Identification of an Aurora-A/Pins^{LINKER}/ Dlg Spindle Orientation Pathway using Induced Cell Polarity in S2 Cells

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SUMMARY

Asymmetric cell division is intensely studied because it can generate cellular diversity as well as maintain stem cell populations. Asymmetric cell division requires mitotic spindle alignment with intrinsic or extrinsic polarity cues, but mechanistic detail of this process is lacking. Here, we develop a method to construct cortical polarity in a normally unpolarized cell line and use this method to characterize Partner of Inscuteable (Pins; LGN/AGS3 in mammals) -dependent spindle orientation. We identify a previously unrecognized evolutionarily conserved Pins domain (Pins^{LINKER}) that requires Aurora-A phosphorylation to recruit Discs large (Dlg; PSD-95/ hDlg in mammals) and promote partial spindle orientation. The well-characterized Pins^{TPR} domain has no function alone, but placing the Pins^{TPR} in *cis* to the Pins^{LINKER} gives dynein-dependent precise spindle orientation. This "induced cortical polarity" assay is suitable for rapid identification of the proteins, domains, and amino acids regulating spindle orientation or cell polarity.

INTRODUCTION

Asymmetric cell division (ACD) occurs when one cell divides to generate two molecularly distinct daughter cells. ACD requires precise alignment of the mitotic spindle with the intrinsic or extrinsic polarity axis so that cellular components, such as fate determinants, are partitioned into different daughter cells (Knoblich, 2008). Recent work has highlighted the importance of ACD in generating cell diversity during early embryogenesis and in maintaining stem cell pool size (Doe, 2008). Thus, characterization of the ACD mechanisms, including cell polarization and spindle orientation, is important for understanding many aspects of development and disease.

Drosophila neural progenitors, or neuroblasts, are an excellent model system for studying ACD. Neuroblasts have apical/basal

polarity, and they align their mitotic spindle with this polarity axis to generate a self-renewed apical neuroblast and a differentiating basal daughter cell. Cell fate determinants have been identified that partition into the apical neuroblast to maintain its fate (e.g., atypical protein kinase C [aPKC]) and that partition into the basal daughter cell to induce differentiation (e.g., Miranda/Prospero) (reviewed in Doe, 2008). Progress has also been made on identifying proteins required for apical/basal spindle orientation. These include proteins that form an apical cortical crescent over one spindle pole, such as Inscuteable (Insc; mlnsc in mammals), Partner of Inscuteable (Pins; LGN/ AGS3 in mammals), Gα_{i/o}, Discs large (Dlg), Scribble, and Mushroom body defect (Mud; NuMA in mammals; LIN-5 in C. elegans); proteins enriched on centrosomes, such as Centrosomin, Sas-4 (CenpJ in mammals), Dynein, Dynactin, Lis1; and the mitotic kinases Aurora-A and Polo (Knoblich, 2008). Virtually all of these proteins are evolutionarily conserved and have a similar function in regulating spindle alignment in yeast, C. elegans, and mammals (Siller and Doe, 2009).

Despite progress in identifying components involved in cortical polarity and spindle orientation, much remains unknown. Here, we describe a system for generating cell polarity and spindle orientation in the normally unpolarized *Drosophila* S2 cell line and use this system to test individual proteins, protein domains, and amino acids for their role in spindle orientation.

RESULTS

Induced Cell Polarity in the Drosophila S2 Cell Line

Transfection-induced expression of the homophilic cell adhesion molecule Echinoid (Ed) can induce cell-cell adhesion in S2 cells and restrict Ed protein to the site of cell-cell contact (Bai et al., 2001). We adapted this method to induce cortical polarity of a heterologous protein by fusing the protein of interest to the Ed cytoplasmic terminus. Ed:green fluorescent protein fusions (Ed:GFP) formed distinct cortical crescents containing the majority of the cortical protein (Figures 1A and 1C). Ed fusion proteins are also detected in cytoplasmic vesicles (Figure 1A, asterisk), as expected for transmembrane proteins, but the presence of vesicles had no effect on the cortical polarity or spindle

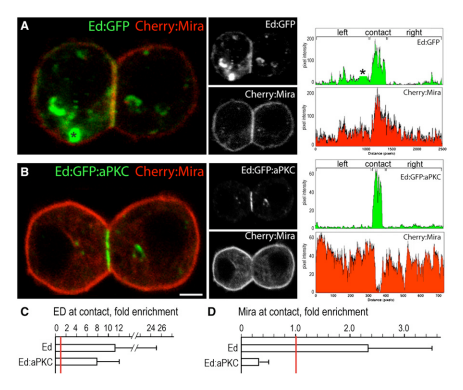


Figure 1. Induced Cell Polarity in Drosophila S2 Cells

(A) The Echinoid (Ed) induced cortical polarity in S2 cells. Cortical polarization of Ed:GFP fusion protein (green); there is also cytoplasmic vesicle staining (asterisk); Cherry:Miranda (Mira) protein (red) shows less cortical enrichment. Right: quantification of cortical pixel intensity (vesicle staining removed from plot).

(B) Induced functional cortical polarity in S2 cells. Polarized Ed:aPKC can exclude Cherry:Miranda from the cortex, thereby inducing Cherry:Mira cortical polarization. Right: quantification of cortical pixel intensity. The scale bar represents 3 µm.

(C) Ed and Ed:aPKC are enriched at the site of cell-cell contact. Pixel intensity was measured for left (L), right (R), and contacting (C) cortical domains, and fold enrichment at the contact site was calculated (C/L+R). The red line indicates a ratio of 1.0 (no enrichment or exclusion). n = 17 (Ed) and 18 (Ed:aPKC).

(D) aPKC excludes Miranda from the cortex. Cherry:Mira pixel intensity levels were measured as described in (C). The red line indicates a ratio of 1.

orientation function of Ed fusion proteins (Table S1 available online). To test whether this system could be used to generate functional cortical polarity matching that of neuroblasts, we generated aPKC cortical polarity by expressing an Ed:GFP: aPKC fusion protein. We observed that cortical aPKC was necessary and sufficient to exclude Miranda from the cortex in S2 cells (Figures 1B and 1D), similar to larval neuroblasts (Rolls et al., 2003). We conclude that the Ed cell adhesion molecule can be used to induce functional cortical polarity in S2 cells.

Induced Pins Cortical Polarity Promotes Spindle Orientation

Drosophila neuroblasts show tight alignment of the mitotic spindle with the apical cortical domain, and many apical proteins are required for proper spindle orientation (Knoblich, 2008). However, it is unknown which domain of each protein is essential for spindle orientation function, nor is it known which provide cortex-to-microtubule links or which act indirectly by regulating cortical polarity. Here, we sought to test individual apical proteins for their ability to promote spindle orientation in Drosophila S2 cells, with the goal of understanding the function of individual proteins, protein domains, and specific amino acids.

We generated protein crescents as described above and measured the angle of the mitotic spindle relative to the center of the Ed cortical crescent (Figure 2B). We measured the spindle axis using two different methods—staining for the spindle marker α -tubulin or for the spindle pole marker Cnn—and both gave identical results (Table 1). Precise alignment of the spindle with the crescent would generate a mean spindle angle near 0° , whereas random spindle orientation would give a mean spindle angle near 45° (see the Experimental Procedures for details). Control experiments showed that polarization of Ed:FLAG or

Ed:GFP resulted in nearly random spindle orientation with a mean spindle angle of 53° and a standard deviation of 23° (Figure 2C; Table 1). The large standard deviation results from the random distribution. We estimated the standard error of the spindle angle measurements to be ±3° based on the analysis of multiple independent experiments (see the Experimental Procedures). These control experiments allowed us to conclude that the spindle orientation is not affected by Ed, FLAG, GFP, or potential cell shape changes due to cell-cell adhesive contacts. We next tested various proteins required for spindle orientation in neuroblasts for their ability to induce spindle orientation in S2 cells. Induced cortical crescents of Insc, EB1, or aPKC failed to orient the mitotic spindle (Table 1; although aPKC could efficiently displace Miranda from the cortex, see above). In contrast, full-length Pins (Ed:PinsFL) could partially orient the mitotic spindle (27° ± 15°; Figure 2D; Table 1), consistent with its requirement for spindle orientation in Drosophila neuroblasts (Parmentier et al., 2000; Rebollo et al., 2007; Schaefer et al., 2000; Siegrist and Doe, 2005; Yu et al., 2000).

