, Inc., Lancaster, CA, USA, ³Frank Laboratory of Neutron Physics, Joint Institute for Nuclear Research, Dubna, Russian Federation, ⁴Department of Physical Chemistry of Drugs, Comenius University, Bratislava, Slovakia.

Detailed information on lipid bilayers structure is very important for understanding the processes in cellular membranes. Nowadays X-ray and neutron scattering experiments are widely used as a source of lipid bilayers structural parameters (area per lipid, thickness, etc.), but in contrast to scattering patterns from crystal structures, lipid membranes produce very broad peaks in scattering intensity profiles due to their fluid nature. This makes the process of retrieving structural details indirect and introduces approximations to convert experimental structure factors to electron or neutron scattering profiles. On the other hand, molecular dynamics simulations are able to provide direct sub-atomic level information on lipid bilayers structure, but can produce unreliable data because of inaccuracy of the force fields. Simulation to Experiment (SIMtoEXP) software is designed to facilitate direct, model free, comparison of MD simulations to experimentally measured structure factors for lipid bilayers. We present SIMtoEXP version 2.0 - an updated version of the original software. The new version of this program is written in C++ using the Qt GUI library, instead of the original C/Perl implementation. New features, including a MD trajectory reading module, were added to simplify the workflow for users. In future releases SIMtoEXP software will support structure factor calculations from simulations with the coarse-grained Martini force field. The Martini force field provides significant computational speedup while retaining significant chemical detail of the lipids. This allows simulating large lipid-protein aggregates on longer time-scales and investigate the structure and lateral organization of cell membranes. Incorporation of Martini support in SIMtoEXP will allow direct comparison of coarse-grained simulations to experiments, which will benefit both Martini force-field parametrization and the interpretation of experimental results.

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Modeling Ethers with Molecular Dynamics: Updated CHARMM Force Field Parameters for Ethers in Model Compounds and Lipid Membranes

Alison M. Leonard.

UMCP, Crofton, MD, USA.

Small ether molecules such as polyethylene glycol (PEG) have extensive industrial and medical applications. Additionally, phospholipids containing ether linkages make up 30% of the glycerophospholipids in the human brain. Preliminary research suggests that the CHARMM All-Atom Additive Force Field does not accurately recreate experimental results for bilayers composed of ether lipids such as 1,2-di-O-hexadecyl-*sn*-glycero-3-phosphocholine (DHPC). 100-ns simulations of pure DHPC bilayers do not reproduce essential target data such as surface area per lipid and electron density profile, which describe the structural properties of the membrane. Additionally, the initial parameterization of ethers tested a limited population of linear ethers and did not accurately reproduce potential energy scans about the O-C-C-O dihedral (*J. Chem. Theory Comput.*, **3**(3):1120 - 1133). We have used the MP2 density and Dunning diffuse aug-cc-pVQZ basis sets to compute the charge distributions of model ethers, including various lengths of PEG. We have also used Hybrid Methods for Interaction Energies (HM-IE) to compute potential energy scans about the O-C-C-O angle effectively at the CCSD(T)/aug-cc-pVQZ level (*J. Phys. Chem. A*, **108**: 107 - 112). We found that the charge on ether oxygens is substantially more negative than the current CHARMM force field suggests. Adjusting this charge allows more water to penetrate a DHPC bilayer, increasing the surface area per lipid and improving the bilayer's structural representation. Additionally, our new dihedral parameters for the central O-C-C-O angle more accurately reproduce the quantum potential energy scan and bring