pattern can be enhanced by exposure to agonist concentrations (1-5 μM) of ryanodine in OT terminals only. RyR antagonists, 8-Br-cADPR or higher concentrations (10 μM) had the opposite effect; significantly reducing the amount of OT associated with the membrane area. Additionally, Ca²⁺-evoked NP release from permeabilized terminals was increased by agonist concentrations of ryanodine and conversely, decreased by antagonist concentrations of this drug. Agonist concentrations of ryanodine were also able to increase the asynchronous phase of low frequency electrically stimulated capacitance increases from isolated NH terminals. Thus, the ryanodine-sensitive mobilization of secretory granules seems to have a functional role in modulating secretion of neuropeptides from NH terminals. [Supported by UMass Grant P60037094000000 (SOM) and NIH Grant NS29470 (JRL)]

1583-Pos
Reactive Cysteines of Ryanodine Receptor Type 1 Influence Function and Response to Oxidative Stress
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Redox modulation of the skeletal muscle ryanodine receptor1 (RyR1) plays a key role in determining the responsiveness of the Ca²⁺ release channel to physiological modulation. The sensitivity of RyR1 to redox stress may be conferred by previously unidentified hyper-reactive cysteines (1040, 1303, 2426, 2606, 2611, 2625 and 3635). Wild type RyR1 (∼RyR1), and seven hyper-reactive cysteine mutations of RyR1 were stably expressed in HEK-293 cells and their contribution to RyR1 function evaluated. Addition of RyR1 activator 4-chloro-m-cresol (4-CMC) elicited an increase in [Ca²⁺], in the ∼RyR1 cells but failed to produce a Ca²⁺ response in the C1030S, C2606S, C2436S and C7S (expressing all seven cysteine mutations) expressing cells, while the C1040S and C2611S mutations significantly attenuated 4-CMC mediated Ca²⁺ response. Microsomal fractions isolated from C1040S, C2611S, C2436S and C3635S bound to [H]ryanodine and C1303S showed significantly lower, levels of RyR-binding, although significantly above preparations from RyR-null HEK 293. The sensitivity of RyR1 to 1, 4-naphthoquinone (NQ) appears to depend on the expression of RyR1 and the presence of reactive cysteines. Sensitivity to NQ-induced cytotoxicity was determined by the multi-Tox fluorescence assay. NQ decreased cell viability in a dose-dependent manner, but the ∼RyR1 cells were less sensitive than C2606, C1040S, C2611S and the C7S mutants. These data indicate the role of hyper-reactive cysteines in regulating RyR1 function and its response to oxidative stress. Supported by NIH AR43140.

1584-Pos
Triclosan Uncouples Excitation-Contraction Coupling in Skeletal Myotubes Without Blocking RyR1
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The chlorinated diphenylethers are a class of broad-spectrum antimicrobial agents. One of the most potent and widely used member of this group is triclosan (TCS; 2,4,4’-trichloro-2’-hydroxydiphenylether). We studied the effects of TCS in primary myotube cultures using Ca²⁺ imaging with Fluo 4 and whole-cell voltage clamp. Acute perfusion with 10 mM TCS resulted in a significant but transient elevation in cytosolic Ca²⁺ in unstimulated (resting) myotubes, an effect not seen in RyR null (dyspedic) cells. TCS caused a rapid decline in the amplitude of electrically evoked Ca²⁺ transients culminating in complete loss of Ca²⁺ transients. Upon failure of excitation-contraction (EC) coupling, RyR1 remained responsive to application of caffeine (20mM). Caffeine-induced release of SR Ca²⁺ in the presence of TCS was comparable to, or greater than, that measured in the control period indicating that the release channels remained functional and the SR stores were replete with prolonged TCS exposure. Acute submicromolar TCS (0.5 μM) enhanced Ca²⁺ transient amplitude at 0.1 Hz stimulus, whereas pre-incubation of myotubes with TCS for 24 hr was sufficient to alter the relationship between stimulus frequency and Ca²⁺ transient amplitude across the entire stimulation frequency range. TCS (10 μM) also completely inhibited depolarization-triggered extracellular Ca²⁺ entry and suppressed DHPR mediated Ca²⁺ current to that observed in dyspedic cells. These uncoupling effects were observed without any influence on the magnitude of store-operated Ca²⁺ entry (SOCE) in myotubes. These results are the first to identify that TCS (and possibly related structures) impairs EC coupling by uncoupling orthograde and retrograde signaling between RyR1 and DHPR in skeletal muscle. Supported by NIH AR0535104 (K.G.B.), AR43140 and ES011269 (I.N.P.), and MDA4319 (K.G.B.) and MDA4155 to (R.A.B.).

