Molecular pathways regulating mitotic spindle orientation in animal cells

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Summary
Orientation of the cell division axis is essential for the correct development and maintenance of tissue morphology, both for symmetric cell divisions and for the asymmetric distribution of fate determinants during, for example, stem cell divisions. Oriented cell division depends on the positioning of the mitotic spindle relative to an axis of polarity. Recent studies have illuminated an expanding list of spindle orientation regulators, and a molecular model for how cells couple cortical polarity with spindle positioning has begun to emerge. Here, we review both the well-established spindle orientation pathways and recently identified regulators, focusing on how communication between the cell cortex and the spindle is achieved, to provide a contemporary view of how positioning of the mitotic spindle occurs.

Key words: Spindle, Microtubules, Oriented cell division, Polarity, Mitosis, Centrosome

Introduction
All multicellular animals are tasked with two fundamental developmental challenges: generating cellular diversity and forming three-dimensional tissues, both of which initiate from a single-celled zygote. Cellular diversity is spawned by cell divisions yielding non-identical daughters, and tissue morphogenesis is established through the precise three-dimensional arrangement of cell divisions that form the overall architecture of the organism. Both of these essential challenges are resolved in part through oriented cell division, which regulates embryogenesis, organogenesis and cellular differentiation. Notably, oriented cell divisions, and hence asymmetric cell divisions, remain crucial throughout adulthood as well, functioning as the basis for tissue homeostasis during growth and wound repair. One primary feature of oriented cell division is the proper positioning of the mitotic spindle relative to a defined polarity axis. In principle, spindle orientation is achieved through signaling pathways that provide a molecular link between the cell cortex and spindle microtubules. These pathways are thought to elicit both static connections and dynamic forces on the spindle to achieve the desired orientation prior to cell division. Although our knowledge of the signaling molecules involved in this process and our understanding of how they each function at the molecular level remain limited, collective efforts over the years have shed light on the importance of spindle orientation to animal development and function. Moreover, emerging evidence shows an association between improper spindle orientation and a number of developmental diseases as well as tumor formation. The study of spindle orientation is therefore fundamental to both developmental biology and human disease.

Over a century ago, Oscar Hertwig discovered that sea urchin embryos biased the orientation of the mitotic spindle along their long axis, which led to a model (the ‘Hertwig Rule’) in which cells orient divisions in response to mechanical forces (Hertwig, 1884). The Hertwig model stated that mechanical regulation was the primary determinant of spindle orientation as cells sensed shape changes in response to external forces. Later discoveries, including those in ascidian embryos, in which cell division orientation correlated with differential daughter cell fate and size, suggested a molecular basis for spindle orientation (Conklin, 1905). Conklin reasoned that the ability to alter division orientation to achieve autonomous fate specification during development must rely upon internal molecular regulation rather than external mechanics. The modern molecular era has now vastly expanded our understanding of oriented cell division, with studies showing support for both models.

Genetic identification of the first spindle orientation regulators occurred nearly two decades ago (Cheng et al., 1995; Etemad-Moghadam et al., 1995; Kraut et al., 1996; Zwaal et al., 1996). Since then, the ‘parts list’ of proteins required for proper spindle orientation has grown tremendously, and includes components from several signaling pathways that couple the mitotic spindle to cortical polarity complexes in a variety of cell types from a diverse group of organisms. More recent studies utilizing genetic screening technologies and improved cell culture-based systems have expanded that list further. Moreover, these recent studies have begun to view spindle orientation through a more molecularly focused lens that allows better insight into how these parts fit together during oriented cell divisions.

The ability to bias the orientation of cell division via regulated spindle positioning is conserved from yeast to mammals (Siller and Doe, 2009). For the scope of this Review, we have chosen to focus on selective, well-studied examples from metazoan model organisms. We first highlight several developmental processes that rely on regulated spindle orientation/asymmetric cell divisions and provide an overview of how cell polarity, and hence spindle orientation, is first established in these examples. A particular focus will be placed on the mechanisms by which cortical polarity cues ‘capture’ spindle microtubules, as this process represents an early step in the spindle orientation process. Although microtubule capture itself is likely to be a conserved and generalized aspect of spindle orientation, the underlying molecular pathways appear to be diverse, highlighting the importance of further understanding of the mechanisms involved. We describe several well-established spindle orientation components as well as newly identified factors that regulate communication between the cell cortex and the mitotic spindle, with an aim to provide readers with an updated view of how different cell types regulate the position of the mitotic spindle.

Oriented/asymmetric cell divisions during metazoan development
It should be noted that oriented/asymmetric cell division, as discussed throughout this Review, is defined by two mutually
coupled events: (1) establishment of a cortical polarity axis by the unequal distribution of polarity proteins and cell fate determinants; and (2) alignment of the mitotic spindle with respect to this polarity axis. In certain cases, the polarity axis coincides with a tissue or body axis of the organism. We will focus our discussion primarily on how these two processes are linked at the molecular level using four examples: the Caenorhabditis elegans zygote, Drosophila neuroblasts, Drosophila sensory organ precursors and mammalian epidermal cells. As gene and protein names often vary between species, please see Table 1 for the naming of orthologs.

The first series of divisions in the C. elegans zygote produces siblings that are asymmetric in both size and fate (Fig. 1A). Early genetic studies revealed a set of evolutionarily conserved partitioning-defective (PAR) proteins that are necessary for establishing an anterior-posterior (A-P) cortical polarity axis prior to the first zygotic division (Kemphues et al., 1988). Subsequent studies demonstrated that the PAR complex also regulates spindle orientation and the unequal force generation exerted on the spindle poles (Etemad-Moghadam et al., 1995; Grill et al., 2001; Kemphues et al., 1988). Defects in PAR complex genes result in mispositioning of the mitotic spindle, loss of daughter cell asymmetry and, ultimately, non-viable animals. As shown in Fig. 1A, two successive divisions with A-P spindle orientations produce a four-cell embryo containing a blastomere known as the EMS blastomere. Spindle orientation along the A-P axis and subsequent asymmetric cell division of the EMS cell then generates an E daughter cell, which will give rise to endodermal lineages, and a MS daughter cell, which will form mesodermal lineages. Spindle orientation in the one-cell stage and the P2 blastomere cell requires the activity of GPR-1/2 and LIN-5, which constitute an evolutionarily conserved non-canonical G-protein signaling network (Colombo et al., 2003; Gönczy, 2008; Werts et al., 2011). Mutations in Wt signaling pathway components result in misalignment of the EMS spindle and mis-specification of germ cell layers (Schlesinger et al., 1999; Walston et al., 2004) (Fig. 1A). By manipulating contact sites at the four-cell stage, Goldstein showed that cell-cell contacts establish a site that captures centrosomes via emanating microtubules to orient cell divisions (Goldstein, 1995). Actin-rich contact sites between the EMS and P2 cells determined spindle orientation and influenced partitioning of fate information necessary for gut specification (Goldstein, 1995; Waddle et al., 1994). Collectively, these studies suggest that spindle orientation is an essential determinant of cell fate specification during C. elegans development.

