Roles of the cytoskeleton in regulating EphA2 signals

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The lateral organizations of receptors in the cell membrane display a tremendous amount of complexity. In some cases, receptor functions can be attributed to specific spatial arrangements in the plasma membrane. We recently found that one member of the largest subfamily of receptor tyrosine kinases (RTKs), EphA2, is organized over micrometer length scales by the cell’s own cytoskeleton, and that this can regulate receptor signaling functions. Spatial organization of the receptor was found to be highly associated with invasive character, and mechanical disruption of receptor organization altered key down-stream events in the EphA2 signaling pathway. In this Addendum article, we put forth possible models for why EphA2 and other receptors may employ mechanical and spatial inputs mediated by the cytoskeleton. We speculate that this class of input may be common, and contributes to the intricacies of cellular signaling.

Receptor Spatial Organization

The spatial organization of receptors in the cell membrane spans multiple length scales, from the molecular to the size of the cell itself. Signaling assemblies consisting of tens, to tens of thousands of molecules can apparently function as cooperative units. Hierarchical organization of signaling receptors can directly feed into signaling pathways to regulate collective cell signaling outcomes.\textsuperscript{1,3} For example, T-cell receptor activation was found to be dependent on the spatial organization within the immunological synapse,\textsuperscript{1,4-9} and in a recent report we recently found that the EphA2 receptor tyrosine kinase (RTK) pathway can be modulated based on receptor translocation.\textsuperscript{3}

There are distinct biophysical mechanisms that regulate receptor spatial organization and associated biochemical functions. The most commonly studied is direct protein-protein interaction. For example, the ligand-induced dimerization of RTKs is widely considered as the prototypical mechanism for their activation.\textsuperscript{2,10-13} Another effecter that influences protein organization is lipid-membrane driven separation of proteins into discreet assemblies. The formation of such lipid membrane compartments may be based on the immiscibility of specific lipid components in the plasma membrane\textsuperscript{14-16} or mechanical bending effects at an intermembrane junction.\textsuperscript{17} A third cellular regulator of protein organization is the network of cytoskeleton filaments which can act as scaffolds with the aid of adaptor proteins for corolling or directly moving receptors across the cell membrane.\textsuperscript{1,3,7,18} The interplay between these mechanisms exerts hierarchal and dynamic control of receptor organization and cell function. The role of the cytoskeleton is typically studied in the context of adhesion proteins such as integrins, and its role in the arrangement of free floating membrane proteins is poorly defined.\textsuperscript{18,39} This is because the connectivity between free floating receptors and the cytoskeleton is not clear and little is known about these associations.

The EphA2 Signaling Pathway

RTKs play important roles in receiving and amplifying signals from other cells.
and from the immediate environment. The Eph family of receptors constitute the largest subfamily of RTKs, and these contribute to cellular development and morphogenesis in a wide range of tissues. Abnormal expression and function of the EphA2 receptor is implicated in a range of human malignancies including breast, lung and ovarian cancers. In particular, 40% of human breast cancers overexpress EphA2, which is associated with a poor prognosis and the development of drug resistance.20-22

The ligand to EphA2 is a membrane-associated GPI-linked protein expressed on the surface of adjacent cells.22,23 Because both the ligand and receptor are in membranes, EphA2 binding and activation can only proceed through direct physical contact between cells. Structural studies of EphA receptors indicate that ligand-binding can lead to dimerization and the formation of higher order aggregates.22,23 Clustering of Eph-ephrin complexes is thought to be enhanced by specific domains.24 These include the fibronectin type III repeats, the SAM domain of the Eph receptors and by PDZ domain proteins.25 Ligand-induced clustering of the EphA2 receptor results in autophosphorylation and recruitment of downstream signaling molecules through Shc and Grb2 adaptor proteins. Receptor activation leads to stimulating the PI3K, Akt and MAPK adaptor proteins. Receptor activation leads to the activation of Erk and metalloprotease 10 (ADAM10) which regulate signaling through receptor degradation.

As is the case with most studies on such juxtacrine signaling systems, activation of EphA2 is often achieved with soluble ligands that are pre-clustered. We hypothesized that ephrin-A1 bound to synthetic lipid membranes would provide for a better mimic of the natural cell-cell junction geometry and might reveal additional features of this signaling process.3 This interface presents active ephrin-A1 ligand molecules that are fluid in two dimensions and thus captures some of the native geometry. We found that membrane-bound ephrin-A1 triggers the EphA2 receptor on living cells and allows for quantifying receptor translocation. Such quantitative measurements are difficult in live cell couples, and therefore have not been explored in detail.

Seeking Signals: Cytoskeleton Transport of Ligand-Bound EphA2

Using live-cell fluorescence microscopy techniques we found that the EphA2 receptor rapidly formed clusters as a result of ligand binding. Clusters grew and coalesced until they were transported to the center of the cell-supported membrane junction. The motion of the EphA2-ephrin-A1 clusters was highly correlated to the motions of the actin cytoskeleton. This was measured using two-color total internal reflection fluorescence microscopy (TIRFM) tracking of ephrin-A1 and enhanced green fluorescent (EGFP)-actin. Eph receptors are known to play a role in remodeling the actin cytoskeleton and to elicit actomyosin contraction through the Rho family of guanosine triphosphate hydrolases (GTPases).24 Ephrin-A1 stimulation of EphA2 is reported to lead to RhoA-dependent actomyosin contractility, which is in agreement with the observed cellular phenotypes in our experiments.3,26,27 Interestingly, we found that the translocation of ligand-bound EphA2 followed that of the actomyosin contractility, thus suggesting a physical association between them. In order to identify the mechanism of actin reorganization and its connection with EphA2 transport, we used the Rho-kinase inhibitor Y27632 to block the cytoskeleton contraction pathway.27 Analysis of the results revealed that the mechanism of ligand-receptor transport was mediated through a RhoA-dependent pathway that actively transports EphA2 receptor clusters.