Full-length Pins exists by default in a "closed" or autoinhibited state because of intramolecular binding between its seven N-terminal tetratricopeptide repeat (TPR) domains and its three C-terminal Gai-binding (GoLoco) motifs; binding of Gai to the C-terminal GoLoco motif region can "open" Pins (Du and Macara, 2004; Nipper et al., 2007). Gai levels are low in S2 cells (Figure S1), raising the possibility that Ed:Pins FL is not fully active due to autoinhibition. To test this hypothesis we coexpressed Gai and Ed:Pins FL, and observed a clear improvement in the mean spindle angle to $17^{\circ} \pm 15^{\circ}$ (n = 55; Figure 2E; Table 1). Single amino acid mutations (Arg > Phe) to each GoLoco motif, which prevent Pins from binding Gai (Willard et al., 2004), were found to block Gai-induced improvement of spindle orientation

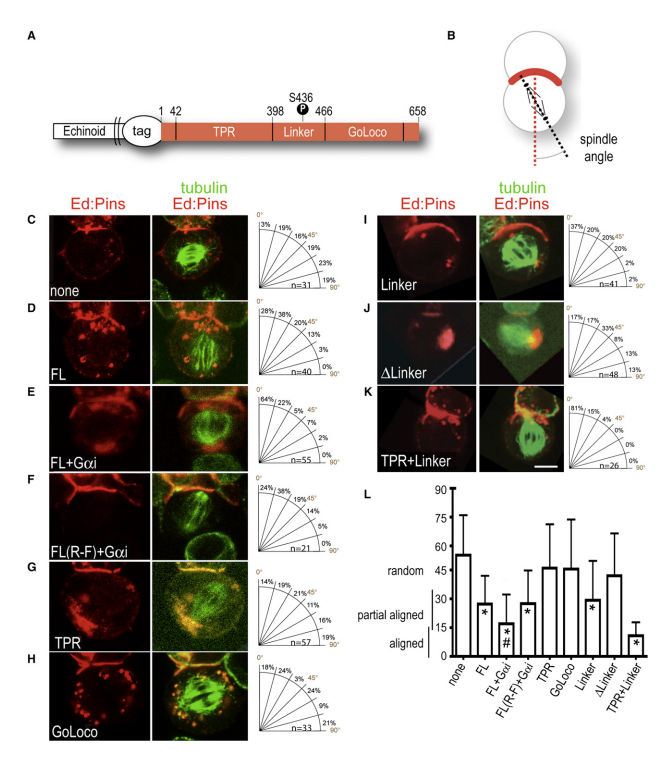


Figure 2. The Pins TPR and Linker Domains Synergistically Orient the Mitotic Spindle in *Drosophila* S2 Cells
(A) N-terminal TPR domain (amino acids 42–398), central linker region (399–466), and "GoLoco domain" containing three GoLoco motifs (467–658). Serine 436 (S436; black ball).

(B) Spindle angle assay. A vector perpendicular to the Ed crescent (red dashed line) and along the spindle axis (black dashed line) indicates spindle angle. (C–K) Pins domains were fused in frame to Echinoid tagged with a FLAG epitope (Ed-FLAG) and transfected into S2 cells. Cells were stained for FLAG (red) and α -tubulin (green). FL, Pins full length; FL+G α i, PinsFL cotransfected with G α i (G α i is colocalized with Pins, not shown); FL(R-F)+G α i, PinsFL with a mutated G α i binding domain cotransfected with G α i (G α i is delocalized, not shown); TPR, Pins TPR domain alone; GoLoco, Pins GoLoco domain alone; Linker, Pins Linker domain alone; ALinker, Pins full-length protein lacking the linker domain; TPR+Linker, Pins TPR and Linker domains. The scale bar represents 3 μ m.

 $(26^{\circ}\pm24^{\circ}; n=27;$ Figure 2F; Table 1). This mirrors the situation in wild-type neuroblasts, where $G\alpha i$ is required for tight alignment of the spindle with the Pins crescent (Nipper et al., 2007). We conclude that $G\alpha i$ is required for Pins to accurately orient the mitotic spindle in S2 cells, similar to the role of $G\alpha i/Pins$ in neuroblasts.

We next tested different fragments of Pins for their ability to promote spindle orientation: the N-terminal TPR region (residues 1-398), the central linker (residues 399-466), and the C-terminal GoLoco region (residues 467-658). The Pins^{TPR} domain is the best candidate for providing Pins spindle orientation function, because it binds the microtubule-binding protein Mud, which is required for spindle orientation in Drosophila and C. elegans (Bowman et al., 2006; Izumi et al., 2006; Lorson et al., 2000; Siller et al., 2006). Surprisingly, we found that the Pins^{TPR} domain alone had no ability to orient the mitotic spindle in our S2 assay $(46^{\circ} \pm 25^{\circ}; Figure 2G; Table 1)$. The GoLoco region also failed to orient the spindle (46° \pm 28°; Figure 2H; Table 1). The only region of Pins that promoted spindle orientation was the previously uncharacterized linker region (29° ± 20°; Figure 2I; Table 1), and deletion of the linker from full length Pins abolished spindle orientation activity (42° ± 25°; Figure 2J; Table 1).

Pins^{LINKER} has spindle orientation activity that is similar to the "closed" Pins full-length protein, but not as good as the "open" (Gai bound) Pins full-length protein. This suggests that the Pins LINKER is active when Pins is in the "closed" form and that "opening" of Pins reveals a second domain that can improve Pins^{LINKER}-mediated spindle orientation. What is the second domain? Pins^{LINKER+GOLOCO} did not improve on Pins^{LINKER} spindle orientation, and Pins^{GOLOCO} alone has no spindle orientation function (Table 1). In contrast, addition of the Pins^{TPR} domain (Pins^{TPR+LINKER}) showed striking improvement in spindle orientation over Pins^{LINKER} alone ($10.7^{\circ} \pm 7.1^{\circ}$; Figure 2K; Table 1). In fact, PinsTPR+LINKER spindle orientation was comparable to Pins^{FL} + Gαi (17° ± 15°; Table 1) or neuroblast spindle orientation in vivo (Nipper et al., 2007; Siller et al., 2006; Siller and Doe, 2008), despite the absence of the GoLoco domain and endogenous $G\alpha i$. We draw two conclusions from these data. First, the Pins GOLOCO domain and Gai are not essential for Pinsmediated spindle orientation; their function appears limited to Pins "opening" and cortical tethering. Second, the Pins TPR can improve the spindle orientation ability of the Pins^{LINKER} domain to a level matching Gai-activated full length Pins.