Asymmetric division of Drosophila neuroblasts, the stem cells of the developing fly brain, regulates development of the fly central nervous system (Fig. 1B). Neuroblasts polarize along an apical-basal (A-B) axis and divide in a stem cell-like manner to produce a self-renewed neuroblast and a ganglion mother cell (GMC) that produces differentiated neurons and glia (Doe, 2008). Thus, a relatively small number of neuroblasts can supply the vast number of self-renewed neuroblasts that participate in astral microtubule formation possibly through a dynein-dependent transport complex (Delaval et al., 2011; Thoma et al., 2009) and LKB1, regulate spindle orientation (Table 1), suggesting that spindle misorientation might contribute to tumor development (Pease and Tirnauer, 2011).

Neurogenesis requires control of cell division orientation in neuroprogenitors at specific developmental stages (Lancaster and Knoblich, 2012) to balance proliferative and neurogenic outcomes (Konno et al., 2008; Peyre et al., 2011). Mutations in the spindle orientation regulators LIS1 (PAFAH1B1) and HTT manifest in Type I lissencephaly and Huntington’s disease, respectively. Interestingly, both LIS1 and HTT regulate the function of cytoplasmic dynein, a potentially universal regulator of spindle orientation (Table 1; see main text). Whether a causative molecular link exists between spindle orientation and neurodevelopmental disorders will be an important future research endeavor.

Components of the intraflagellar transport (IFT) machinery in cilia have been linked to misoriented cell divisions possibly underlying polycystic kidney disease (PKD) (Delaval et al., 2011; Fischer et al., 2006; Hildebrandt and Otto, 2005). Several IFT proteins localize to centrosomes, and mutations in IFT88 result in spindle misorientation (Delaval et al., 2011), leading to improper spatial arrangement of nephron epithelia and cyst development. Interestingly, IFT88 participates in astral microtubule formation possibly through a dynein-dependent transport complex (Delaval et al., 2011). However, mutations in other IFT components appear to drive cystogenesis through mechanisms independent of spindle misorientation (Jonasson et al., 2012).

Intestinal crypt stem cells regulate spindle orientation to produce differentiated progeny. Heterozygous mutations in the tumor suppressor protein APC, which is mutated in ~85% of colorectal cancer cases (Markowitz and Bertagnolli, 2009), results in spindle misorientation and altered cell shape (Quyn et al., 2010). However, this dependence of intestinal stem cells on division orientation/asymmetric division has been challenged recently and remains controversial (Schepers et al., 2011).
Table 1. Identified regulators of spindle orientation

