Coordinating cytoskeletal tracks to polarize cellular movements

Atsuko Kodama, Terry Lechler, and Elaine Fuchs

For many years after the discovery of actin filaments and microtubules, it was widely assumed that their polymerization, organization, and functions were largely distinct. However, in recent years it has become increasingly apparent that coordinated interactions between microtubules and filamentous actin are involved in many polarized processes, including cell shape, mitotic spindle orientation, motility, growth cone guidance, and wound healing. In the past few years, significant strides have been made in unraveling the intricacies that govern these intertwined cytoskeletal rearrangements.

In this report, we highlight the actin–microtubule crosstalk that occurs during directed cell movements. For each step in the process, we review the regulatory mechanisms that underlie both the independent and the interdependent cytoskeletal rearrangements of actin filaments and microtubules that coordinate cellular locomotion in a polarized fashion. We discuss how external directional cues activate Rho GTPases at specific cellular sites to initiate localized cytoskeletal rearrangements, and we focus on the signaling pathways that cause either actin or microtubule rearrangement and the regulatory interactions between these cytoskeletons. We also consider “search and capture” mechanisms involving structural interactions between F-actin and microtubules near the leading edge of cells. Finally, we spotlight spectraplakins, able to directly bind both F-actin and microtubules. Recently, spectraplakins have emerged as candidates for coordinating the two cytoskeletons in directional migration.

Establishment of the cortical platform at the leading edge of a cell

Cells migrate by coordinating cytoskeletal-mediated extensions and contractions concomitantly with making and breaking contacts to an underlying substratum. To orchestrate directional movements, cells must activate a specific site(s) at the membrane periphery in response to a polarized external cue (Fig. 1; top left). A particular locale then becomes a “cortical platform” for the transmission of converging internal signals that are necessary to elicit subsequent cytoskeletal responses. The outcome is dependent upon the cell type and the precise signaling pathways that are engaged, and can range from the polymerization and/or reorganization of actin to the polarized capture and stabilization of microtubules and their associated microtubule organizing center (MTOC).

A cortical platform can facilitate crosstalk between F-actin and microtubules by functioning as a transducer/amplifier of the internal cellular signals that orchestrate both cytoskeletons. Small GTPases such as Cdc42, Rac, and Rho have long been implicated in these processes, but precisely how their activities are temporally and spatially regulated at cortical platforms has often been obscure. Some insights have come from studying cultured mammalian cells, including epithelial cells, neurons, astrocytes, and fibroblasts, all of which use transmembrane integrin heterodimers to adhere to, organize, and migrate on a substratum of ECM (Hood and Cheresh, 2002; Fukata et al., 2003).

Referred to as “directional sensors” or “compasses,” the internal cellular modules able to sense extracellular directional gradients have been particularly well studied in chemotactic neutrophils. The engagement of G protein–coupled receptors and activation of Gβγ at the neutrophil surface triggers a complex signaling cascade that culminates in cytoskeletal reorganization and directed migration (Meili and Firtel, 2003). Recent reports reveal that Gβγ binds p21-activated kinase 1, which recruits and activates a guanine nucleotide exchange factor referred to as PIXα. Once activated, PIXα then associates with the small GTPase Cdc42, which upon activation can stimulate actin polymerization (Li et al., 2003).

Positive reinforcement of the process appears to occur through the added ability of Gi to recruit and activate phosphatidylinositol 3 kinase, which promotes the accumulation of phosphatidylinositol (3,4,5)P3 (PIP3). PIP3 then serves as a docking site for PH domain harboring proteins such as guanine nucleotide exchange factors for Rho family GTPases, including not only Cdc42 but also Rac. In this way, the positive-feed back loop involving PIP3 results in increased levels of activated Rac/Cdc42 at the leading edge of the cell (Meili and Firtel, 2003). In the meantime, Rho appears to be activated at the rear of the neutrophils via G12 and G13 (Xu et al., 2003). A recent but likely not final twist to these complexities in directional

Correspondence to Elaine Fuchs: fuchslb@rockefeller.edu

Abbreviations used in this paper: +TIP, plus end–interacting protein; MTOC, microtubule organizing center.
sensing mechanisms comes from analyses on T cells, dendritic cells, and fibroblasts that implicate glycosylphosphatidylinositol-anchored proteins in lipid rafts in the activation of small GTPases (del Pozo et al., 2004; Jakstis et al., 2004; Krautkramer et al., 2004; Palazzo et al., 2004).

Localized activation of small GTPases appears to be a unifying and early step in orchestrating the downstream rearrangements in cytoskeleton necessary to polarize cell motility (Fig. 1; top right). Fluorescence resonance energy transfer experiments using probes tailored for individual family members have provided suggestive evidence that different Rho GTPases might function in transmitting unique signals from different cortical platforms, and such specificity may be operative at least in some situations (Kraynov et al., 2000; Gardiner et al., 2002; Itoh et al., 2002).