Pins^{LINKER} Acts through Dlg and Khc-73 to Promote Spindle Orientation

In this section we explore the mechanism of Pins^{LINKER} -mediated spindle orientation, and in the following section we investigate how the Pins^{TPR} domain improves Pins^{LINKER} activity. Pins binds the tumor suppressor protein Dlg (Bellaiche et al., 2001; Sans et al., 2005), and Dlg is required for spindle orientation in *Drosophila* neuroblasts (Siegrist and Doe, 2005). Here, we test

whether Dlg is required for Pins^{LINKER}-mediated spindle orientation. S2 cells transfected with Ed alone or with Ed-Pins lacking the linker domain localize endogenous Dlg primarily in the cytoplasm (Figures 3A and 3B). In contrast, we observed clear Dlg recruitment to Pins^{LINKER} or Pins^{TPR+LINKER} cortical crescents (Figures 3C and 3D). Moreover, dlg RNA interference (RNAi) reduced Dlg levels and abolished Pins^{LINKER}-mediated spindle orientation (42° ± 24°; Figure 3F; Table 1), as well as blocked spindle orientation conferred by full-length Pins and Pins^{TPR+LINKER} (Table 1); Pins^{TPR+LINKER} cells treated with control RNAi constructs (bazooka or insc) had no effect on spindle orientation (Table 1). If Pins^{LINKER} recruits DIg to promote spindle orientation, we should be able to bypass Pins by inducing Dlg cortical polarity. We previously showed the Dlg guanylate kinase-like (DlgGK) domain was necessary for proper spindle orientation in neuroblasts (Siegrist and Doe, 2005); here, we test whether the Dlg^{GK} domain is sufficient for spindle orientation in S2 cells. We used Ed to form a DIgGK cortical crescent and found that it had spindle orientation ability similar to that of Pins LINKER (28 $^{\circ}$ \pm 20 $^{\circ}$; Figure 3G; Table 1). We conclude that the Pins recruits Dlg to the cortex, where the Dlg GK domain promotes spindle orientation.

How does the DIgGK interact with spindle microtubules? A good candidate is Khc-73 (Kif13b in mammals), a microtubule plus end-directed kinesin motor that can bind both microtubules and the DlgGK domain (Hanada et al., 2000) and is required for proper spindle orientation in Drosophila neuroblasts (Siegrist and Doe, 2005). We found that expression of a dominant-negative Khc-73 fragment prevents Pins^{LINKER}-mediated spindle orientation (51.9° ± 23.5°; Figure 3I; Table 1). Similarly, RNAi directed against Khc-73 completely blocked Pins^{TPR+LINKER}mediated spindle orientation (40.5° ± 21.9°; Figure 3J; Table 1), as well as Dlg^{GK}-mediated spindle orientation (46.7° ± 30.5°; Figure 3H; Table 1). khc-73 RNAi does not block recruitment of Dlg to the Pins^{LINKER} cortical crescent (Figure 3E), consistent with Khc-73 acting downstream of cortical Dlg, rather than transporting Dlg to the cortex as proposed for mammalian cells (Hanada et al., 2000). We conclude that Khc-73 acts downstream of Dlg to promote Pins^{LINKER}/Dlg^{GK}-mediated spindle

Pins^{TPR} Recruits Mud and Acts in *cis* to Improve Pins^{LINKER}-Mediated Spindle Orientation

In this section we investigate how Pins^{TPR} can significantly improve the spindle orientation ability of the Pins^{LINKER} domain, despite having no spindle orientation function on its own. The Pins^{TPR} binds Insc and Mud, which are both required for spindle orientation (Bowman et al., 2006; Izumi et al., 2006; Kraut et al., 1996; Siller et al., 2006). S2 cells express detectable levels of Mud but not Insc (Figure S1), and Ed:Insc has no spindle orientation ability in S2 cells (Table 1), so we tested the role of Mud in Pins^{TPR+LINKER} spindle orientation. Pins^{TPR+LINKER} was

⁽L) Quantification. Mean spindle angle and standard deviation are shown (sampling error is estimated at $\pm 3^\circ$; see the Experimental Procedures). The Pins TPR, GoLoco, and Δ Linker proteins show no spindle orientation (30° – 60° spindle angles); Pins FL, FL(R-F)+G α i, Linker proteins show partial spindle orientation ($\sim 15^\circ$ – 30° spindle angles); and Pins FL+G α i and TPR+Linker proteins show good spindle orientation ($\sim 0^\circ$ – 15° spindle angles). *, highly significant compared to Ed alone "none" (p < 0.01); #, significant spindle orientation compared to Pins FL (p < 0.05). Other proteins showed no difference from Ed alone (p > 0.05). Error bars represent standard deviation from the mean for all experimental values.

Table 1. Echinoid Fusion Protein Spindle Orientation Ability					
Echinoid:FLAG Fusion Partner	Average Angle	Standard Deviation	n	Pathway	
Pins ^{TPR+LINKER} (in <i>cis</i>)	10.7	7.1	26	TPR+LINKEF	
^a Pins ^{TPR+LINKER} (in <i>cis</i>)	11.2	5.6	29	"	
Pins ⁽¹⁻⁴⁴⁶⁾	13.4	13.3	25	"	
Pins ^{TPR+LINKER} (S436D) +	13.6	12.9	35	"	
aur-A RNAi	10.0	12.5	33		
Pins ^{TPR+LINKER} (S436D)	13.7	11.3	50	cc .	
Pins ^{TPR+LINKER} + baz RNAi	13.7	9.3	24	"	
Pins ^{FL} + Gα	17.2	15.4	55	"	
Pins ^{FL} (Arg-Phe)	25.9	23.8	27	LINKER	
Pins ^{FL}	27.2	14.8	40	"	
Pins ^{TPR} + Pins ^{LINKER}	27.2	18.1	34	"	
(in <i>trans</i>)	21.2	10.1	04		
^b Pins ^{LINKER}	27.3	15.6	33	"	
Pins ^{FL} (Arg-Phe) + Gαi	27.5	17.3	21	"	
Dlg ^{GK}	27.5	20.2	41	"	
Pins ^{TPR+LINKER} + <i>dlc</i> RNAi	27.9	22.5	33	"	
Pins ^{TPR+LINKER} + <i>mud</i> RNAi	29.1	18.1	28	ш	
Pins ^{LINKER}	29.2	20.5	41	u	
Dlg ^{GK} + <i>mud</i> RNAi	29.3	18	29	"	
Pins ^{LINKER} + dlc RNAi	29.9	17.5	33	"	
Dlg ^{GK} + <i>dlg</i> RNAi	30.5	22.6	16	"	
Pins ^{TPR+LINKER} + <i>lis1</i> RNAi	31.1	25.9	22	"	
Pins ^{TPR+LINKER} (N226F)	31.4	23.8	26	"	
°Pins ^{LINKER}	32.4	17.0	30	"	
Pins ^{TPR+LINKER} +	40.5	21.9	41	NONE	
khc-73 RNAi					
Pins ^{FL} + <i>dlg</i> RNAi	40.8	27.1	48	u	
Pins ^{TPR+LINKER} +	41.0	24.5	53	"	
aur-A RNAi					
Pins ⁽¹⁻⁴²⁶⁾	41.1	23.1	21	"	
Pins ⁽¹⁻⁴³⁵⁾	41.3	25.1	24	"	
Pins ⁽¹⁻⁴⁰⁶⁾	41.7	24.1	28	"	
Pins ⁽¹⁻⁴¹⁶⁾	41.9	25.2	26	"	
Pins ^{LINKER Deleted}	42.0	24.6	48	"	
Pins ^{TPR+LINKER} + dlg RNAi	42.1	23.7	23	"	
Pins ^{TPR+LINKER} (S436A)	43.0	20.8	44	"	
Pins ^{TPR+LINKER} +	43.5	25	39	"	
Nocodazole					
Pins ^{LINKER} + dlg RNAi	43.7	27.5	48	"	
EB1	44.5	29.7	27	"	
Dlg ^{GK} + Nocodazole	45.2	25	25	"	
Pins ^{GoLOCO}	45.7	28.2	33	"	
Pins ^{TPR}	46.1	25.2	57	"	
Dlg ^{GK} + <i>khc-73</i> RNAi	46.7	30.5	21	"	
dInsc ⁽²⁵²⁻⁵⁸³⁾	48.9	27.6	15	"	
aPKC	50.5	22.5	19	"	
Pins ^{TPR+LINKER} + Khc-	51.9	23.5	11	"	
73 ^{MBS}					

Table 1. Continued				
Echinoid:FLAG Fusion Partner	Average Angle	Standard Deviation	n	Pathway
None	52.8	23.3	31	"
^a None	46.5	27.5	33	"