<table>
<thead>
<tr>
<th>Protein name</th>
<th>Proposed localization and function</th>
<th>References</th>
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<tr>
<td><strong>Dm/Hs: InsCuteable</strong></td>
<td>Cortically polarized. Recruits and stabilizes Pins at the cortex.</td>
<td>Du et al., 2001; Kraut et al., 1996; Schaefer et al., 2000; Schober et al., 1999; Woodarz et al., 1999</td>
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<tr>
<td><strong>Dm: Pins (Raps)</strong></td>
<td>Cortically polarized. Recruits Mud-Dynein and Dlg-Khc-73 complexes to connect to spindle MTs.</td>
<td>Colombo et al., 2003; Du et al., 2001; Johnston et al., 2009; Lechler and Fuchs, 2005; Schaefer et al., 2000</td>
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<td><strong>Ce: GPR-1/2</strong></td>
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<td><strong>Hs: LGN (GPM2)</strong></td>
<td>Cortically polarized, centrosomes, spindle MTs. Binds Pins TPR domains; associates with Dynein-Dynactin complex; participates in cortical force generation on the spindle.</td>
<td>Bowman et al., 2006; Colombo et al., 2003; Du and Macara, 2004; Gönçzy et al., 1999; Izumi et al., 2006; Johnston et al., 2009; Siller et al., 2006; Siller and Doe, 2008</td>
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<td><strong>Dm: Mud</strong></td>
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<td><strong>Ce: LIN-5</strong></td>
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<tr>
<td><strong>Hs: NuMA (NUMA1)</strong></td>
<td>Spindle MTs. Minus-end-directed motor protein; transduction of cortical pulling forces.</td>
<td>Gönçzy et al., 1999; Kotak et al., 2012; Johnston et al., 2009; Siller and Doe, 2008</td>
</tr>
<tr>
<td><strong>Dm/Ce: cytoplasmic dynein</strong></td>
<td>CAP-Gly domain, MT plus-end binding. Promotes MT rescue from catastrophe, resulting in MT growth towards cortex; yeast homolog may activate dynein.</td>
<td>Akhmanova et al., 2001; Komarova et al., 2002; Miller et al., 2006; Sheerman et al., 2003</td>
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<td><strong>Dm: CLIP-190</strong></td>
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<td><strong>Hs: CLIP-170 (CLIP1)</strong></td>
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<td><strong>Dm: Glued</strong></td>
<td>CAP-Gly domain, MT plus-end binding. Required for centrosome orientation in C. elegans; Glued regulates spindle assembly; enhances Dynein processivity.</td>
<td>Culver-Hanlon et al., 2006; Gönçzy et al., 1999; Siller et al., 2005</td>
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<td><strong>Ce: DNC-1</strong></td>
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<td><strong>Hs: p150-Dynactin (DCTN1)</strong></td>
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<td><strong>Dm: Ctp</strong></td>
<td>Spindle MTs and centrosome. Component of the Dynein complex; binds the centrosomal protein Ana2; promotes cortical Pins-Mud complex.</td>
<td>Wang et al., 2011</td>
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<td><strong>Ce/Hs: DLC</strong></td>
<td>Apical cortex enrichment in neuroblasts. Binds phosphorylated Pins; associates with Khc-73; provides ‘MT capture’ pathway at cortex.</td>
<td>Johnston et al., 2009; Johnston et al., 2012; Siegrist and Doe, 2005</td>
</tr>
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<td><strong>Dm: Dlg (Dlg1)</strong></td>
<td>MT plus-ends. Plus-end tracking MT motor; associates with Pins-Dlg complex; provides static link from cortex to spindle MT plus-ends.</td>
<td>Hanada (2000); Johnston et al., 2009; Johnston et al., 2012; Siegrist and Doe, 2005</td>
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<td><strong>Dm: Khc-73</strong></td>
<td>Cortically polarized. Functions downstream of Wnt/Fz morphogenic signaling; binds Mud-Dynein complex similar to Pins; regulates activation of APC; may involve additional non-canonical pathways (e.g. activation of Rho superfamily GTPase).</td>
<td>Schlesinger et al., 1999; Ségalen et al., 2010; Walston et al., 2004</td>
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<tr>
<td><strong>Dm: Dsh</strong></td>
<td>Cortically polarized. Mud-Dynein complex similar to Pins; regulates activation of APC; may involve additional non-canonical pathways.</td>
<td>Johnston et al., 2009; Siller and Doe, 2008</td>
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<td><strong>Ce: DSH</strong></td>
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<td><strong>Dr: Dvl</strong></td>
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<td><strong>Dm: LIS-1</strong></td>
<td>Probably spindle-associated. Associates with and regulates Dynein function; participates in MT stabilization; stimulates Dynein-dependent spindle movements.</td>
<td>Godin et al., 2010</td>
</tr>
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<td><strong>Dm: Htt</strong></td>
<td>Spindle poles. Regulates localization of Dynactin, Dynactin and NuMA.</td>
<td>Thoma et al., 2009; Thoma et al., 2010</td>
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<td><strong>Hs: HTT</strong></td>
<td>Spindle microtubules. Loss of VHL leads to unstable astral MTs and loss of spindle checkpoint; inhibits MT catastrophe and promotes rescue frequency.</td>
<td>Rogers et al., 2002; Toyoshima and Nishida, 2007; Wen et al., 2004</td>
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<tr>
<td><strong>Dm/Hs: VHL</strong></td>
<td>Spindle MTs. Stabilizes spindle MTs; functions upstream of Rac in C. elegans gonad development; orients the mother centrosome in mGSCs; controls planar orientation of intestinal epithelial cells.</td>
<td>Cabello et al., 2010; Fleming et al., 2009; Yuan et al., 2007; Wen et al., 2004</td>
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<tr>
<td><strong>Dm: EB1</strong></td>
<td>MT plus-ends. Regulates MT assembly, stability and dynamics; essential for plus-end trafficking of many other components.</td>
<td>Buttrick et al., 2008</td>
</tr>
<tr>
<td><strong>Dm/Ce: APC</strong></td>
<td>MT plus-ends. Stabilizes spindle MTs; functions upstream of Rac in C. elegans gonad development; orients the mother centrosome in mGSCs; controls planar orientation of intestinal epithelial cells.</td>
<td>Buttrick et al., 2008</td>
</tr>
<tr>
<td><strong>Dm: Akt (Akt1)</strong></td>
<td>Cell cortex. Regulates centrosome migration; promotes proper spindle morphology; necessary for cortical APC localization.</td>
<td>Tabler et al., 2010; Wu and Rose, 2007; Yuan et al., 2012</td>
</tr>
<tr>
<td><strong>Dm: Par-1</strong></td>
<td>Putative EB1 interaction (MT plus-ends); cell cortex in C. elegans embryos. Kinase activity helps establish polarity of other components; ensures centrosome orientation in Drosophila mGSCs.</td>
<td>Grill et al., 2001; Tanenbaum et al., 2011</td>
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<td><strong>Ce: PAR-1</strong></td>
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<td><strong>Hs: MARK1</strong></td>
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<td><strong>Dm: MCAK (Kinesin-13; KIF2C)</strong></td>
<td>Putative EB1 interaction (MT plus-ends); MT depolymerase, promotes catastrophe; activity is coupled to spindle force generation.</td>
<td>Cabello et al., 2010; Fleming et al., 2009; Walston et al., 2004; Wen et al., 2004; Yamashita et al., 2003</td>
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<td><strong>Xl/Hs: PAK</strong></td>
<td>Cytoplasmic. Phosphorylates Ran to promote activation at centrosome; Ran promotes Mud localization.</td>
<td>Bompad et al., 2010; Speicher et al., 2008; Wei et al., 2011</td>
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<tr>
<td><strong>Dm: Canoe</strong></td>
<td>Cortical polarized in Drosophila neuroblasts; component of epithelial junctions.</td>
<td>Speicher et al., 2008; Wei et al., 2011</td>
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<td><strong>Hs: Arafadin MLLT4</strong></td>
<td>Binds directly to Pins; recruits activated Ran and promotes cortical Mud localization.</td>
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<td><strong>Dm: Aurora A (aurora kinase A)</strong></td>
<td>Spindle poles. Kinase activated in mitosis; phosphorylates Pins to promote Dlg-Khc-73 pathway activation; also promotes proper cortical polarity.</td>
<td>Johnsen et al., 2009; Lea et al., 2006a</td>
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<tr>
<td><strong>Dm: BNF1A</strong></td>
<td>Spindle poles. Kinase that promotes astral MT organization; Myosin light chain is putative effector; requires phosphorylation by LKB1 to become activated.</td>
<td>Thaiarambila, 2012</td>
</tr>
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<td><strong>Hs: AMPK (PRKAA1)</strong></td>
<td>Spindle poles. Important for astral MT organization and spindle pole development; may act downstream of Wnt signaling; target of phosphorylation currently unknown.</td>
<td>Bei et al., 2002; Nakayama et al., 2012</td>
</tr>
<tr>
<td><strong>Ce: CSK-1</strong></td>
<td>Cytoplasmic. Phosphorylates LGN to promote its polarity; phosphorylates NuMA to promote maintenance of LGN-NuMA complex.</td>
<td>Matsumura et al., 2012</td>
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<tr>
<td><strong>Dm: Dscl41</strong></td>
<td>Cytoplasmic. Phosphorylates LGN, promotes 14-3-3 binding and cortical removal; regulates LGN polarity in epithelial cells.</td>
<td>Hao et al., 2010</td>
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<td><strong>Hs: c-Src (CSK)</strong></td>
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<td><strong>Hs: ABL1</strong></td>
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<td><strong>Dm: PKC-3</strong></td>
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<td><strong>Ce: PkC</strong></td>
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<td><strong>Hs: PRKZ</strong></td>
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cell polarity (PCP) within the imaginal disc tissue. The spindle must be controlled in both an A-B orientation, through the action of the Pins complex, as well as an A-P orientation, which is controlled by a non-canonical Frizzled/Dishevelled (Fz/Dsh) signaling pathway (David et al., 2005; Gho and Schweisguth, 1998; Roegiers et al., 2001) (Fig. 1C). Proper asymmetric division in SOPs further relies on the activity of core PCP components [a discussion of which is beyond the scope of this Review, but see Wallingford (Wallingford, 2012) for an excellent recent review]. The disruption of spindle orientation during asymmetric SOP divisions results in mis-specification of cell lineages and defects in bristle development (Bellaïche et al., 2001; Lu et al., 1999). Thus, regulated spindle orientation plays a central role in the development of multicellular sensory structures.

