Introduction

The unequal distribution of cell fate determinants during asymmetric cell division is a fundamental process that underlies the generation of cell diversity in a variety of multicellular organisms (Yamashita et al., 2007). The positioning of the mitotic spindle relative to the cell polarity axis is critical to mediate asymmetric cell divisions (Siller and Doe, 2009).

Several mechanisms ensure correct spindle alignment in the asymmetrically dividing unicellular organism, budding yeast *Saccharomyces cerevisiae*. First, spindle positioning relative to the intrinsic polarity axis of the cell is ensured by two partially redundant pathways; one reliant upon the function of the conserved protein Kar9, the other upon the microtubule motor protein dynein (Segal and Bloom, 2001). Second, if spindle orientation fails, a surveillance mechanism called the spindle positioning checkpoint (SPOC) pauses cell cycle progression until proper spindle alignment is achieved (Fraschini et al., 2008; Caydasi et al., 2010a).

Under normal cell cycle progression, the mitotic exit network (MEN) initiates mitotic exit and cytokinesis in late anaphase. The MEN is a GTPase-driven signaling pathway whose components associate with the yeast microtubule organizing center, the spindle pole body (SPB). MEN signaling starts with the activation of the GTPase Tem1 that triggers the sequential activation of the protein kinase Cdc15 and the Dbf2–Mob1 kinase complex. Dbf2–Mob1 promotes the full activation of the protein phosphatase Cdc14, which is needed to promote Cdk inactivation, leading to mitotic exit and cytokinesis (Bardin and Amon, 2001). Feedback mechanisms involving Cdc14 and Cdk contribute to MEN activation at the level of Cdc15 and Dbf2–Mob1 (Jaspersen and Morgan, 2000; König et al., 2010). However, the mechanisms by which the activation of Tem1 is controlled remain debatable. Tem1 preferentially associates with the SPB that enters the daughter cell (dSPB; Bardin et al., 2000; Pereira et al., 2000). Based on genetic studies and the presence of the guanine nucleotide exchange factor (GEF) homology domain, the putative GEF Lte1 has been implicated in Tem1 activation (Keng et al., 1994; Shirayama et al., 1994b). Lte1 is confined at the daughter cell cortex and the entrance of the dSPB bound to Tem1 brings Tem1 in close proximity to Lte1, thus promoting Tem1 activation (Bardin et al., 2000; Pereira et al., 2000). However, Lte1 is not essential for mitotic exit at temperatures above 30°C and no GEF activity toward Tem1 has been observed.

The spindle position checkpoint (SPOC) is an essential surveillance mechanism that allows mitotic exit only when the spindle is correctly oriented along the cell axis. Key SPOC components are the kinase Kin4 and the Bub2–Bfa1 GAP complex that inhibit the mitotic exit-promoting GTPase Tem1. During an unperturbed cell cycle, Kin4 associates with the mother spindle pole body (mSPB), whereas Bub2–Bfa1 is at the daughter SPB (dSPB). When the spindle is mispositioned, Bub2–Bfa1 and Kin4 bind to both SPBs, which enables Kin4 to phosphorylate Bfa1 and thereby block mitotic exit. Here, we show that the daughter cell protein Lte1 physically interacts with Kin4 and inhibits Kin4 kinase activity. Specifically, Lte1 binds to catalytically active Kin4 and promotes Kin4 hyperphosphorylation, which restricts Kin4 binding to the mSPB. This Lte1-mediated exclusion of Kin4 from the dSPB is essential for proper mitotic exit of cells with a correctly aligned spindle. Therefore, Lte1 promotes mitotic exit by inhibiting Kin4 activity at the dSPB.

The cortical protein Lte1 promotes mitotic exit by inhibiting the spindle position checkpoint kinase Kin4

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detected with purified proteins (Adames et al., 2001; Geymonat et al., 2009a).