All spindle angle measurements were made using tubulin stains except where noted.

able to recruit Mud to the cortex (Figure 4I), and mud RNAi reduced Pins^{TPR+LINKER} spindle orientation activity to match that of Pins^{LINKER} alone (Figure 4A; Table 1). Thus, Mud is required for Pins^{TPR} enhancement of the Pins^{LINKER} spindle orientation. To confirm that Mud functions via binding the Pins^{TPR} domain, we engineered a TPR(N226F) mutation based on previous structural studies that showed that this amino acid was likely to contact TPR ligands (Goebl and Yanagida, 1991). The N226F mutation blocked Pins-Mud binding in vitro (Figure 4B) without affecting stability of global TPR protein folding (Figure S2). When we placed the $Pins^{TPR(N226F)+LINKER}$ mutant into our S2 spindle assay, we observed a spindle orientation of $31^{\circ} \pm 24^{\circ}$ (Figure 4C; Table 1), nearly identical to Pins^{LINKER} or Pins^{TPR+LINKER} plus mud RNAi (Table 1). Thus, the Pins^{TPR} must bind Mud to improve Pins^{LINKER}-mediated spindle orientation. However, Pins TPR-Mud interaction is not sufficient for spindle orientation: Pins^{TPR} alone can recruit Mud (Figure 4J), but it has no spindle orientation ability (Figure 2G; Table 1). We conclude that the Pins^{TPR} can recruit Mud, which, together with the Pins^{LINKER} domain, promotes optimal spindle

We next asked whether the Pins^{TPR} and Pins^{LINKER} domains must be together in the same protein to provide full spindle orientation function. Whereas a single Pins^{TPR+LINKER} protein gives excellent spindle orientation (10.7° \pm 7.1°; Table 1), expression of the Pins^{TPR} and Pins^{LINKER} as separate Ed fusion proteins resulted in reduced spindle orientation equal to that of Pins^{LINKER} alone (27.2° \pm 18.1°; Table 1). We conclude that Mud binds the TPR domain within the Pins^{TPR+LINKER} protein and that Pins^{TPR}/Mud acts in cis to stimulate the Pins^{LINKER}/Dlg^{GK}/Khc-73 spindle orientation pathway.

Pins^{TPR+LINKER} Spindle Orientation Requires the Lis1/Dynein Complex

Lis1 and the dynein complex are evolutionarily conserved proteins required for spindle orientation from yeast to mammals (reviewed in Siller and Doe, 2009). The dynein complex is a microtubule minus end-directed motor complex that, when anchored at the plasma membrane, can exert pulling force on spindle pole microtubules (reviewed in Siller and Doe, 2009). Moreover, dynein complex proteins are known to interact with the Mudrelated proteins in *C. elegans* and mammals (Couwenbergs et al., 2007; Merdes et al., 2000; Nguyen-Ngoc et al., 2007). Here, we test whether Lis1 or the dynein complex is required

^a Spindle axis determined by Cnn staining.

^b Ed:GFP:Pins^{LINKER}.

^c Ed:Cherry:Pins^{LINKER}.

^d This is the Insc domain necessary and sufficient for spindle orientation (Knoblich et al., 1999).

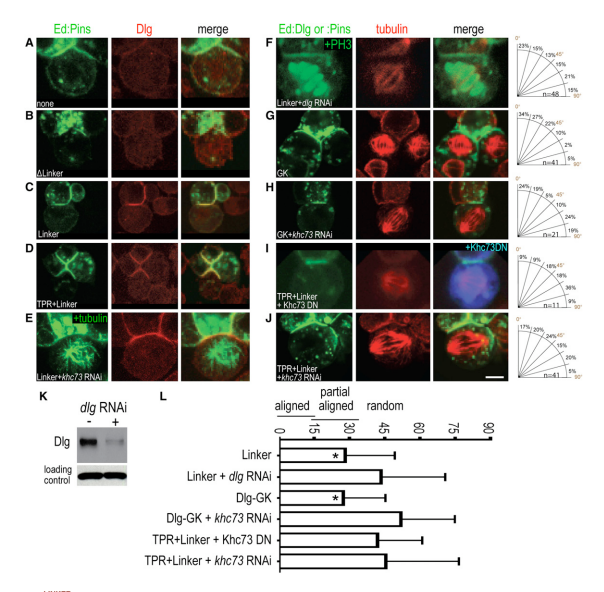


Figure 3. Pins^{LINKER} Requires DIg and Khc73 for Spindle Orientation

S2 cells were transfected with Ed:GFP alone or the indicated Ed:Pins domains (green) and immunostained for endogenous Dlg or Tubulin (red). RNAi knockdown was performed for the indicated genes.

(A–E) Pins^{LINKER} recruits Dlg to the cortex.

(F-L) Pins^{LINKER} requires Dlg and Khc-73 for spindle orientation. The mitotic marker PH3 is shown in (F). Western blot analysis shows RNAi knockdown of Dlg; detection of α-tubulin demonstrates equal loading of lysates (K). The scale bar represents 3 μm for (A)–(J).

(L) Quantification of spindle orientation (mean spindle angle ± standard deviation). *, significantly better spindle orientation compared to Ed alone control (p < 0.01). Other proteins showed no difference from Ed alone control (p > 0.05).

Error bars represent standard deviation from the mean for all experimental values.

for Pins^{TPR+LINKER} or Pins^{LINKER} spindle orientation in S2 cells. We find that dynein light chain (dlc) RNAi reduced endogenous Dlc protein levels (Figure 4F) and decreased PinsTPR+LINKER spindle orientation to levels similar to Pins^{LINKER} alone (Figure 4E; Table 1). Similarly, lis1 RNAi depleted Lis1 protein levels (Figure 4F) and reduced Pins^{TPR+LINKER} spindle orientation to Pins^{LINKER} levels (Figure 4D; Table 1). In contrast, *dlc* RNAi had no effect on Pins^{LINKER} spindle orientation (Table 1), indicating that it acts "downstream" of the PinsTPR domain. We conclude that Pins^{TPR+LINKER} requires Lis1/dynein complex

activity to achieve accurate spindle orientation; in the absence of Lis1 or the dynein complex, only the Pins^{LINKER} pathway appears to be functional.

The Pins^{TPR+LINKER} Induces Rapid, Directional Spindle **Alignment**

We have shown that the Pins^{LINKER} pathway provides partial spindle orientation, while the PinsTPR+LINKER pathway provides full spindle orientation. These differences could be due to many mechanisms that cannot be distinguished by fixed cell

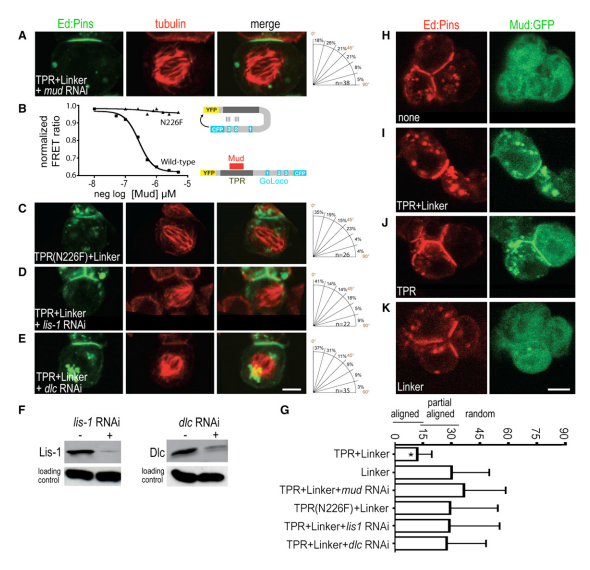
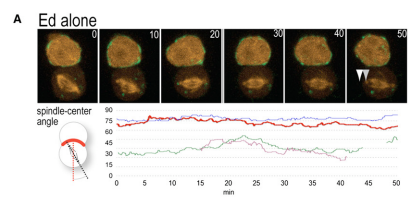


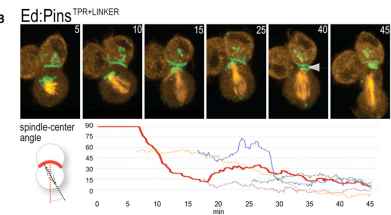
Figure 4. Pins^{TPR+LINKER} Requires Mud and the Dynein/Dynactin Complex to Improve Spindle Orientation Compared to Pins^{LINKER} Alone (A–F) mud RNAi reduces Pins^{TPR+LINKER} spindle orientation to that of Pins^{LINKER} alone (A). The Pins N226F TPR domain mutation abolishes Mud-Pins interaction in a FRET assay (B). The Pins^{TPR(N226F)+LINKER} mutant has spindle orientation equal to that of Pins^{LINKER} alone (C). lis-1 RNAi reduces Pins^{TPR+LINKER} spindle orientation to that of Pins^{LINKER} alone (E). Western blots show RNAi knockdown of Lis-1 and Dlc; detection of α-tubulin demonstrates equal loading of lysates (F). The scale bar represents 3 μm for (A)–(E).