Basal epidermal cells contribute to the architecture of the epidermis, a stratified squamous epithelium of skin that regulates fluid and electrolyte exchange and guards against harmful or
infectious substances, such as microbes. Early in development, these progenitor basal cells preferentially divide within the epithelial plane, producing two symmetric daughter basal cells to expand the undifferentiated basal layer (Fig. 1D). However, at a later developmental stage, basal cell divisions switch to an asymmetric mode, which allows the self-renewal of a proliferative basal cell while also producing a suprabasal cell that is committed to differentiating into the deep tissue layers forming the skin barrier (Lechler and Fuchs, 2005). Strikingly, this asymmetric division switch accompanies a 90° reorientation of the mitotic spindle, from parallel to perpendicular to the epithelial plane, which is dependent on the Par-aPKC and Pins-Insc-Mud polarity complexes (Fig. 1D) (Lechler and Fuchs, 2005; Poulson and Lechler, 2010). Spindle reorientation ensures asymmetric inheritance of Notch, which specifies barrier cell differentiation. Spindle misorientation mutants thus elicit impaired stratification and defects in barrier function, leading to dehydration and death (Williams et al., 2011). These results demonstrate the evolutionary conservation of coupling spindle orientation to decisions of cell fate and tissue structure, and underscore the importance of proper cell division orientation during development and in disease (see Box 1).

**Linking the spindle to the cell cortex: the role of microtubule plus-end binding proteins**

As highlighted above, correct orientation of the spindle with respect to cell polarity is crucial for tissue development and homeostasis. The ability to position statically the mitotic spindle, a large, otherwise dynamic cellular structure, necessitates physical connections between the cell cortex and the plus-ends of astral microtubules emanating from the spindle poles (Fig. 2). In this manner, the polarization of cortical cue(s) affords the cell the ability to position the spindle in a biased orientation via interactions between these polarized proteins and microtubule plus-end binding proteins (+TIPs). In this model, +TIPs would serve as the prey in a 'microtubule capture' mechanism by interacting with cortical bait.

Metazoans have evolved a diverse set of +TIPs (Fig. 2A) with varied domain architectures, protein structures and biological functions (Slep, 2010). For example, members of the XMAP215/CLASP family of +TIPs contain multiple domains known as TOG or TOG-like domains, which demonstrate direct plus-end tracking activity in reconstituted systems and bind α/β-tubulin dimers directly (Fig. 2B) (Al-Bassam et al., 2007). XMAP215 members play an essential role in interphase microtubule polymerization and stabilization, which is necessary for proper formation of the mitotic spindle (Slep, 2009; Tournebize et al., 2000). ZYG-9, a C. elegans XMAP215 member, regulates spindle positioning in one-cell-stage embryos (Bellanger et al., 2007). A second autonomously tracking +TIP, end-binding protein 1 (EB1), serves as the cellular workhorse for plus-end protein localization. EB1 itself binds microtubules through an N-terminal calponin homology (CH) domain, an interaction that is enhanced by an acidic, EEY tripeptide motif at the extreme C-terminus (Fig. 2A,C). EB1 binding induces tubulin polymerization in vitro (Slep and Vale, 2007) and is necessary for proper assembly of the mitotic spindle (Rogers et al., 2002). Depletion of EB1 in Drosophila S2 cells causes a significant reduction in microtubule dynamics by inducing extended phases of no growth or of shrinkage, although the overall morphology of interphase microtubules is not disrupted. By contrast, EB1 depletion in mitotic cells induces shortened astral microtubules and fragmented microtubules at prophase along with compacted spindles, detached spindle poles and unfocused nonastral spindles at metaphase (Rogers et al., 2002). Subsequent studies indicated that EB1 stabilizes microtubules through direct interaction with Adenomatous polyposis coli (APC) and the actin polymerizing formin protein Diaphanos (Dia) (Wen et al., 2004). EB1 regulates planar spindle orientation in nonpolarized, cultured epithelial cells through a microtubule stabilization mechanism downstream of β1-integrin adhesion. This effect also requires myosin X-dependent remodeling of the actin cytoskeleton (Toyoshima and Nishida, 2007). It is also worth noting that the budding yeast homolog of EB1, Bim1, functions in spindle orientation through a complex with Kar9 and the myosin Myo2 in a process that involves the guidance of microtubules along actin filaments (Hwang et al., 2003; Korinek et al., 2000). Thus, XMAP215 (also known as Msp5 in Drosophila) and EB1 appear to regulate spindle orientation through the stabilization of spindle structure (Fig. 2).

