

presumably to a wide variety of couplings to the membrane. We provide viscosity measurements for lipid bilayers with various compositions in the gel and liquid crystalline phases.

#### 2191-Pos Board B210

##### **Mc Model of Lipid Raft Protein Diffusion Matched to Live Cell Measurements with Controlled Chemical Perturbation Experiments**

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Partitioning of certain type extra-cellular or cytosolic leaflet membrane proteins into functioning cholesterol stabilized domains within both leaflets of the membrane structure has long been shown by various studies facilitating different techniques. We have modeled how lipid-stabilized nano-domains influence the diffusion of some membrane proteins. By combining these simulations with non-perturbative experimental measurements of the diffusion behavior for chemically unperturbed and controlled perturbation cases of that diffusion, we were able to quantify the essential parameters describing the domains. The simulations feature four independent parameters, i.e. area fraction of the membrane occupied by domains, size of the domains, probability of the proteins to exit the domains, and effective protein diffusion within the domains. Kinetic Monte-Carlo modeling of proteins' association with membrane nano-domains has been performed over a selected region of the parameter space. Some of these parameters may be modulated at the single cell level during the time of a single experiment, allowing precise matching of all four parameters. We have used our non-perturbative imaging based FCS, bimFCS, to measure the diffusion of several membrane proteins from inner and outer leaflets of the cell-membrane over multiple length scales simultaneously. This allows extracting information of the domains which transiently trap the diffusing proteins.

We studied GFP-tagged GPI anchored proteins for the external leaflet and compared measurements of monomers and induced dimers on the same cells. For the intracellular leaflet, we studied Lyn-anchored GFP and used the Rapamycin induced cross-linking of the FKBP12 and Frb domains. We were able to successfully match experimental data with in-silico modeling by Kinetic Monte Carlo simulations, giving insight about sizes and area fraction of raft domains for each leaflet of the membrane, and also affinity of associated proteins with them.

#### 2192-Pos Board B211

##### **High-Speed Interferometric Scattering Microscopy of Receptor Mobility Reveals Anomalous Diffusion in Model Membranes**

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The diffusive behavior of membrane proteins has been reported to change with the timescale of the measurement. High-speed single-particle tracking studies have revealed that it is Brownian on time scales less than 100  $\mu$ s, confined from 1 to 10 ms, and Brownian at longer times (>10 ms).<sup>1</sup> This behavior has been interpreted as hop diffusion between submicron compartments in the plasma membrane. Due to the fundamental difficulty in achieving simultaneously high spatial and temporal resolution, these results have relied on a single experimental approach. Furthermore, attempts to use a complementary fluorescence technique to study anomalous diffusion have not reproduced the hop diffusion model.<sup>2</sup>

Here we use interferometric scattering microscopy (iSCAT)<sup>3</sup> to track the motion of the GM1 ganglioside receptor binding the B subunit of cholera toxin using 40-nm gold nanoparticle labels, in both supported lipid bilayers (SLBs) and droplet hydrogel bilayers (DHBs).<sup>4</sup> In SLBs, in single trajectories containing >200,000 frames and providing simultaneous 10  $\mu$ s temporal and sub-nm spatial precision, we observe a shift from Brownian diffusion at early time scales (30 - 100  $\mu$ s) to confined diffusion (100  $\mu$ s - 10 ms) back to Brownian diffusion (>10 ms). In contrast, GM1 diffusion in DHBs is Brownian at all timescales. These data suggest that the presence of a structural support on one side of the bilayer is sufficient to produce anomalous diffusion, even for diffusers that do not penetrate the membrane.

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#### 2193-Pos Board B212

##### **Investigation of Temperature Induced Mechanical Changes in Supported Bilayers by Reconstructed Atomic Force Microscopy Tapping Forces**

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Membrane mechanics, such as rigidity, adhesion, and fluidity are known to play an important role in the regulation of many biological and cellular pro-

cesses. These properties can be drastically affected by changes in temperature; increasing temperature can enhance membrane permeability and susceptibility whereas decreasing temperature is known to restrict membrane movement and the intake of essential nutrients. Temperature also plays a key role in determining the phase of a membrane; evidence of nanoscale rearrangement of lipids into raft-like domains has been found at physiological temperatures. Rafts are thought to be a site of importance for signal transduction and membrane protein trafficking. There is even a growing association between rafts and the development of neurodegenerative diseases. Thus, it is evident that temperature has a critical role in the maintenance of membrane biology. In this work, patches of total brain lipid extract bilayer containing 30% exogenous cholesterol were formed in phosphate buffered saline and examined using atomic force microscopy while the temperature was systematically increased from 28-40°C. Tapping mode atomic force microscopy (TMAFM) and scanning probe acceleration microscopy (SPAM) were used to obtain topographical and mechanical information of the sample, respectively. SPAM is relatively new technique that offers the advantage of quickly obtaining mechanical information, while maintaining nanoscale spatial imaging resolution, due to its basis in TMAFM. The principle of SPAM is that the noisy deflection signal is captured during TMAFM imaging and reconstructed to reveal the time-resolved forces between the cantilever tip and the sample surface. These tip/sample forces can be correlated with surface properties such as rigidity and adhesion. With SPAM it is also possible to construct harmonic images, thus providing further insights on subtle surface characteristics.

#### 2194-Pos Board B213

##### **Red Blood Cell Membrane Fluctuations and their Mechanisms: Passive Versus Active**

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Red blood cells (RBCs) are soft and flexible biconcave discs, which are able to pass through capillaries with diameters several times smaller than the RBC size. The RBC deformability also results in noticeable membrane fluctuations, which must be correlated with RBC membrane properties. However, it is still under debate whether RBC membrane fluctuations are simply passive thermal undulations or whether a red cell also experiences active fluctuations which are driven by a metabolic activity or other cell processes. We will present direct evidence that the RBC undulations are not solely passive thermal fluctuations, which has been obtained from a set of different experiments and simulations using a high spatiotemporal resolution: from 10 microseconds to several seconds in time and up to 20 nanometers in space. Experimental results show a violation of the fluctuation-dissipation theorem (FDT) for freshly prepared RBCs indicating the existence of active processes. However, the FDT is satisfied for starved cells demonstrating that the membrane fluctuations are passive when the energy supply is absent. Experiments also show a considerable change in the fluctuation amplitudes for fresh and starved cells. Subsequently, we perform simulations which fully mimic and quantify the experiments. We are able to quantitatively extract RBC membrane properties including shear elasticity, bending rigidity, and membrane viscosity. Furthermore, we test several models for active fluctuations, which mimic different possible mechanisms including spectrin network remodeling, ion pumps, and change in the spontaneous membrane curvature. Simulation results agree well with experimental data and suggest that several processes mentioned above may contribute to active RBC fluctuations. We will discuss which processes are more likely to take place.

#### 2195-Pos Board B214

##### **Combined Stopped-Flow and Electrophysiological Experiments Suggest Direct Sodium Channel Inhibition by Model Fluorobenzene Anesthetics**

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General anesthetics are often proposed to affect membrane properties through interactions with the lipid bilayer. Though recent studies have shown that inhaled anesthetics can interact more specifically with certain membrane proteins such as ion channels. For example, volatile anesthetics inhibit voltage-gated sodium channels to reduce neurotransmitter release. Fluorobenzenes (FBs), once considered for clinical use, were abandoned due to their flammability and toxicity but are still valuable model anesthetics for investigating the molecular mechanisms of anesthetics. We examined the properties of four FB compounds, 1,2-DiFB, 1,4-DiFB, 1,3,5-TriFB and HexaFB on lipid bilayer