(G) Quantification of spindle orientation (mean spindle angle \pm standard deviation). *, significantly better spindle orientation compared to Pins^{LINKER} (p < 0.01). (H–K) The Pins^{TPR} domain is sufficient to recruit Mud to the cortex. Transfected GFP:Mud (residues 1825–2016, the Pins-binding domain) is recruited to the cortex by Pins^{TPR+LINKER} (I) or Pins^{TPR} (J) but not by Ed alone (H) or Pins^{LINKER} (K). The scale bar represents 3 μm for (H)–(K). Error bars represent standard deviation from the mean for all experimental values.

analysis, including timing of spindle orientation (e.g., the Pins^{LINKER} pathway may only be active for part of mitosis), amplitude of spindle rocking (e.g., Pins^{LINKER} could have larger amplitude), or directional spindle movement (e.g., only the Pins^{TPR+LINKER} may induce movement toward the center of the crescent). To distinguish among these possibilities, we tagged Ed fusion proteins with GFP and the mitotic spindle with Cherry: α -tubulin and performed live imaging of spindle orientation in S2 cells expressing Ed alone, Ed:Pins^{LINKER}, and Ed:Pins^{TPR+LINKER} (Figure 5).

We observed that Ed:GFP control cells had mitotic spindles that drifted relative to the Ed:GFP crescent (Figure 5A); the spindle never showed rapid movement (Figure 5D), and there was typically a gap between spindle poles and the cell cortex (Figure 5A, arrowheads; Figure 5E). This matches spindle movements in untransfected S2 cells (Echard et al., 2004; Goshima et al., 2007). A similar result was observed for Ed:Pins^{TPR} cells (Figures 5D and 5E, and data not shown). In contrast, the Ed: Pins^{TPR+LINKER} cells often showed rapid, directional spindle movement to align the mitotic spindle with the center of the





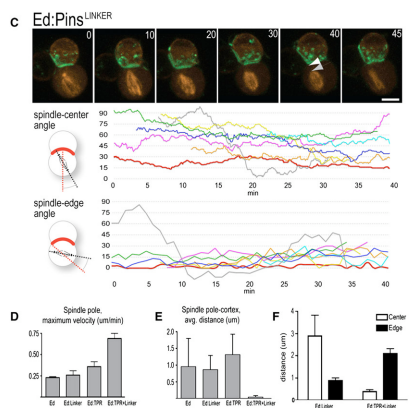
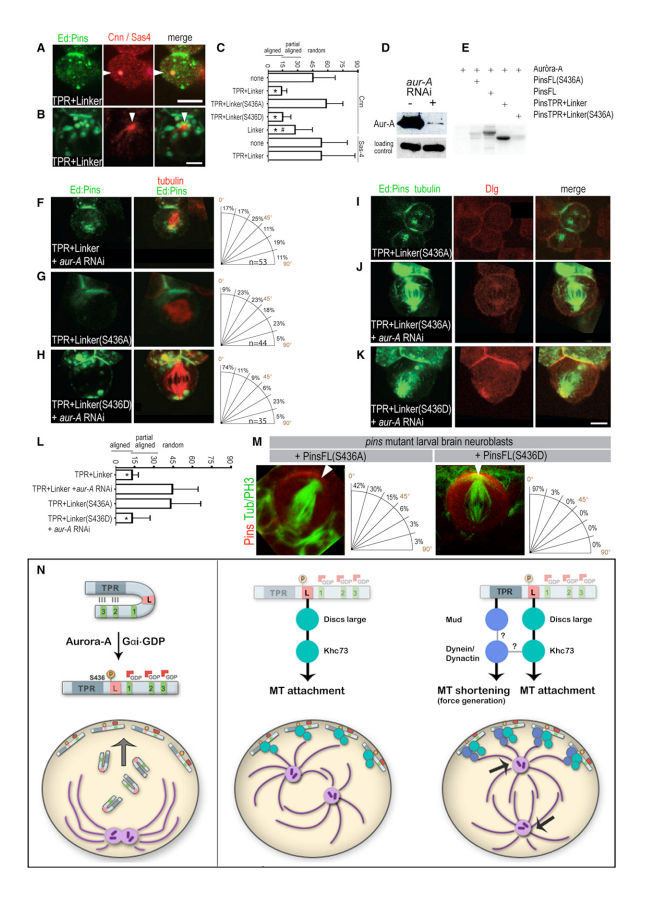


Figure 5. Live Imaging of Spindle Orientation Shows Static Spindles in PinsLINKER Cells but Dynamic Spindle Movement in PinsTPR+LINKER Cells To focus on the mechanism of spindle alignment, we analyzed only Ed:Pins cells that started metaphase with misaligned spindles (>15°).

(A-C) Top: (A)-(C) correspond to Movies S1-S3. Time is in minutes. Arrowheads, spindle pole-to-cortex distance. Each line is a different cell. Red line, cell shown in the still frames and Movies S1-S3. Bottom: quantification of spindle orientation angle to crescent center or crescent edge (y axis) over time (x axis). Ed:GFP control cells have slowly drifting spindles (n = 4) (A). Ed:Pins $^{TPR+LINKER}$ cells have dynamic spindle orientation (n = 5) (B). Ed: Pins^{LINKER} cells have poor spindle orientation to the crescent center (top tracing) but good spindle orientation to the crescent edge (bottom tracings; n = 8) (C). The scale bar represents 2 μm .

- (D) Ed:Pins^{TPR+LINKER} cells have the greatest maximum spindle pole velocity.
- (E) Ed:Pins^{TPR+LINKER} cells have the closest spindle poleto-cortex distance during the 10 min interval prior to anaphase onset.
- (F) Ed:Pins^{LINKER} cells show close spindle pole to edge of the crescent association, whereas $\operatorname{Ed:Pins}^{\operatorname{TPR+LINKER}}$ cells show close spindle pole to center of the crescent association

For panels (D)-(F), error bars indicate standard deviation from the mean for three to five independent live cell division imaging experiments.



Pins crescent (Figures 5B and 5F). Quantification of the maximum velocity of spindle movement confirms that the Ed: Pins^{TPR+LINKER} induces a significantly greater spindle velocity compared to Ed alone, Pins^{LINKER}, or Pins^{TPR} (Figure 5D); similarly, the Ed:Pins^{TPR+LINKER} cells showed a much tighter association with the spindle pole and the cell cortex than Ed alone. Pins^{LINKER}, or Pins^{TPR} (Figure 5E). We conclude that the Pins^{TPR+LINKER} has the ability to induce rapid spindle movement toward the center of the cortical crescent.