To elucidate the relation between EphA2 receptor motions and specific signaling cascades we used the “spatial mutation” strategy.5,3,7,8 In this approach, physical barriers fabricated onto the underlying substrate guide mobility of molecules in the supported membrane. These structures additionally impede the lateral motion of cell surface receptors through their action on bound ligands in the supported membrane. The technique is highly specific because only ligand-bound EphA2 receptors expressed on the cell surface are spatially reorganized. Interestingly, cells with spatially mutated EphA2 receptor organization showed: (a) altered f-actin morphology and (b) a decrease in the recruitment and the localization of the ADAM10 metalloprotease. Given the roles of secretases (such as ADAM10), and the cytoskeleton, the physical reorganization of the EphA2 receptor may have wide implications across multiple signaling pathways. ADAM10, for example, plays an important role in EphA receptor signaling since it takes part in ephrin ligand shedding; thus allowing for release of the physical tether between adjacent cells engaged in juxtacrine signaling. The spatial mutation strategy avoids off-target effects that are common when using pharmacological or genetic inhibition and thus it provided a direct link between receptor transport, ADAM10 recruitment and actin dynamics.

Why Mechanical Force? Biological Roles for Receptor Transport

The direct consequences of EphA2 transport are two-fold (see Fig. 1). The first is altering the size and the distribution of EphA2-ephrin-A1 clusters across the cell-cell junction. One may be tempted to draw parallels with the immunological synapse where the location of the T-cell receptor (TCR) can affect phosphorylation states.5,7,8 This is not the case, and we did not observe clear differences in EphA2 phosphorylation as a function of receptor spatial organization.5 The cellular mechanism of receptor transport may be similar, but signal outputs are different. A second consequence of EphA2 transport is the potential to apply mechanical strain on the EphA2-ephrin-A1 complex. If the ligand or the receptor encounters resistance to lateral transport, then the complex will experience tension and may undergo a conformational switch. The type of signaling mechanism (mechano-transduction model) is commonly cited for proteins involved in cellular adhesion such as the integrin family.28-31

An important question pertains to why the EphA2 pathway might incorporate sensitivity to force. The formation of complex tissues with controlled form and tissue homeostasis implies that cells can
sence and react to very subtle changes in the mechanical properties of their environments.\textsuperscript{28,32,33} In this regard, much attention has been focused on the integrins and associated focal adhesion proteins as master regulators of force.\textsuperscript{31,34} However, our work suggests that other receptors may incorporate sensitivity to force and we propose that mechanotransduction may be a common motif in signaling pathways. Mechanical aspects of the cellular microenvironment will change the spatial organization and the tension forces acting on all receptors whose ligands are surface associated. Correspondingly, it is likely that natural selection processes have explored this component of signal regulation.

In the case of EphA2, the increased ADAM10 recruitment found in the unrestricted EphA2 receptor clusters suggests that there may be enhanced levels of ligand cleavage and endocytosis of the ligand-receptor complex. The accepted mechanism for termination of EphA2 forward signaling involves the regulated cleavage of ligands by the ADAM10 protease.\textsuperscript{35,36} Enhanced rates of EphA2 endocytosis would consume available ephrin-A1 ligand and may bias the input-response function of the entire system. Importantly, the EphA2 receptors interact with other signaling pathways, such as the chemokine receptors, integrins and cadherins, which suggests that increased EphA2 receptor endocytosis may affect other signaling cascades.

Irrespective of the actual biological purpose for these behaviors, the experimental tools developed—the spatial mutation strategy—offer a route to uncovering mechanisms and signaling roles for receptor spatial organization. The technique enables single cell manipulations and facilitates quantitative characterization.\textsuperscript{8} Importantly, this approach differs fundamentally from conventional tools employed for deconstructing the signaling roles of the cytoskeleton. For example, the most widely used approach consists of drug targeting of the cytoskeleton, which affects many signaling pathways and thus lacks specificity.

An open question to be addressed is the mechanism mediating the physical association of EphA2 to the cytoskeleton. The ERM family of intracellular proteins (which includes ezrin, radixin and moesin) are possible candidates since they are known to mediate dynamic binding between actin filaments and the cytoplasmic face of several transmembrane proteins.\textsuperscript{38} ERM proteins are known to display diversity in their functions across different cell lines. For example, ezrin and moesin play an active role in the human T cell activation pathway by influencing the spatial organization of the immunological synapse.\textsuperscript{1,7,17,39} A multidisciplinary approach that combines advances in biophysical chemistry, optical microscopy/nanoscopy and cell biology will be required to identify and characterize the proteins mediating this coupling.

Given the mechanical sensitivity of the EphA2 pathway, it seems plausible that many other receptor pathways are susceptible to mechano-regulation. We speculate that receptor transport is a general mechanism used by a range of cells and receptors and, in different contexts, may be used to achieve specific goals. The advent of physical methods, such as the spatial mutation strategy, marks a clear path toward investigating the subtleties of mechanical/spatial transduction, and one that involves the confluence of biophysics, surface chemistry and cell biology.

Figure 1. Scheme depicting the mechanical coupling of ligand bound EphA2 clusters and the actin cytoskeleton. This physical coupling may alter the EphA2 pathway by: (i) changing the size and distribution of clusters, and (ii) imposing mechanical tension on the EphA2-ephrin-A1 complex. See text for details.

References