In addition to its role in the formation and stabilization of spindle microtubules through direct plus-end binding, EB1 recruits an expanding list of indirect +TIPs. Several of these have been shown to regulate spindle orientation, and they fall into two groups that act through distinct EB1-interacting motifs (Kumar and Wittmann, 2012; Slep, 2010). The first group, which consists of the proteins CLIP-170 (also known as CLIP1; CLIP-190 in Drosophila) and p150-dynactin (also known as DCTN1; Glued in Drosophila), bind homodimeric EB1 C-termini through a conserved N-terminal CAP-Gly domain (Fig. 2A,C). CLIP-170 promotes microtubule growth, which has been shown to occur selectively towards the cell cortex and may contribute to directional migration of motile cells (Akhanova et al., 2001; Komarova et al., 2002). The budding yeast homolog of CLIP-170, Bik1p, regulates spindle orientation, possibly through Num1p-mediated dynein activation and/or asymmetric polarization of Kar9 (Miller et al., 2006; Sheeman et al., 2003). As clear orthologs of Num1p and Kar9 are nonexistent or as yet undiscovered, the role of CLIP-170 in spindle orientation in higher eukaryotes remains unclear. A role for p150-dynactin in spindle orientation, however, has been demonstrated. In C. elegans zygotes, p150 (DNC-1) is necessary for proper centrosome rotation along the longitudinal axis and thus spindle orientation (Skop and White, 1998). A subsequent study demonstrating a more robust reduction of p150 expression revealed additional roles in pronuclear migration and centrosome separation (Gönczy et al., 1999). The Drosophila ortholog, Glued, has been shown to regulate both spindle assembly (Siller et al., 2005) and orientation (Siller and Doe, 2008) in larval neuroblasts. These functions of Glued occur via enhanced processivity of dynein, an essential spindle orientation component (discussed in detail below) (Culver-Hanlon et al., 2006). Notably, alterations in Huntingtin (Htt), the protein mutated in the neurodegenerative Huntington’s disease, result in mislocalization of p150 and subsequent spindle misorientation. Htt associates with p150 at spindle poles during mitosis (Gauthier et al., 2004); knockdown of Htt results in a partial loss of spindle-associated p150 along with shortened spindles and unfocused spindle poles (Godin et al., 2010). This loss of Htt-mediated spindle orientation in vivo causes mis-specification of neural progenitor cells in mice, providing evidence for a potential link between spindle orientation, cell fate and disease (see Box 1) (Godin et al., 2010).

Members of the second family of EB1-interacting +TIPs contain the short SxIP polypeptide motif (Fig. 2A,C). The SxIP motif binds in a slightly bent conformation to a hydrophobic cavity and adjacent ‘polar rim’ formed at the EB1 C-terminus (Honnappa et al., 2009). This mode of EB1 interaction, via a short polypeptide motif, allows for a vast diversity of otherwise unrelated proteins to...
Recent evidence suggests this interaction might be regulated by phosphorylation of the SxIP motif or flanking residues, pointing to a potential for cell cycle-dependent regulation (Buey et al., 2012; Honnappa et al., 2009). As noted above, EB1 binds the tumor suppressor protein APC, which contains a clearly defined SxIP motif, and this interaction is required for plus-end APC localization (Fig. 2A,C) (Honnappa et al., 2009). APC has a defined spindle orientation role in several model systems. Firstly, in distal tip cells of the *C. elegans* gonad, APC functions upstream of Rac-mediated spindle orientation, which is required for proper gonadal development (Cabello et al., 2010). APC may also control the timing of spindle rotation in the ABar, a cell present at the eight-cell stage, during *C. elegans* blastomere development (Walston et al., 2004). Secondly, in *Drosophila*, male germline stem cells (GSCs) adhere to a hub cell within the testis niche and undergo asymmetric divisions to produce a proximal self-renewed stem cell and a differentiated spermatogonial cell. Local signals from the...
niches induce self-renewal only in the proximal cell, thus necessitating precise orientation of division perpendicular to the niche cell. Mutations in APC result in mispositioning of the mother centrosome and subsequent defects in spindle orientation (Yamashita et al., 2003). Neuroepithelial cells in the developing *Drosophila* brain also rely on APC for proper spindle orientation; APC RNA interference (RNAi) results in altered planar spindle orientation and defects in proper symmetric divisions of these cells (Lu et al., 2001). Finally, several recent studies have demonstrated a role for APC-mediated spindle orientation in mammalian small intestinal crypt stem cells. For example, an APC truncation mutant causes spindle misorientation that manifests as a switch from A-B to planar orientation (Quyn et al., 2010). Moreover, analysis of intestinal tumors in *APC<sup>min</sup>* (multiple in neoplasia; containing a nonsense truncation mutation at codon 850 in the APC N-terminus) mice demonstrate more severe spindle orientation phenotypes in intestinal crypt cells than in ‘normal-appearing’ heterozygous cells (although these cells show misorientation relative to wild type) (Fleming et al., 2009), suggesting the potential role of proper spindle orientation as a tumor suppressor mechanism (see Box 1). These studies highlight a key role for the tumor suppressor protein APC in spindle positioning and suggest that its EB1 plus-end association is instrumental for promoting interactions with cortical orientation cues (Fig. 2; Table 1).

A recent proteomic study identified microtubule affinity-regulating kinase 1 (MARK1; also known as PAR-1), a serine/threonine kinase, as another SxIP-containing partner of EB1 (Fig. 2A,C) (Jiang et al., 2012). In *C. elegans*, PAR-1 regulates the cortical polarization of factors involved in positioning of the mitotic spindle. Loss of PAR-1 results in embryos that fail to undergo posterior displacement of the mitotic spindle and do not maintain proper A-P spindle orientation (Wu and Rose, 2007). It should be noted, however, that whether the role of *C. elegans* PAR-1 in these processes occurs through regulation of microtubule dynamics per se has not been firmly established (Pellettieri and Seydoux, 2002). Studies in the *Xenopus* embryo found that PAR-1 induces ‘vertical’ spindle orientation in the neuroepithelium, which promotes the generation of deep, differentiated neuronal progeny through asymmetric division patterns (Tabler et al., 2010). Interestingly, PAR-1 has also been shown to function in centrosome orientation checkpoint, a crucial process that ensures the proper oriented division of *Drosophila* male GSCs (Yuan et al., 2012). Although the precise mechanism through which PAR-1 acts in these systems remains unclear, these studies suggest that it may achieve a similar spindle orientation outcome through diverse molecular mechanisms. Moreover, whether EB1 binding and plus-end localization is required for PAR-1 function in spindle orientation has not been examined.