Live imaging of the Ed:Pins^{LINKER} cells revealed a phenotype that helped explain its partial spindle orientation phenotype. Ed:Pins^{LINKER} cells showed no dynamic spindle movements (Figure 5D), no movement of the spindle pole toward the cortex (Figure 5E), and preferential alignment with the edge of the Pins crescent (Figures 5C and 5F). We conclude that slow spindle drift brings the initially misaligned spindle into contact with the edge of the crescent, where it becomes stabilized, leading to partial spindle orientation relative to the center of the crescent.

The Pins Spindle Orientation Pathway Is First Active at Prophase

To test for Pins function at interphase, we used the centriolar marker Sas-4, because mature centrosomal markers such as Centrosomin (Cnn) are not present during interphase in S2 cells (Goshima et al., 2007). We found that PinsTPR+LINKER has no effect on centriole positioning during interphase relative to Ed alone control (Figures 6A, 6C, and S3). We next determined whether Pins^{TPR+LINKER} can anchor centrosomes during prophase by measuring centrosome position relative to the center of the Pins crescent (see the Experimental Procedures). We found that the Pins^{TPR+LINKER} showed excellent centrosome alignment at prophase (12° \pm 7°; Figures 6B, 6C, and S3), indistinguishable from the angle of spindle orientation at metaphase (Figures 2K and 2L). Similarly, all other Pins constructs showed prophase centrosome alignment that matched metaphase spindle alignment: for example Ed alone showed no centrosome alignment ($45^{\circ} \pm 22^{\circ}$) and Pins^{LINKER} showed intermediate alignment (28.2° ± 16.2°; Figure S3). Thus, Pins can anchor centrosomes at prophase, and this foreshadows the spindle orientation observed at metaphase. This does not mean that one centrosome always maintains a position near the Pins crescent throughout mitosis, however, as movies show centrosome separation displacing both centrosomes from the Pins^{TPR+LINKER} cortex followed by rapid movement of the bipolar spindle to restore spindle alignment (Figures 5C and 5C', and data not shown). We conclude that the Pins spindle orientation pathway is first activated at the start of prophase.

Aurora-A Activates the Pins Spindle Orientation Pathway by Phosphorylating S436 within the Pins^{LINKER} **Domain**

Lack of Pins function during interphase could be due to immature centrosomes that nucleate few microtubules (Goshima et al., 2007) and/or due to activation of the Pins pathway at the onset of mitosis. The Aurora-A kinase is activated at the start of prophase (Hutterer et al., 2006) and is required for neuroblast spindle orientation (Lee et al., 2006; Wang et al., 2006); here, we test whether it is required to activate the Pins spindle orientation pathway in S2 cells. We used aurora-A RNAi to reduce endogenous Aurora-A protein levels in S2 cells (Figure 6D) and found that this blocked Pins^{TPR+LINKER} from orienting the mitotic spindle (41° ± 24.5°; Figure 6F; Table 1). A second aurora-A RNAi construct gave the same effect (data not shown).

Aurora-A could be acting to promote centrosomal maturation (Berdnik and Knoblich, 2002), or more directly to phosphorylate a member of the Pins spindle orientation pathway. Pins contains three predicted Aurora-A consensus phosphorylation sites (Ferrari et al., 2005), so we performed a deletional analysis to identify which site, if any, warranted further characterization. The Pins^{TPR+LINKER} protein extends to amino acid 466; truncation down to amino acid 446 (Pins 1-446) retained excellent spindle orientation (13.4° \pm 13.3°), whereas truncations down to amino acid 435 (Pins¹⁻⁴³⁵) completely blocked spindle orientation $(41.3^{\circ} \pm 25.1^{\circ})$, as did all shorter Pins constructs (Table 1). We conclude that the Pins amino acids 436-446 are essential for its spindle orientation function. Interestingly, this small domain is evolutionarily conserved from sea urchin to mammals and contains a single predicted Aurora-A phosphorylation site (S436) (Figure S4). We used recombinant Aurora-A kinase to show that this site is a direct target of Aurora-A in vitro and that mutation of this site (S436A) completely blocked Aurora-A phosphorylation of Pins (Figure 6E); additional predicted Aurora-A sites outside this essential domain (S479 and S571) were not phosphorylated (Figure 6E). Thus, Aurora-A specifically phosphorylates S436 within the Pins^{LINKER} domain.

Figure 6. Aurora-A Directly Phosphorylates S436 in the Pins Linker to Recruit the Dlg/Khc-73 Complex and Regulate Spindle Orientation

(A) Interphase S2 cell transfected with Ed:FLAG:Pins^{TPR+LINKER} stained for the centriolar marker Sas-4 showing no centriole anchoring to the crescent.

⁽B) Prophase S2 cell transfected with Ed:FLAG:PinsTPR+LINKER stained for the centrosomal marker Cnn showing centrosome anchoring to the crescent.

⁽C) Quantification of mean centrosome (Cnn) or centriole (Sas4) angle ± standard deviation. *, significant compared to Ed alone control (p < 0.01).

⁽D) Western blot demonstrates partial RNAi knockdown of Aurora-A; alpha-tubulin blot demonstrates equal loading of lysates.

⁽E) Aurora-A phosphorylates Pins on residue 436 in the Linker domain. Recombinant Aurora-A kinase and [γ-32P]-ATP were incubated alone or in the presence of wild-type or S436A versions of full-length or TPR+Linker Pins.

⁽F-K) Aurora-A phosphorylation of Pins S436 is required for DIg recruitment and spindle orientation. The scale bar represents 3 µm for (C)-(H).

⁽L) Quantification of spindle orientation (mean spindle angle ± standard deviation). The number of cells scored for each genotype is 26, 34, 28, 26, 8, 22, and 19 (from top to bottom). *, highly significant versus Ed alone (p < 0.01); #, significantly different from Pins^{TPR+LINKER} (p < 0.05 when assayed by either Cnn or Sas-4). (M) Pins Linker Aurora-A pathway is required in vivo for neuroblast spindle orientation. Larval neuroblasts in a pins62 homozygous mutant expressing the unphosphorylatable PinsFL(S436A) protein show defective spindle orientation (left; note alignment to edge of crescent), whereas the phosphomimetic PinsFL(S436D) protein show spindle pole alignment to the center of the Pins crescent (right). Red, Pins; green, α-tubulin.

⁽N) A model for Pins-mediated spindle orientation. See the main text for details.

Error bars represent standard deviation from the mean for all experimental values.

We next tested the role of S436 phosphorylation in spindle orientation and found that a nonphosphorylatable mutant protein, Pins $^{TPR+LINKER(S436A)}$, failed to orient the mitotic spindle (43° \pm 20.8°; Figure 6G), similar to the effect of aurora-A RNAi (41° \pm 24.5°; Figure 6F). In contrast, the phosphomimetic S436D mutation resulted in excellent spindle orientation (13.7° \pm 11.3°), similar to Pins $^{TPR+LINKER}$ (Table 1). Importantly, the phosphomimetic Pins $^{TPR+LINKER(S436D)}$ protein was able to orient the mitotic spindle even after nearly complete RNAi depletion of Aurora-A (Figures 6D and 6H), showing that the role of Aurora-A in spindle orientation is to phosphorylate Pins, rather than to promote centrosomal maturation.

How does S436 phosphorylation activate the Pins spindle orientation pathway? The similarity of *aurora-A*, *dlg*, and *khc-73* RNAi phenotypes led us to test whether S436 phosphorylation was required to recruit Dlg to the Pins^{TPR+LINKER} cortical crescent. Indeed, the nonphosphorylatable Pins^{TPR+LINKER(S436A)} failed to recruit endogenous Dlg (Figure 6I). RNAi depletion of Aurora-A also blocked recruitment of Dlg to the wild-type Pins cortical domain (Figure 6J), but it did not have any effect on recruitment induced by the phosphomimetic Pins^{TPR+LINKER(S436D)} protein (Figure 6K). We conclude that Aurora-A phosphorylates the Pins^{LINKER} at S436, triggering recruitment of Dlg protein and the activation of the Pins^{LINKER}/Dlg/Khc-73 spindle orientation pathway at the onset of mitosis.