**Actin and microtubule regulatory signals transmitted from the cortical platform**

After polarized activation of Rho family small GTPases at cortical platforms, cells transmit downstream signals that are responsible for two distinct processes—motility and polarity. Cells respond to activated GTPases by mobilizing their actin cytoskeletal network and changing their morphology (Hall, 1998). This polarized rearrangement of actin-based structures provides the driving force for “motility,” resulting in the GTPase-dependent induction of filopodia, lamellipodia, and stress fibers (Hall, 1998). However, without polarization of the microtubule cytoskeleton as well, cells cannot sustain the directionality of their movements.

Recent reports reveal that activated Rho promotes the stabilization of microtubules through its downstream target effector mDia (Wen et al., 2004). This stabilization has been visualized by immunofluorescence through the aid of an antibody that binds to the exposed COOH-terminal glutamine residue in long-lived tubulin (Palazzo et al., 2001). Microtubule dynamics can also affect the activity of Rho GTPases and the ability of cells to migrate (Waterman-Storer et al., 1999; Rodriguez et al., 2003). In particular, microtubule disassembly results in Rho activation, yielding an increase in focal adhesions (Krendel et al., 2002; Kirchner et al., 2003), and conversely, microtubule targeting to focal adhesions appears to promote focal contact disassembly (Kaverina et al., 1999). The ability of Rho GTPases to impact on both actin and microtubule cytoskeletons suggests an underlying interdependency upon what had long been surmised to be separate cytoskeletal networks. However, in this case the microtubule–actin crosstalk arises not from structural interactions, per se, but rather from alterations in the regulatory signals that modulate these two cytoskeletons (Wehrle-Haller and Imhof, 2003).

**Capturing microtubules at cortical platforms**

As postulated by Kirschner and Mitchison nearly two decades ago, microtubules “search” cytoplasmic space by continuously growing and shrinking from their plus ends, which project outward to the cell periphery (Kirschner and Mitchison, 1986). Microtubules are then “captured” and transiently stabilized at specific membrane target sites through plus end–interacting proteins (+TIPs), such as EB-1 and CLIP-170 (Schuyler and Pellman, 2001; Gundersen et al., 2004). +TIPs are thought to act in part by protecting the growing ends of microtubules from catastrophe proteins that might otherwise bind to and initiate the depolymerization of the microtubule (Komarova et al., 2002; Tirnauer et al., 2002). Interestingly, activated Cdc42 and Rac may impact on the growth and dynamics of microtubules at the cell periphery by a PAK signaling pathway that most likely inhibits the microtubule-destabilizing protein Op18/stathmin (Daub et al., 2001; Wittmann et al., 2003, 2004).
+TIPs not only participate in the stabilization of microtubules, but also in targeting microtubules to specific locales. For example, RNA interference knockdown analyses in *Drosophila* and mammalian cells have unveiled functions for EB-1 not only in microtubule dynamics, but also chromosome segregation (Rogers et al., 2002; see also Louie et al., 2004). +TIPs can also interact with members of protein complexes at cortical platforms. One such protein is APC, which through independent binding domains has the capacity to bind to both EB-1 and cortical proteins (Barth et al., 2002).

A particularly powerful system for dissecting the sequence of molecular events involved in cellular polarization is the introduction of scratches or wounds into a monolayer of adherent mammalian cells in culture. In wounded astrocyte cultures, for instance, the small GTPase Cdc42 is activated at the leading edge, a process that triggers the binding of a polarizing protein Par6, which in turn activates PKCζ, which then phosphorylates and inactivates GSK3β (Etienne-Manneville and Hall, 2001, 2003). This cascade has been proposed to enable APC to then associate with microtubule tips and allow the selective capturing and stabilization of microtubules at the leading edge of the migrating front.

Some individual cells move in a random but polarized fashion, which at first glance appears to be analogous to the polarization process described for a wound response. However, in one report the downstream effector was IQGAP, which is also a direct binding partner for activated Cdc42 at the leading edge (Fukata et al., 2002). Additional direct interactions between IQGAP and the +TIP CLIP-170 then appeared to link the temporal capture of the microtubule plus ends to this activated cortical platform. Interestingly, expression of a mutant IQGAP that could not bind activated GTPases resulted in multiple protrusion sites (Fukata et al., 2002). This provides further evidence that coupling microtubule stabilization to a cortical platform is required for sustaining polarity.

Additional direct GTPase targets such as mDia have also surfaced as binding partners or regulators for the +TIP EB-1 (Wen et al., 2004). Other proteins that localize to and are likely to be involved in these types of F-actin–microtubule connections include the minus end–directed microtubule motor protein dynein, the CLIP-associated proteins (or CLASPs), and the gigantic spectraplakin protein ACF7 (Leung et al., 1999; Karakesisoglou et al., 2000; Kodama et al., 2003; Gundersen et al., 2004).

Whether in isolation or as an adhering sheet, cells also often polarize their MTOC in the direction of migration. In wounded astrocyte cultures, dominant-negative disruption of dynein function abrogates the MTOC reorientation process (Etienne-Manneville and Hall, 2001), and this and other reports implicate dynein/dynactin in the signaling pathway that leads to MTOC positioning (Burakov et al., 2003). Although a direct connection between dynein/dynactin and the Cdc42/Par6/PKCζ/GSK3β pathway has not yet surfaced, increasing evidence points to the view that the cortical platform that develops at a wound edge can act as a scaffolding complex. This concept sets the scene for multiple +TIPs to encounter many different receptor proteins that may converge at this platform.