Finally, the kinesin-13 MCAK (also known as KIF2C), belonging to the kinesin family of microtubule motor proteins that typically move toward plus-ends (Hirokawa et al., 2009), also localizes to MT plus-ends via an EB1-SxIP interaction (Fig. 2A,D) (Honnappa et al., 2009; Tanenbaum et al., 2011). MCAK functions as a potent microtubule depolymerase, promoting catastrophe events and thus regulating microtubule length and morphology. Interestingly, a recent report using an elegant reconstitution system demonstrated that the plus-end depolymerase activity of MCAK was coupled to local force production (Oguchi et al., 2011), raising the intriguing possibility that depolymerases could participate in spindle force generation involved in spindle orientation (Fig. 2D) (Grill et al., 2001) (discussed below). This EB1-dependent localization and depolymerase activity is conserved in invertebrates, such as with the *Drosophila* ortholog, Klp10A (Mennella et al., 2005). Loss of Klp10A exacerbates microtubule polymerization, which results in increased density, abundance and length of spindle astral microtubules (Morales-Mulia and Scholey, 2005). Notably, proper regulation of astral microtubules is essential for mitotic spindle orientation in *Drosophila* neuroblasts (Siller and Doe, 2008). Although no evidence yet exists for a direct role of Kinesin-13 depolymerases in spindle orientation, the results discussed above suggest these microtubule enzymes, along with other related proteins (Loughlin et al., 2011), might serve an essential regulatory function at the interface between plus-ends and cortical spindle orientation complexes. Future studies using high resolution, real-time imaging to examine astral MT dynamics directly at the cortex during oriented cell division will be extremely valuable in understanding this process. One potential model would be that shortening of astral MT ends by depolymerase enzymes is necessary to generate gaps that cortical force generators then close through spindle movements coupled to reorientation (Fig. 2D).

In summary, a vast number of diverse protein families localize to plus-ends through distinct mechanisms. This network of protein complexes appears to regulate spindle orientation by different mechanisms, most of which are unknown and will require further studies to resolve completely. More importantly, understanding how these varied mechanisms function together at the cortex/+TIP interface will be an important future research focus. For example, do components of the cortical spindle orientation complex affect the activity of MT stabilizing agents at the +TIP? Conversely, do stable MTs provide regulatory feedback for activity of cortical protein complexes? Does the cortical actin cytoskeleton play an intervening role between the cortex and MTs, perhaps by both maintaining cortical polarity and interacting with spindle MTs? The actin cytoskeleton has been shown to regulate leading edge MT turnover in migratory cells (Gupton et al., 2002). The mechanism(s) by which the actin cytoskeleton contributes to spindle orientation in animal cells remains an area of emerging investigation. Local changes in cortical actin polymerization, induced by the activity of +TIPs, such as APC and Dia, could provide a site for MT capture and spindle stabilization. In turn, stable MTs might provide a site for actin-MT crosslinking protein interaction (Rodriguez et al., 2003). Alternatively, +TIP-induced actin regulation could play an important role in stabilizing the cortical localization and polarity of other spindle orientation regulators (Li and Gundersen, 2008). Regulation of the cortical actin network through external forces (Fink et al., 2011) or integrin-mediated adhesion (Toyoshima and Nishida, 2007) can bias spindle orientation in cultured cells. Also, the Spire and Formin families of actin regulators have been shown to cooperate in the asymmetric positioning of the meiotic spindle during oocyte division by establishing a cortical actin network necessary for spindle movement (Pfendler et al., 2011). Finally, how do MT enzymes that regulate growth and catastrophe dynamics contribute to +TIP architecture, and how do MT dynamics regulate coupling to cortical complexes? It seems plausible that the geometry of the MT plus-ends relative to the cell cortex is vital for establishing productive interactions that promote spindle movements (Su et al., 2012). We suggest that a significant role of +TIPs is to shape MT architecture dynamically to provide effective cortical coupling necessary for spindle orientation, possibly through enhancement of cortical force generators (Fig. 2D) (Kozlowski et al., 2007). Although a significant effort has been made to understand the
cortical cues involved in spindle orientation, further studies that
dress the role of the microtubule plus-end will help to elucidate
the molecular interplay involved in cortical coupling.

**Bridging the gap between polarity and spindle positioning: Pins-Dlg-Khc-73, an MT-capturing complex**

One fundamental aspect of spindle orientation is the ability of
cortical cues to capture the spindle physically, probably as an initial
step in dictating its ultimate position. Our discussion of +TIPs
above demonstrates the variety of potential microtubule prey in this
capture event. What then is the cortical bait that secures the spindle
connection? How are these cortex-microtubule interactions
established, maintained and regulated? What are the functional
consequences of spindle capture on other aspects of the spindle
orientation process? Static spindle-cortex connections could allow
for proper localization and/or activation of pathways that act in
subsequent steps; for example, the microtubule-mediated
offloading of force-generating molecules to the cortex (Markus and
Lee, 2011; Pecreaux et al., 2006). Below, we discuss the regulation
of a recently characterized cortical spindle-capturing complex and
present a model for how it cooperates with an evolutionarily
conserved force-generating complex.

As discussed earlier, the evolutionarily conserved polarity protein
Pins regulates spindle orientation in a variety of cell types within
diverse organisms to achieve distinct biological outcomes (David et
al., 2005; Du et al., 2001; Schaefer et al., 2001). Recently, we
identified a ‘microtubule capture’ pathway in *Drosophila* that
involves a Pins-Dlg-Khc-73 complex (Fig. 3A). Discs large (Dlg)
directs its linear motif to the ‘Linker’ domain of Pins
and is required for proper spindle orientation in both an induced
cell polarity S2 cell system and neuroblasts (Johnston et al., 2009;
Siegrist and Doe, 2005). Interestingly, this interaction is dependent
upon phosphorylation of Pins by the mitotic, centrosomal kinase
Aurora-A (Aurka in mammals; *Aur in Drosophila*), as is Pins-
mediated spindle orientation (Johnston et al., 2012; Johnston et al.,
2009; Lee et al., 2006a). In this model, Dlg associates with the cell
cortex and localizes with polarized Pins (Siegrist and Doe, 2005).
Dlg then interacts with Khc-73, a kinesin motor that moves
towards and localizes at microtubule plus-ends (Hanada et al., 2000;
Huckabat et al., 2011; Siegrist and Doe, 2005); Dlg binding appears
to enhance Khc-73 activity in vitro (Yamada et al., 2007). Dlg
binding occurs through a unique domain of Khc-73, termed the
maguk binding stalk (MBS). Thus, Dlg serves as an adaptor protein
that links polarized Pins at the cell cortex to the microtubule plus-
end motor, Khc-73, and loss of any of these three components
results in spindle orientation defects (Fig. 3A). Interestingly, both
the Pins Liniker and Khc-73 MBS domains bind to the guanylate
kinase (GUK) domain of Dlg (Johnston et al., 2009; Siegrist and
Doe, 2005), suggesting that they bind at distinct sites within the
GUK domain. The structural basis for phosphorylation-dependent
Pins-Dlg complex formation has recently been detailed (Johnston et
al., 2012; Zhu et al., 2011a) but a structural analysis of the MBS
domain alone or in complex with Dlg is lacking. It remains to be
determined whether Khc-73 motor activity per se is required for
spindle orientation, although misexpression of the MBS domain
alone functions as a dominant negative, suggesting that other
domains are indeed required (Johnston et al., 2009; Siegrist and
Doe, 2005).