The Pins^{LINKER} Pathway Is Required for Spindle Orientation in Larval Neuroblasts

We next tested whether the Pins LINKER/Aurora-A pathway is active in larval neuroblasts in vivo. We generated transgenic flies that allow inducible expression of full-length Pins with either phosphorylation blocking or mimicking mutations—Pins $^{FL(S436A)}$ and Pins $^{FL(S436D)}$ —in a $pins^{62}$ mutant background. We find that the Pins $^{FL(S436D)}$ phosphomimetic protein provides full spindle orientation function in larval brain neuroblasts lacking endogenous Pins (8.2° \pm 4.0°, n = 32; Figure 6M). In contrast, the non-phosphorylatable Pins $^{FL(S436A)}$ protein shows spindle orientation defects in larval brain neuroblasts (23.2° \pm 17.8°, n = 30; Figure 6M) that closely match the $pins^{62}$ mutant alone (23.2° \pm 23.1°, n = 12) (Siegrist and Doe, 2005). Thus, the single amino acid substitution in Pins $^{FL(S436A)}$ blocks most or all Pins-mediated spindle orientation function in larval brain neuroblasts, replicating and verifying our findings in the S2 assay.

We also assessed whether Pins^{FL(S436A)} protein has a dominant-negative function. Indeed, expression of Pins^{FL(S436A)} in a wild-type background resulted in modest defects in spindle orientation (17.3° \pm 13.6°, n = 56), compared to wild-type controls (6.2° \pm 3.4°, n = 48). In contrast, expression of the Pins^{FL(S436D)} phosphomimetic protein did not significantly affect spindle orientation (6.3° \pm 3.5°, n = 13). We conclude that the Pins^{LINKER}/Aurora-A spindle orientation mechanism is active in vivo and required for proper spindle orientation in larval neuroblasts.

DISCUSSION

We have developed an "induced cell polarity" system using the Drosophila S2 cell line that permits rapid testing of individual proteins, domains, and amino acids for their role in cell polarity and spindle orientation. We use this system to dissect the role of the evolutionarily conserved Pins protein in spindle orientation. We identify a previously unknown Pins^{LINKER} pathway, show that it functions in larval neuroblasts, and show that the Pins^{TPR} acts in *cis* to the Pins^{LINKER} to improve spindle orientation. A model summarizing our results is shown in Figure 6N.

The Pins^{LINKER}/Dlg/Khc-73 Pathway

A surprising result of our studies is the importance of the Pins^{LINKER} domain for spindle orientation in the S2 assay and within neuroblasts in vivo. Only this domain is sufficient for spindle orientation, and a single point mutation in the linker domain (S436A) results in spindle orientation defects in larval neuroblasts that closely mimic the pins null mutant phenotype. On the basis of domain mapping and epistasis analysis, we have identified a linear pathway from cortical PinsLINKER to the plus ends of astral microtubules: (1) Aurora-A phosphorylates Pins^{LINKER} on a single amino acid, serine 436, (2) the phosphorylated Pins^{LINKER} binds and recruits Dlg, (3) the kinesin Khc-73 moves to astral microtubule plus ends using its motor domain and may be anchored at the plus ends by its Cap-Gly domain (Siegrist and Doe, 2005), and (4) the Khc-73^{MBS} domain binds the cortical Dlg^{GK} domain, thereby linking Khc-73⁺ astral microtubule plus ends to the Dlg cortical domain (Figure 6N). Interestingly, this pathway is active in both directions during mitosis. Cortical Pins acts through Dlg and Khc-73 to regulate spindle orientation (this work), and spindle-associated Khc-73 acts through Dlg and Pins to induce Pins/Gai functional cortical polarity in neuroblasts (Siegrist and Doe, 2005).

Why does the Pins^{LINKER} pathway provide only partial spindle orientation function? Live imaging rules out several possible explanations, such as Pins^{LINKER}-induced spindle rocking variability, or that Pins^{LINKER} only functions during a narrow window during mitosis. Live imaging shows that in Pins^{LINKER} cells, the spindle drifts until it is immobilized at the edge of the crescent (Figures 5B and 5D). This is consistent with the fact that Khc-73 is a plus end-directed motor protein, and thus unable to generate pulling forces to bring the centrosome closer to the cell cortex; at best, it would provide a static link between astral microtubules and the cell cortex.

The Pins^{TPR}/Mud/Dynein Pathway

The Pins^{TPR} domain can improve the Pins^{LINKER} spindle orientation to a level matching wild-type neuroblasts. We propose that the Pins^{TPR} domain directly binds Mud and that Mud interacts with the dynein/dynactin/Lis1 complex to enhance Pins^{LINKER} spindle orientation. This model is based on five observations. First, the Pins^{TPR} domain binds Mud in vitro and the two proteins coimmunoprecipitate from in vivo lysates (Bowman et al., 2006; Izumi et al., 2006; Siller et al., 2006) (Figure 4B); this interaction is conserved in the related *C. elegans* and mammalian proteins (Couwenbergs et al., 2007; Merdes et al., 2000; Nguyen-Ngoc et al., 2007). Second, the Pins^{TPR} and Pins^{TPR+LINKER} but not the Pins^{LINKER} can recruit Mud to the cortex of S2 cells. Third, Pins^{TPR+LINKER}-mediated spindle orientation requires the dynein complex proteins Dlc and Lis1. Fourth, Pins^{TPR+LINKER}-mediated spindle orientation spindle

movement toward the center of the Pins cortical crescent, similar to dynein-dependent spindle orientation in Drosophila neuroblasts (Siller and Doe, 2008). Fifth, PinsTPR+LINKER-mediated spindle orientation leads to dynein-dependent movement of the spindle pole close to the cell cortex, consistent with dynein minus end-directed pulling of astral microtubules, as observed in other cell types (Siller and Doe, 2009).

If Pins^{TPR} recruits Mud, and Mud recruits the dynein complex, then why doesn't Pins^{TPR} have spindle-orienting function on its own? The simplest model is that PinsTPR/Mud alone is unable to recruit or activate the dynein complex. Alternatively, the Pins^{LINKER} pathway could be required for "presenting" microtubule plus ends to an active Pins^{TPR}/Mud/Dynein complex, which fits with the requirement for Pins^{TPR} and Pins^{LINKER} acting in *cis*. In summary, our data show that the Pins^{TPR} and Pins^{LINKER} domains provide distinct functions, both of which are required for optimal spindle orientation (Figure 6N). Interestingly, spindle orientation in S2 cells does not show "telophase rescue"a phenomenon whereby spindles that are partially oriented in metaphase/anaphase neuroblasts become aligned with the cell polarity axis by telophase (Siller and Doe, 2008) - consistent with the absence of redundant spindle orientation pathways in this assay.

Regulation of Pins Spindle Orientation Pathways

The Pins^{TPR} pathway is regulated by Gαi binding to the GoLoco domain, relieving intramolecular TPR-GoLoco interactions, and making the TPR domain accessible for intermolecular interactions (Du and Macara, 2004; Nipper et al., 2007). In addition, Gai is required to recruit Pins to the cell cortex, where it can interact with regulator and effector proteins (Yu et al., 2003). In our S2 spindle orientation assay, we can bypass a requirement for Gai by simply deleting the GoLoco domain (thereby freeing the TPR for intermolecular interactions) and tethering the Pins^{TPR+LINKER} protein to the cortex by fusion with the Ed transmembrane protein. Thus, Gai is important to activate and localize full-length Pins, but not as an effector of Pins-mediated spindle orientation.