During an unperturbed cell cycle the bipartite GTPase-activating protein (GAP) complex, Bub2–Bfa1, maintains Tem1 in an inactive state until Bfa1 is inactivated through phosphorylation by the polo-like kinase Cdc5. In late mitosis, this phosphorylation decreases Bub2–Bfa1 GAP activity, promoting mitotic exit (Hu et al., 2001; Geymonat et al., 2003). If the spindle becomes misaligned, the SPOC kinase Kin4 phosphorylates Bfa1, blocking the activity of Cdc5 to inactivate Bfa1 (D’Aquino et al., 2005; Pereira and Schiebel, 2005; Maekawa et al., 2007; Caydasi and Pereira, 2009).

Both Kin4’s localization and kinase activity are important for its control of Bub2–Bfa1 function. The catalytic activity of Kin4 is regulated by the bud neck–associated kinase Elm1. Elm1 phosphorylates a conserved threonine residue in the conserved activation loop (T-loop) of Kin4 (Caydasi et al., 2010b; Moore et al., 2010). However, activation of Kin4 by Elm1 is not sufficient to provide SPOC function. Another prerequisite is Kin4 localization to the mother cell cortex and SBPs. Kin4 SBP3 and cortex association are regulated by the activity of the phosphatase PP2A subunit Rts1 via an unknown mechanism (Chan and Amon, 2009; Caydasi et al., 2010b).

Whereas deletion of KIN4 has only minor consequences upon mitotic progression under normal growth conditions, excessive production of Kin4 transcripts from artificial promoters blocks cell cycle progression in late anaphase in a Bub2–Bfa1-dependent manner (D’Aquino et al., 2005). Similarly, placing a mutated Kin4 variant within daughter cells also causes mitotic exit delays (Chan and Amon, 2010). Thus, it is tempting to speculate that Kin4 kinase activity must be kept high inside the mother cell to promote Kin4’s function if the spindle is misoriented; on the other hand, Kin4 kinase activity must be kept low within the daughter cell to allow mitotic exit. The inhibitory mechanisms that restrain Kin4 kinase activity locally are unknown. Here, we established that Lte1 physically interacts with the catalytically active form of Kin4. In vivo studies showed that Lte1 acts as an inhibitor of Kin4 catalytic activity toward Bfa1. Furthermore, we established that Lte1 is necessary to promote Kin4 hyperphosphorylation and exclusion from the dSPB during anaphase. We therefore propose that Lte1 promotes mitotic exit by inhibiting the activity and dSPB localization of the MEN inhibitor Kin4.

**Results**

**Kin4 and Lte1 physically interact in vivo and in vitro**

To identify Kin4-interacting proteins, we purified Kin4 from yeast cell lysates using the tandem affinity purification (TAP) strategy (Puig et al., 2001). Mass spectrometric (MS) analysis of the composition of the Kin4–TAP complex identified the known Kin4 interactor, Bfa1, and components of the SPB (Fig. 1 A and Fig. S1 A; Pereira and Schiebel, 2005). In addition, we identified a large number of peptides corresponding to Lte1 in the Kin4–TAP complex (Fig. S1 A). Likewise, when we purified Lte1–TAP complexes we identified Kin4 alongside the known Lte1-interacting proteins Kel1, Kel2, Ras1, and Ras2 (Fig. 1 A and Fig. S1 A; Höfken and Schiebel, 2002; Yoshida et al., 2003). Co-purification of Lte1 and Kin4 was unexpected because Kin4 preferentially localizes at the mother cell cortex, whereas Lte1 is mostly associated with the cortex of the bud (Bardin et al., 2000; Pereira et al., 2000; D’Aquino et al., 2005; Pereira and Schiebel, 2005).

To confirm the physical association between Kin4 and Lte1, we performed immunoprecipitation experiments using functional hemagglutinin (HA) and Myc-tagged fusion proteins. Kin4-9Myc coprecipitated with Lte1-6HA in HA specific pull-downs (Fig. 1 B) and, vice-versa (Fig. S1 B). We considered the possibility that the interaction between Kin4 and Lte1 arose from copurification of large subfragments of the cell cortex. However, this was not the case, as neither Kin4-6HA nor Lte1-6HA coprecipitated with a plasma membrane protein of the daughter cell, Ist2-3Myc (Fig. S1 C) (Takizawa et al., 2000), nor did Lte1-6HA coprecipitate the mother cortex–associated protein, Sfk1-9Myc (Fig. S1 D) (Audhya and Emr, 2002). We thus conclude that Lte1 and Kin4 are companions found within common complexes.