Motions of the Cell Surface Molecules

1585-Pos
Method for In-Vitro Studies of Cellular Interactions at the Interface of Two Tissues
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We describe a simple and reliable experimental technique that enables one to create a high fidelity linear interface between two opposing cell layers. The method employs a custom designed lid that fits a standard 3cm cell culture dish. During cell plating, the dish is divided by a 200 micron thick separator that is part of the lid. The separator is covered in a thin layer of parafilm that forms a hermetic seal with the underlying coverslip and creates a temporary gap between the two cell plating environments. After cells attach, the custom lid is replaced with a standard lid and cells are allowed to grow under standard cell culture conditions. When expanding cell layers fill the gap, a linear interface is formed between the two opposing fields. Paracrine factors released from an approaching cell front as well as direct physical and molecular interactions between two cell types affect intercellular orientation, individual cell morphology, and the degree of cells invasion into the opposing layer. The local interface appearance thus depends on a specific cell pair and may vary dramatically. We describe several types of such interfaces for different cell pairs, including cardiomyocytes, fibroblasts, melanocytes, endothelial cells and colon canceroma cell lines. The method serves as a practical in vitro tool to study cell growth and invasion that occur on the interface of two neighboring tissues.

1586-Pos
A Zoo of Dynamic Pattern Formation by Bacterial Cell Division Proteins
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Min proteins of the Escherichia coli cell division system oscillate between the cell poles in vivo. In vitro on a solid-surface supported lipid bi-layer these proteins exhibit a number of interconverting modes of collective ATP-driven dynamic pattern formation including not only the previously described propagating waves, but also near uniform in space surface concentration oscillation, propagating filament like structures with a leading head and decaying tail, and moving and dividing amoeba-like structures with sharp edges. We demonstrate that the last behavior most closely resembles in vivo system behavior. The simple reaction-diffusion models previously proposed for the Min system fail to explain the results of in vitro self-organization experiments. We propose hypotheses that initiation of MinD binding to the surface is controlled by counteraction of initiation and dissociation complexes; the binding of MinD is stimulated by MinE and involves polymerization-depolymerization dynamics; polymerization of MinE over MinD oligomers triggers dynamic instability leading to detachment from membrane.

Fig. 1. MinD (green) and MinE (red) wave transforms into running and dividing amoebas. Frames were taken every 40 s, the scale bar is 5 μm.

1587-Pos
Active Re-Modelling of Cortical Actin Regulates Spatiotemporal Organization of Molecules on a Living Cell Surface
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Cell surface proteins such as lipid-tethered GPI-anchored proteins, Ras-proteins and several glycoproteins, are distributed as monomers and nanoclusters on the surface of living cells. The spatial distribution and dynamics of formation and breakup of these nanoclusters is unusual and controlled by the active remodeling dynamics of the underlying cortical actin (CA). To explain these results, we propose a novel mechanism of nanoclustering, based on the active hydrodynamics of the CA and its coupling to local membrane composition. In addition, our theory makes a falsifiable prediction – GPI-APs must exhibit anomalous concentration fluctuations resembling those at criticality; we confirm this using a fluorescence-based assay. Our work addresses a central issue.
in cell biology, namely the nature of molecular organization and its spatiotemporal regulation on the plasma membrane.