In contrast, the Pins LINKER pathway is not regulated by $G\alpha i,$ because full-length Pins in the absence of Gai provides equal spindle orientation to Pins^{LINKER}, suggesting that the Pins^{LINKER} is active when Pins is in the "closed" form. The Khc-73 mammalian ortholog GAKIN transports hDlg to the cell cortex (Hanada et al., 2000), but there are several reasons to think that this mechanism does not activate the Pins LINKER pathway. First, cortically tethered Dlg^{GK} domain requires Khc-73 for spindle orientation, which rules out a role for Khc-73 in merely transporting Dlg to the cortex; second, khc-73 RNAi does not block the ability of Pins^{LINKER} to recruit Dlg to the cortex.

We have shown that Aurora-A kinase activates the Pins^{LINKER} spindle orientation pathway by phosphorylating S436 in the linker domain and that this pathway is required for accurate spindle orientation in vivo for larval neuroblast asymmetric cell division. Neuroblasts expressing the nonphosphorylatable form of Pins (S436A) have a weaker spindle orientation phenotype than aurora-A null mutants (Lee et al., 2006; Wang et al., 2006), as expected because of Aurora-A regulation of multiple Pinsindependent processes required for spindle orientation, such as centrosome maturation, cell-cycle progression, and cell polarity in flies (Barr and Gergely, 2007; Berdnik and Knoblich, 2002; Hutterer et al., 2006; Lee et al., 2006; Wang et al., 2006; Wirtz-Peitz et al., 2008). However, we show that a Pins phosphomimetic mutant (S436D) allows spindle orientation even after RNAi depletion of Aurora-A levels, suggesting that Aurora-A phosphorylation of PinsS436 is essential for Pins-dependent spindle orientation in the S2 cell assay. Furthermore, our finding that the PinsS436A protein has no spindle orientation activity in pins mutant larval neuroblasts, and has dominant-negative activity in the presence of endogenous Pins, shows that the Aurora-A/Pins^{LINKER} pathway is required for spindle orientation in larval neuroblasts as well.

The Pins spindle orientation pathway is cell-cycle regulated: interphase S2 cells that have polarized Pins TPR+LINKER do not capture centriole/centrosomal microtubules. There are at least two reasons for the lack of Pins interphase activity. First, the level of the Aurora-A kinase is low during interphase, and we have shown that Aurora-A phosphorylation of Pins S436 is essential for Pins-mediated spindle orientation. Second, interphase centrosomes are immature, lacking Cnn and nucleating few microtubules (Goshima et al., 2007). Expression of the Pins S436D protein, which is fully functional during mitosis even after Aurora-A depletion, still has no ability to capture centrioles during interphase. Thus, both centrosome maturation and Aurora-A activation are required for Pins-mediated spindle orientation in S2 cells.

Concluding Remarks

Here, we have induced cell polarity and spindle orientation in a cultured cell line. We used this system to identify two pathways regulating spindle orientation, to establish molecular epistasis within each pathway, and to identify the target of the mitotic kinase Aurora-A that coordinates cell-cycle progression with spindle orientation. In the future, this system should be useful for characterizing spindle orientation pathways from other Drosophila cell types or from other organisms, identifying the mechanisms that control centrosome or spindle asymmetry, and characterizing the establishment and maintenance of cortical polarity. In each of these cases, our induced polarity system should be useful for rapid protein structure/function studies and high-throughput drug or RNAi loss-of-function studies.

EXPERIMENTAL PROCEDURES

Plasmid Construction, Cell Culture, and RNAi

Ed:FLAG, Ed:GFP, and Ed:Cherry constructs were made in pMT-V5 (Invitrogen, Carlsbad, CA), replacing the Ed cytoplasmic domain with the visualization tag and the protein of interest at the C terminus (e.g., Ed:FLAG:Pins). Proteins with different visualization tags conferred similar spindle orientation (Table 1). Standard methods were used to grow S2 cells (Goshima et al., 2007). Cells were seeded at $\sim 1-3 \times 10^6$ cells per well and transfected with 0.4-1 ug total DNA with Effectene (QIAGEN, Germantown, MD), and gene expression was induced by 500 μ M CuSO₄ for 24–48 hr. Cell clustering was induced by rotation at \sim 175 RPM for 1–3 hr.

For RNAi treatment, primers that amplify $\sim\!\!400\text{--}600$ base pairs were designed at http://www.dkfz.de/signaling2/e-rnai/ with T7 promoter tags. PCR-amplified sequences were reverse transcribed with the Megascript T7 kit (Ambion, Austin, TX). S2 cells were seeded at 1×10^6 cells per well in 1 ml of serum-free Schneider media and incubated with 100 μ l (\sim 10 μ g). After 1 hr, 2 ml of serum-containing media was added and cells were incubated for an additional 3 days.

Live Imaging, Immunostaining, and Western Blots

For live imaging, cells were plated on poly-L-lysine-coated chambered coverslips (Nalge). Three focal planes spaced 2 μm apart were collected every 5 s with Volocity software (Improvision) on a McBain spinning disc microscope with a 60× 1.4 NA lens and a Hamamatsu EM-CCD camera. Volocity was used to XY crop, Z merge with maximum pixel intensity, and export as a TIFF series; ImageJ was used to generate Quicktime movies.

For immunostaining, S2 cells were fixed for 20 min in 4% paraformaldehyde, stained (Goshima et al., 2007), and imaged on a Leica SP2 confocal microscope with a 60×1.4 NA lens. For in vivo imaging of larval neuroblasts, we used neuroblast-specific *worniu-gal4* to express $UAS-pins^{FL(S436A)}$ or $-pins^{FL(S436D)}$ in a $pins^{62}$ homozygous mutant background (Nipper et al., 2007). For western blots, $20~\mu g$ of S2 cell extracts were used per lane. The localization of endogenous polarity proteins in S2 cells is shown in Figure S1. Antibodies and dilutions were as follows: rat Lis1, 1:1000 (Siller et al., 2005), rabbit $G\alpha_i$, 1:1000 (Nipper et al., 2007), mouse Dlg, 1:250 (Developmental Studies Hybridoma Bank, lowa); rabbit Sas-4, 1:500 (Basto et al., 2006); rabbit Aurora-A, 1:200 (Berdnik and Knoblich, 2002); rabbit centrosomin, 1:1000 (Megraw et al., 1999); rat Pins, 1:500 (Yu et al., 2000); mouse FLAG, 1:500 (Sigma); rat tubulin, 1:1000 (Abcam); mouse tubulin, 1:1000 (Covance)

Measuring Cortical Polarity, Spindle Orientation, and Centrosome Alignment

Pixel intensity was measured in ImageJ. Ed crescents over 90° cell diameter were excluded. Spindle angles were measured with a vector perpendicular to the center of the Ed crescent and a vector matching the spindle or connecting the spindle poles. To estimate the spindle angle measurement error, we compared the mean angle from multiple independent experiments of Ed-Pins^{LINKER} and Ed-Pins^{FL} + Gαi (three trials each). On the basis of this analysis, and assuming that the sources of error were consistent, we estimated a standard error of $\pm 3^\circ$ in spindle angle measurement in our experiments.

In Vitro Kinase Assays

Recombinant Aurora-A kinase was purchased from Millipore (Billerica, MA). Pins constructs (10 μ g) and Aurora-A (100 ng) were diluted in ice-cold assay buffer (20 mM Tris [pH 7.4], 100 mM NaCl, 1 mM DTT, 10 mM MgCl₂, and 10 μ M ATP). ATP- γ - 32 P (5 μ Ci) was added at 30°C for 30 min. Reactions were quenched by addition of SDS loading buffer. Samples were resolved by SDS-PAGE and analyzed with a Storm 860 and Image Quant 5.1 (Molecular Dynamics).

FRET Assays

Sensor proteins were diluted to 200 nM in 20 mM Tris (pH 7.5), 100 mM NaCl, and 1 mM DTT with or without MudPBD and excited at 433 nm, and the YFP (525 nm) to CFP (475 nm) emission ratios were measured.

SUPPLEMENTAL DATA

Supplemental Data include Supplemental Experimental Procedures, one table, four figures, and three movies and can be found with this article online at http://www.cell.com/supplemental/S0092-8674(09)00978-7.

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