Additionally, Kel1 peptides were also found in the Kin4 purification and a fraction of Kin4-6HA coimmunoprecipitated with Kel1-9Myc (Fig. 1 C). This interaction was specific for Kel1, as Kin4 did not coimmunoprecipitate with the closely related molecule Kel2 or the Lte1 interactors Ras1 and Ras2 (Fig. S1, A and E; and unpublished data). To better characterize Lte1–Kin4–Kel1 interactions, we asked whether the absence of any one component would influence the association between the others. The association of Lte1 with Kin4 was unaffected by loss of Kel1 (Fig. 1 D), whereas deletion of LTE1 impaired binding of Kel1 to Kin4 (Fig. 1 E). This suggests that Lte1 bridges the interaction between Kel1 and Kin4.

We next asked whether Lte1 directly interacts with Kin4. Here, we used purified, recombinant Lte1 and Kin4 proteins. In contrast to full-length Lte1, N- and C-terminal truncated Lte1 constructs were soluble and could be purified from *Escherichia coli* cells (Fig. 1, F and G). Both 6His-Lte1-N and 6His-Lte1-C proteins bound GST-Kin4 but neither bound GST alone (Fig. 1, F and G; lanes 8–11). We therefore concluded that Lte1 and Kin4 are present in common complexes in vivo and physically associate in vitro.

**Misplacement of Lte1 but not Kel1 to the mother cell cortex promotes mitotic exit of cells with misaligned spindle**

We next asked how KEL1 and LTE1 are functionally linked to KIN4. Overexpression of KIN4 is toxic (D’Aquino et al., 2005). We found that co-overexpression of LTE1, but not of KEL1, rescued the toxicity of Kin4 overproduction (Fig. 2 A), suggesting a functional interplay between Lte1 and Kin4 but not Kel1 and Kin4 in vivo.

Overexpression of LTE1, expression of a mutant variant of LTE1 that accumulates in the mother cell cortex, or disruption of the mother–daughter cell diffusion barrier (defective bud neck) causes SPOC deficiency in cells with misaligned spindles (Bardin et al., 2000; Castillon et al., 2003; Geymonat et al., 2009a). To test whether misplacement of Kel1 would impair SPOC function in a similar manner to Lte1, we established a strategy that
Fusion of Sfk1 with GBP altered neither cell cycle progression nor SPOC proficiency of kar9Δ cells (Fig. 2 B; unpublished data). Lte1-GFP and Kel1-GFP were efficiently enriched in the mother cell cortex in strains carrying the mother cell cortex protein Sfk1 tagged with GBP (Sfk1-GBP; Fig. 2, B–E).

Figure 1. Lte1 interacts with Kin4 in vitro and in vivo. (A) Kin4 and Lte1-interacting partners found by MS analysis. (B–E) Kin4 interacts with Lte1 and Kel1. Immunoprecipitations using anti-HA or anti-Myc beads as indicated. (F and G) In vitro binding assay using bacterially expressed KIN4 and LTE1. (F) Lte1 truncations used in G; numbers represent amino acid positions. (G) Immunoblotting of bacterially expressed GST (lane 1) and GST-Kin4 (lane 2) bound to glutathione-Sepharose beads. Protein extracts of E. coli expressing 6His (lane 3), 6His-Lte1-N (lane 4), and 6His-Lte1-C (lane 5) were incubated with GST (lanes 6, 8, and 10) or GST-Kin4 (lanes 7, 9, and 11) for 2 h at 4°C.
the late anaphase arrest induced by SPOC, which requires both Tem1 inhibition and Kin4 function (D’Aquino et al., 2005; Pereira and Schiebel, 2005), nocodazole-induced metaphase arrest does not require Kin4; however, it can be bypassed by Tem1 activation (Chan and Amon, 2009). Thus, it is most likely that mother cell–enriched Lte1 promotes mitotic exit of cells with misaligned spin-
dles by inhibiting Kin4 rather than activating Tem1.

form a complex with Kin4 (Fig. S1), had no inhibitory effect on SPOC in SFK1-GBP kar9Δ cells (Fig. 2, B and E). Thus Lte1, but neither Kel1 nor Kel2, bypassed the SPOC once improperly located in the mother cell compartment.