1588-Pos
Membrane Anchor Dependent Colocalization in Cellular Membranes Observed by Fluorescence Cross-Correlation Spectroscopy
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Membrane anchors exist on many proteins in a variety of combinations of enzymatically attached fatty acids and glycosylation. These anchors play a part in protein trafficking within cells and in associating proteins with cell membranes. They are also frequently found on well-known signaling proteins. Given the variety of anchor composition, we question whether these anchors play a more significant role in the lateral sorting or dynamic colocalization of proteins within cell membranes. To observe this in vivo, we create fusion proteins of red and green fluorescent proteins with the consensus protein lipidation motif of various signaling proteins and express both red and green constructs in HEK293T cells. The dynamic colocalization of red and green fluorescent proteins, and therefore the dynamic colocalization of membrane anchors, can be directly observed using Fluorescence Cross-Correlation Spectroscopy (FCCS). FCCS allows us to observe dynamic colocalization on the nanometer length scale. Unlike FRET, FCCS can detect positive colocalization regardless of orientation and at lengths larger than 10nm. Recent results will be discussed.

1589-Pos
Growth of the E. Coli Outer Membrane
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The outer membrane (OM) of E. coli is composed of four elements: lipopolysaccharide (LPS), phospholipids, OM proteins, and lipoproteins. Together these elements form a continuous protective layer, defending the bacterium against harsh environments and toxic chemicals. Maintenance of an intact OM requires that synthesis and insertion of new OM components keep pace with bacterial growth. At present, little is known about where new OM is incorporated, or how its growth is regulated. We use video microscopy to examine the behavior of fluorescently labeled LPS and specific OM proteins on the surfaces of growing bacteria. Initially, labeled LPS and OM proteins in an individual cell exhibit a uniform peripheral distribution. As the bacterium elongates, fluorescent spots emerge, subsequently drift apart from one another, and occasionally bifurcate. Arresting bacterial growth with Rifampin halts the motion of the fluorescent spots, resulting in a fluorescence pattern which remains stable over a period of hours. We hypothesize that the appearance and divergence of these fluorescent spots of labeled OM is due to insertion of newly synthesized, unlabeled OM components. We track the motion of these spots on the surfaces of E. coli, and measure the convergence and divergence of adjacent tracks on the periphery of the cell. Our data suggest that new OM is incorporated in patches and distributed non-uniformly, with the bulk of the new material inserted along the lateral walls of the cell and lower rates of insertion in the poles of the cell.

1590-Pos
Single-Molecule Study of the Dynamics of Lipid-Like Molecules in the E. Coli Outer Membrane
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While there have been many studies on the diffusion of membrane lipids in eukaryotic cells, which have given insight into the structure and organization of these membranes, little is known about their role in bacterial membranes, specifically the Gram negative bacteria, Escherichia coli. The E. coli outer envelope consists of inner and outer lipid membranes that are separated by a periplasmic space containing the cell wall. The outer membrane is unique in that it is thinner than mammalian plasma membranes and consists of a phospholipid inner leaflet with a predominantly lipopolysaccharide (LPS) outer leaflet. Here we look at the diffusion of the fluorescent lipid analog 3,3-dioctadecyldimethylammonium iodide (DiI(C16)) and Alexa488-LPS in the outer membrane of live E. coli cells using single molecule imaging/tracking techniques. The diffusion coefficient of DiI(C18) was found to be (3.2 ± 0.2) \times 10^{-11} \text{cm}^2/\text{sec} time scales of 0.33 sec. By contrast, the diffusion coefficient of DiI(C18) in human colon cancer cells of the nasopharynx (KB) is found to be (1.94 ± 0.2) \times 10^{-8} \text{cm}^2/\text{sec}, in good agreement with previously measured diffusion coefficients of DiI(C18) in other mammalian cells. The mobility of LPS in the outer membrane and the implications of the slow diffusion of DiI(C16) on the structure of the outer membrane of E. coli will be discussed.