Using the induced polarity assay in *Drosophila* S2 cells, we
found that although full-length Pins could robustly orient the
mitotic spindle, the Linker domain alone was sufficient for a
‘partial orientation’ phenotype. We found that this intermediate
activity of the isolated Linker domain typically oriented spindles
precisely at the distal edge of the cortical Pins crescent (as opposed
to the center of the crescent with full-length Pins). Live-cell
imaging experiments revealed that mitotic spindles with poles
contacting non-Pins-expressing regions of the cell cortex rotated
slowly before becoming secured at the edge of the Linker crescent
(Johnston et al., 2009). These results suggest that the Pins-Dlg-
Khc-73 complex functions to capture the mitotic spindle statically
by establishing a physical connection between cortical polarity and
microtubule plus-ends (Fig. 3A, Fig. 4).

Whether such a capture mechanism occurs with other polarity
proteins remains to be determined; the plethora of microtubule
plus-end binding proteins discussed above suggests this may
indeed be a widespread component of spindle orientation. Also
unclear is the precise function of this spindle-capturing activity.
Does static spindle capture allow for activation of additional
pathways at specific sites? Are other regulators localized or
offloaded to the cortex more efficiently from the astral
microtubules of a stationary spindle? Do cortical force generators
act with higher fidelity on a ‘captured’ spindle? Alternatively, the
capture of microtubules at the edge of a cortical polarity complex
may be sufficient for spindle orientation in certain cellular contexts.

**Linking polarity to spindle force generation: the role of the Mud-dynein complex**

Perhaps one of the most well characterized effectors in spindle
orientation pathways is the Mud-dynein complex (Figs 3, 4)
(Gönczy, 2008; Siller and Doe, 2009). *Drosophila* Mud is a large
(>2000 amino acid) coiled-coil protein originally identified as a
regulator of neuroblast proliferation (Guan et al., 2000). Mud
shares overall weak sequence homology with the *C. elegans*
and mammalian functional orthologs LIN-5 and NuMA, respectively
(Siller et al., 2006). However, these proteins share a relatively
higher conservation in a small C-terminal domain termed the Pins-
binding domain (PBD). The PBD directly interacts with the Pins
tetratricopeptide repeat (TPR) domain, and this interaction requires
prior Ga protein-mediated relief of an autoinhibitory conformation
between the TPR and GoLoco domains of Pins (Du and Macara,
2004; Nipper et al., 2007). This interaction recruits Mud to the Pins
cortical crescent, thereby polarizing Mud localization (Fig. 3A).
Notably, Mud also localizes to spindle poles and microtubules,
independent of Pins, where it functions in the establishment and
proper maintenance of focused microtubules (Bowman et al., 2006;
Izumi et al., 2006; Silk et al., 2009; Siller et al., 2006). Additional
studies have shown that the scaffold protein, Canoe (Cno),
promotes formation of the cortical Pins-Mud complex through
several small GTPases (e.g. Ran) and is important for proper
spindle orientation (Fig. 3A) (Speicher et al., 2008; Wee et al.,
2011).

The function of Mud in Pins-mediated spindle orientation
appears to be that of a cortex-to-microtubule adaptor protein,
analogous to the role played by Dlg (Fig. 4). Mud associates
with cytoplasmic dynein, which serves as the sole minus-end-directed
microtubule motor protein during mitosis. Dynein is a large protein
complex consisting of core light, intermediate and heavy chains,
the latter of which possesses the ATPase activity necessary for
movement along microtubules. The non-catalytic subunits serve as
binding sites for additional regulatory proteins, some of which are
essential for dynein activity and, thus, spindle orientation (e.g.
p150-dynactin and LIS-1) (Kardon and Vale, 2009; Siller and Doe,
2008). Precisely how Mud interacts with dynein remains unclear,
although recent evidence suggests that Mud associates with the dynein light chain subunit, Cut up (Ctp), which is required for centrosomal localization of Mud and spindle orientation in neuroblasts (Fig. 3A) (Wang et al., 2011). Recent evidence also suggests that a cortically localized component of dynein, via specific interaction with Mud, is required for proper spindle orientation (Kotak et al., 2012). Thus, Mud facilitates a link between the dynein motor complex and cortically polarized Pins.

In contrast to the ‘spindle capture’ mechanism proposed for Khc-73, dynein is thought to provide the ‘force generation’ required for rapid spindle repositioning (Gönczy, 2008). The minus-end directional movement of dynein along astral microtubules (towards the spindle pole) results in net pulling forces on the spindle towards cortical Pins tethered by Mud (Gusnowski and Srayko, 2011) (Fig. 3A). Spindle-severing experiments in C. elegans embryos have established a role for the Pins-Mud-dynein complex in generating the force necessary for spindle positioning (Colombo et al., 2003; Nguyen-Ngoc et al., 2007). Additionally, the polarized localization of the Pins-Mud complex results in unequal pulling forces at one spindle pole, which probably contributes to the asymmetry in spindle morphology and subsequent size of daughter cells (Bowman et al., 2006; Izumi et al., 2006; Siller et al., 2006).