To illustrate the myriad of potential interactions afforded by such a scaffold, APC can bind to the adherens junction protein β-catenin (Dikovskaya et al., 2001). β-Catenin in turn can bind to members of the dynein/dynactin complex (Ligon et al., 2001), although so can EB-1 (Berrueta et al., 1999), and EB-1 in turn can bind to CLIP170 and Lis1 (Coquelle et al., 2002; Goodson et al., 2003). As if one of these various circuitous routes weren’t sufficient to recruit dynein/dynactin to polarized sites, yeast two-hybrid analyses have uncovered a direct association between the p150glued dynactin subunit and the ezrin/moesin/radixin domain of a neuronal spectraplakin that also possesses binding sites for EB-1, as well as F-actin and microtubules (Liu et al., 2003; Subramanian et al., 2003).

In summary, although a common core pathway seems likely for polarizing cell movements, the mind-boggling opportunities for direct and indirect interactions between +TIPs and cortical platform proteins seem to reflect a tailoring of this process to suit the particular needs of different cells and tissues. A comparison of this process across the eukaryotic kingdom supports the notion that a general mechanism underlies the integration of polarization processes with microtubule search and capture dynamics. The molecular twists that appear to be superimposed upon this theme seem likely to exist for the purpose of coordinating these dynamics in a regulated fashion. Additional factors may be structural, using multiple protein complexes to modify or reinforce the strength of microtubule–actin interactions. Finally, the length of time during which a localized GTPase is activated might also influence the degree to which a cortical platform amplifies microtubule retention at a polarized site (Fig. 1; bottom).

One final issue worth considering is the impact of the evolutionary spectrum on the mechanisms underlying microtubule plus end capturing by cortical platforms. In this regard, the significantly larger size of mammalian cells compared with single-cell eukaryotes such as yeasts necessitates the production of longer and more stable microtubules to span the cytoplasm. In addition, when yeast cells polarize to divide, the orientation and establishment of actin–microtubule connections is exquisitely linked to the regulation of yeast’s rapid cell cycles. As such, the tethering of microtubules to the cortical platform is both transient and cyclic. Similarly, the mating process in yeast needs only to sustain the polarization machinery for several hours. By contrast, higher eukaryotes must often maintain their tethering machinery for extended periods in order to accommodate more protracted polarization processes such as epithelial wound closure.

**Spectraplakins: scaffolds for direct cross-linking of actin filaments and microtubules**

The need to prolong microtubule–actin anchorage provides a potential explanation for why mammals have developed more efficient machineries to strengthen interactions between microtubule- and actin-based structures. In this regard, the spectraplakins have emerged as higher eukaryotic scaffolding proteins, which have direct binding sites for +TIPs, F-actin, and microtubules (Leung et al., 1999; Yang et al., 1999; Karakesisoglou et al., 2000; Goodson et al., 2003). As if one of these various circuitous routes weren’t sufficient to recruit dynein/dynactin to polarized sites, yeast two-hybrid analyses have uncovered a direct association between the p150glued dynactin subunit and the ezrin/moesin/radixin domain of a neuronal spectraplakin that also possesses binding sites for EB-1, as well as F-actin and microtubules (Liu et al., 2003; Subramanian et al., 2003).

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they could act early in polarization by directly tethering the process is switched off. Whether the leading edge of a wound environment. Embedded within this issue are how polarized cortical mechanisms are tailored to enable cells to perform this intricate process in response to specialized cues from their localized environment. Recent reports have contributed greatly to our understanding of how directed cell migration is orchestrated through cytoskeletal scaffolds. A special tribute goes to the Fuchs lab members, past and present, who contributed to the scientific foundation that inspired us to write this review. We are also grateful to our colleagues throughout the scientific community who have developed this field to the exciting state that it has become. Finally, we would like to thank Dr. Markus Schober for his critical reading of this review. E. Fuchs is an Investigator of the Howard Hughes Medical Institute. T. Lechler is a postdoctoral fellow supported by the Jane Coffin Childs Foundation. Past work contributing to this review has been supported by the Howard Hughes Medical Institute and by a grant from the National Institutes of Health (R01 AR27883).

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Conclusions and prospects
Recent reports have contributed greatly to our understanding of how directed cell migration is orchestrated through cytoskeletal rearrangements triggered by polarized activation of small GTPases. A challenge for the future is to understand how these mechanisms are tailored to enable cells to perform this intricate process in response to specialized cues from their localized environment. Embodied within this issue are how polarized cortical platforms are sustained for different lengths of times and how the process is switched off. Whether the leading edge of a wound site or a cell–cell or cell–substratum junction, a polarized cortical platform presents a molecular galaxy in which to integrate a constellation of signal transduction pathways with cytoskeletal rearrangements. The science underlying this field is likely to hold many new insights into cell biology and is likely to keep researchers concentrated in this area for many years to come.

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