1591-Pos
Investigation of the Confining Potential of Toxin Receptors in Membrane Microdomains by Single Molecule Tracking with Lanthanide-Doped Nanoparticles
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We coupled costable and non-blinking Yb0.4Eu0.6VO4 nanoparticles to epsilon toxins produced by Clostridium perfringens type B and D, which bind to a specific receptor on MDCK cells. Single-molecule tracking using these labels shows that the toxin receptor exhibits confined motion within microdomains. To analyze the receptor trajectories, we introduced a novel approach based on an inference method [1]. Our only assumption is that the receptor moves according to the Langevin equation of motion. This method fully exploits the information of the ensemble of the trajectory (Fig. A), in contrast to the usual mean square displacement analysis, which focuses only on a single observable, the second-order moment. Applying both techniques to collected trajectories, we can highlight the difference in extracted parameters. From the shape of the confining potential (Fig. C), which is obtained by mapping the forces (Fig. B) inside domains, we can deduce information about the confinement of the membrane. In combination with experiments on cholesterol depletion and cytoskeleton depolymerization, this technique will shed light into the nature of the membrane micropatterning.


1592-Pos
Mechanisms Regulating the Diffusion of the Lipid Raft Marker Cholera Toxin B Subunit
The B subunit of cholera toxin (CTXB) is generally accepted as a marker of lipid rafts. Compared to other raft markers or lipid-anchored proteins, CTXB exhibits relatively slow diffusion. A variety of mechanisms could potentially account for this slow diffusion of CTXB, including crosslinking of small raft domains, confinement by the actin cytoskeleton, association with caveolae, incorporation into actively maintained domains, or molecular crowding effects in response to elevated membrane protein density. We evaluated the role of each of these mechanisms in controlling the lateral diffusion of CTXB in the current study by employing fluorescence recovery after photobleaching (FRAP) of fluorescently labeled CTXB following actin depolymerization, ATP depletion, cholesterol depletion, labeling across a range of CTXB concentrations, or in caveolin-1 knockout MEFs. Of these conditions, only cholesterol depletion significantly altered the diffusion mobility of CTXB. Furthermore, we tested whether the slow diffusion of CTXB is an intrinsic property of its receptor by examining the effects of CTXB on the diffusion of a fluorescent GM1 analog. The results of this experiment showed that CTXB slows the diffusion of its receptor. However, binding of CTXB to cells did not affect the diffusion of another raft marker (YFP-GL-CP), a non-raft marker (YFP-GT46), or a fluorescent lipid analog (DiC16). Taken together, these data suggest that CTXB diffusion is not limited by actin corals, caveolae, molecular crowding effects, or the intrinsically slow diffusion of GM1. In addition, they suggest that crosslinking of small rafts induced by CTXB binding does not substantially alter the dynamics of membrane domains enriched in other types of raft or non-raft proteins or lipids.

1593-Pos
Direct Observation of Hop Diffusion of Lipid and Protein Molecules in the Plasma Membrane by High-Speed Single Fluorescent-Molecule Imaging
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Spatio-Temporal Regulation of RhoGTPases Signaling by Myosin II. Selwin K. Wu1,2* and Rashmi Priya3*. (B) Model for self-organization of the biomechanical cortical network at the lateral adherens junction and the medial apical cortex of cells. Similarly, contractile stress-induced disassembly of tensile actomyosin cables can also relax the actomyosin network (Wu et al., 2014a,b; Jodoin et al., 2015). In addition, myosin II can scaffold signaling molecules to regulate RhoGTPase activity (Conti and Adelstein, 2008). Myosin II has a head domain that binds and exerts forces on F-actin, and also rod domains which self-associate to form bipolar filaments of Myosin molecules (Figure 2). Strikingly, myosin II can regulate signaling via its rod domain. Active Remodeling of Cortical Actin Regulates Spatiotemporal Organization of Cell Surface Molecules. Theory. Active Remodeling of Cortical Actin Regulates Spatiotemporal Organization of Cell Surface Molecules. We propose a model for nanoclustering based on active hydrodynamics, wherein cell surface molecules bound to dynamic actin are actively driven to form transient clusters. This consistently explains all of our experimental observations. In summary, our theoretical framework invoking dynamic actin filaments in conjunction with motor activity provides a consistent explanation of all of the observations of the distribution and dynamics of passive and inert molecules on the plasma membrane from previously published experimental data of Goswami et al. (2008). Download Citation | Active Re-Modelling of Cortical Actin Regulates Spatiotemporal Organization of Molecules on a Living Cell Surface | Cell surface proteins such as lipid-tethered GPI-anchored proteins, Ras-proteins and several glycoproteins, are distributed as monomers and...