The requirement of the Mud-dynein complex for proper spindle positioning has been firmly established in several systems. However, with our ‘induced polarity’ assay, we found that this complex was not sufficient to regulate spindle orientation in isolation (Johnston et al., 2009). Instead, Mud-dynein functioned synergistically with Dlg-Khc-73 downstream of Pins. Whether direct crosstalk exists between these two complexes, or if they perform otherwise independent functions that diverge below Pins, remains to be determined. Kinesin family members play an important role in plus-end delivery of various cargo proteins (Hirokawa et al., 2009), suggesting a model in which Khc-73 aids in localization of Pins-Mud pathway components. Live-cell analysis revealed that the Pins<sup>TPR</sup>-Mud-dynein component was required the for rapid, dynamic spindle movement necessary for robust orientation, further strengthening the model of Mud-dynein complex-mediated force generation. An important question for future investigation will be how the minus-end Mud-dynein pathway synergizes with that of the plus-end Dlg-Khc-73. Interestingly, cooperative involvement between opposite end-directed motor protein pathways has also been seen in meiotic spindle positioning (Ellefson and McNally, 2009).

Recently, the Mud-dynein complex was shown to regulate spindle orientation mediated through the Wnt-Frizzled (Fz) effector Dishevelled (Dsh) in both Drosophila and zebrafish (Fig. 3B) (Ségalen et al., 2010). The Fz-Dsh complex had long been known to influence planar spindle orientation in the pl division of Drosophila SOPs; however, a bona fide downstream pathway was unknown (Bellaïche et al., 2001; Gho and Schweisguth, 1998). Both genetic and biochemical evidence demonstrated that Mud acts directly downstream of the Dishevelled-Egl10-Pleckstrin (DEP)
domain of Dsh (Fig. 3B). Although this model is analogous to that of Mud acting downstream of the Pins TPR domain, the DEP and TPR domains share no similarities in primary sequence or tertiary structure. These results, therefore, highlight a striking example of convergent evolution of a shared signaling pathway downstream of divergent upstream activators. Whereas the minimal TPR-binding peptide has been identified in Mud and its structure determined (Smith and Prehoda, 2011; Zhu et al., 2011b), the precise molecular basis for the Dsh-Mud complex remains to be determined. Whether additional pathways act downstream of Dsh is unclear, and a complete molecular model for Dsh-mediated spindle orientation remains to be described (Fig. 3B).

Despite ample research establishing an important role for the Mud-dynein complex in spindle orientation, many questions remain regarding its molecular function. How does Mud binding affect dynein activity? Structural and biophysical data are beginning to illuminate the conformational changes dynein might undergo to achieve its movement downstream of microtubules (Carter et al., 2011; Redwine et al., 2012). Does Mud actively alter these structural transitions to enhance dynein function? How is synergy achieved between Mud-dynein and Dlg-Khc-73 complexes downstream of Pins? Despite diverse sequence determinants for binding, do Pins and Dsh regulate Mud-dynein function in similar ways? Does the Mud-dynein pathway function together with a secondary pathway in Dsh-mediated spindle orientation?

**Phosphoregulation of spindle orientation: an assortment of kinases regulates spindle orientation**

Intense efforts have recently been focused on identifying additional regulators of spindle orientation. Notably, a network of protein kinases has now been shown to influence spindle positioning through various mechanisms, including regulation of microtubule-architecture (e.g. LKB and c-Src), regulation of cortical cue-adapter complexes (e.g. ABL1 and AurA) and localization of microtubule regulators such as Ran (gray) (e.g. PAK).

**Fig. 4. A summary of the mechanisms that contribute to spindle orientation.** Cortical spindle orientation cues, such as Pins and Dsh (dark blue), polarize to specific regions of the cell cortex. Cortical adapters, such as Dlg and Mud/NuMA (light blue), are then recruited by direct interactions with cortical cues. Polarized cortical cues and adapters interface with spindle microtubules through interactions with various plus-end localized microtubule-binding proteins, including motor proteins (e.g. Khc-73 and Dynein; red) and additional regulatory complexes (e.g. EB1-APC; green). These elements provide microtubule capture and force generation effects that coordinate spindle positioning. Additional plus-end binding proteins aid in microtubule dynamics, such as polymerases (e.g. XMAP215, dark green) and depolymerases (e.g. MCAK, light green), which are necessary for proper spindle assembly and function. An assortment of protein kinases (yellow) have been shown to regulate spindle orientation through various mechanisms, including regulation of microtubule-architecture (e.g. LKB and c-Src), regulation of cortical cue-adapter complexes (e.g. ABL1 and AurA) and localization of microtubule regulators such as Ran (gray) (e.g. PAK).
Protein-protein interactions underlying the macromolecular structure and the molecular mechanisms involved in phosphorylation-dependent spindle orientation are unanswered questions awaiting further research in this area. What are the specific residues that are phosphorylated in each of these components? What are the spatiotemporal constraints of phosphoregulation? Finally, what are the phosphatases that direct dephosphorylation and how are they regulated throughout the cell cycle?

Conclusions

The ability to orient cell division allows for the generation of cellular diversity and for regulated spatial placement of daughter cells within tissues. Nature has evolved sophisticated signaling pathways that allow cortical cues to communicate with the mitotic spindle to influence its position prior to cell division (Fig. 4). The microtubule plus-end is a node for the localization of diverse +TIPs that serve as spindle-capture prey. Whether autonomously localized proteins, such as XMAP215 and CLIP-170, or cargo of the essential +TIP EB1, such as APC, these assorted messengers regulate microtubule organization and aid in the cortical connection of the spindle. The cortex, by contrast, contains orientation cues, such as Pins and Dsh, that organize protein complexes to function as spindle-capturing bait as well as force generators for the rapid spindle motions required for repositioning during cell division. In this Review, we have summarized many of the seminal findings describing the molecular basis for spindle orientation. Although significant progress has been made on this front, it is apparent that the surface has merely been scratched. What additional +TIPs might serve as spindle-capturing prey? How do microtubule dynamics and the regulatory proteins involved relate to spindle capture and force generation? What other cortical complexes serve as spindle orientation regulators, and in what cell types do these function? Is dynein the lone force generator involved in spindle orientation, and how is the dynein complex regulated at the molecular level? How do divergent spindle capturing events cooperate with the conserved force-generating complex? Finally, does the association between spindle orientation defects and human disease play a causal role, and can spindle orientation regulators represent potential therapeutic targets? We excitedly await further analysis of this fascinating biological process.

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