Moreover, mother cortex–located Lte1 did not impair the metaphase arrest induced by treatment of cells with the micro-
tubule-depolymerizing drug, nocodazole (Fig. 2 F). In contrast to the late anaphase arrest induced by SPOC, which requires both Tem1 inhibition and Kin4 function (D’Aquino et al., 2005; Pereira and Schiebel, 2005), nocodazole-induced metaphase arrest does not require Kin4; however, it can be bypassed by Tem1 activation (Chan and Amon, 2009). Thus, it is most likely that mother cell–enriched Lte1 promotes mitotic exit of cells with misaligned spin-
dles by inhibiting Kin4 rather than activating Tem1.
Interaction between Kin4 and Lte1 requires active Kin4

To investigate the requirements for the Kin4–Lte1 interaction in vivo, we asked whether the binding of Kin4 to Lte1 was restricted to a particular phase in the cell cycle. We found Kin4 and Lte1 interacting in all phases of the cell cycle and in cells where the SPOC was active (Fig. S2, A and B). This may indicate that a fraction of Lte1 and Kin4 interacts throughout the cell cycle, although the majority of both proteins were kept in different compartments.

Using our GBP strategy, we found that bringing Lte1 to the mother cell cortex led to a 30–40% increase in the amount of Lte1 interacting with Kin4 in immunoprecipitations (Fig. S2 C), supporting the idea that Kin4–Lte1 complexes might be facilitated if Lte1 and Kin4 were present in the same subcellular compartment. We therefore pursued the idea that Kin4 and Lte1 may, to some extent, colocalize in the cytoplasm during the entire cell cycle. Indeed, inspection of live cells coexpressing functional KIN4-GFP LTE1-3Cherry (Maeder et al., 2007) revealed that although Lte1 and Kin4 accumulated most strongly in the daughter and mother cell bodies, respectively, they were not entirely excluded from the opposite cellular compartments (Fig. 3 A). Interestingly, just after cytokinesis, Kin4 remained associated with the bud neck and the new bud, in which Lte1 accumulated (Fig. 3 A). At the time that Kin4 accumulated at the bud neck, Lte1 was more evenly dispersed (Fig. 3 A). These data suggested that a fraction of Lte1 and Kin4 may indeed associate with one another throughout the cell cycle.

Importantly, Lte1 was unable to bind to the kinase-dead Kin4-T209A and to two other catalytic inactive Kin4 mutants (Fig. 3 B and Fig. S2 D). The interaction between Kin4 and Lte1 was also drastically reduced in eln1Δ cells, in which Kin4 is catalytically inactive (Caydasi et al., 2010b; Moore et al., 2010). The absence of Lte1–Kin4 complexes was not directly related to SPOC deficiency, as Lte1 still associated with Kin4 in rts1Δ cells, which are SPOC deficient with Kin4 catalytic activity similar to wild-type cells (Fig. 3 C; Chan and Amon, 2009; Caydasi et al., 2010b). Furthermore, overexpression of LTE1-3Cherry, using the inducible Gal1 promoter, caused a strong accumulation of Kin4-GFP but not Kin4-T209A-GFP (both expressed endogenously) in the bud (Fig. 3 D), further supporting the notion that the catalytic activity of Kin4 is required for its association with Lte1.

Lte1 inhibits the kinase activity of Kin4 in vitro

To understand the functional relevance of the Kin4–Lte1 interaction described here, we investigated whether Kin4 influenced the phosphorylation or localization of Lte1. When we compared KIN4 wild-type cells with kin4Δ cells, we failed to detect any differences in either the phosphorylation or localization profiles of Lte1 with or without SPOC activation (unpublished data). Furthermore, purified Kin4 was not able to phosphorylate recombinant Lte1 in vitro (unpublished data). We therefore consider it unlikely that Kin4 phosphorylates Lte1 or influences its localization.

The fact that Lte1 physically associates with Kin4 prompted us to ask whether Lte1 regulates Kin4 catalytic activity. For this, Kin4 in vitro kinase activity toward Bfa1 was measured in the presence of increasing levels of purified 6His-Lte1-N or 6His-Lte1-C (Fig. 4). 6His-GFP was used as a control to demonstrate that addition of buffer and an unrelated protein did not influence Kin4 kinase activity (Fig. 4 A, lanes 1 and 2). Purified GST-Kin4 but not the kinase-dead variant (GST-Kin4-T209A), enriched from yeast cell lysates, phosphorylated Bfa1 in vitro (Fig. 4 A, lanes 3 and 4). 6His-Lte1-N significantly inhibited the phosphorylation of Bfa1 by Kin4 in a dose-dependent manner (Fig. 4 B). An inhibition of ～64% in Kin4 specific activity was reached at the highest concentration of 6His-Lte1-N tested (Fig. 4 B). 6His-Lte1-C also inhibited Kin4 kinase activity, although not as efficiently as 6His-Lte1-N (Fig. 4 C; 34% inhibition). Thus, Lte1 inhibits Kin4 catalytic activity in vitro.

Lte1 is an inhibitor of Kin4 function in vivo

Kin4 is catalytically active during every cell cycle even when SPOC activation is not triggered (D’Aquino et al., 2005; Caydasi et al., 2010b). The fact that co-overexpression of LTE1 and KIN4 was not lethal for cell growth (Fig. 2 A), in contrast to KIN4 overexpression alone (Fig. 2 A), supports the notion that cells must maintain a balance between Kin4 and Lte1 activities to achieve normal cell cycle progression. If true, one would expect cells lacking LTE1 to be more sensitive to increasing amounts of Kin4 compared with wild-type cells. Indeed, although wild-type cells were able to grow with additional gene copies of KIN4 supplied by either a centromeric (CEN-KIN4) or a 2µ-based plasmid (2µ-KIN4; Fig. 5 A), the addition of extra copies of KIN4, even from a centromeric plasmid, drastically impaired growth of lte1Δ cells (Fig. 5 A). Deletion of BFA1 rescued this lethality (Fig. 5 A), showing that the toxicity of KIN4 likely arose from an inhibition of the MEN via the Bub2–Bfa1 GAP complex. The accumulation of anaphase cells (indicative of a mitotic exit defect) in lte1Δ strains carrying an additional copy of CEN-KIN4 confirmed this conclusion (unpublished data). In contrast, anaphase cells did not accumulate in lte1Δ bfa1Δ CEN-KIN4 cells (unpublished data). Thus, Lte1 counterbalances Kin4 activity in a dose-dependent manner in vivo.

Considering that Lte1 inhibited Kin4 catalytic activity in vitro (Fig. 4), we reasoned that Kin4 catalytic activity should increase in the absence of LTE1. However, we did not observe any significant difference in the specific kinase activity of Kin4-6HA enriched from wild-type and lte1Δ cells from asynchronous cultures or from cultures whose cell cycle progression was arrested in G1-, S-, meta- and late anaphase or by SPOC activation (Fig. S3, A–C). Importantly, Lte1 was not present in the Kin4 immunoprecipitates used in the kinase assays, due to the stringent washing conditions required to eliminate cross-contaminating activity from other kinases (unpublished data).

To assess the role of Lte1 in the regulation of Kin4 kinase activity in vivo, we investigated the phosphorylation pattern of the established Kin4 substrate Bfa1 in the presence or absence of LTE1. Bfa1 becomes phosphorylated by the polo-like kinase Cdc5 at anaphase onset (Hu et al., 2001); this phosphorylation causes a transient appearance of the most hyperphosphorylated form of Bfa1-3HA (Fig. 5, B and C; asterisk). Kin4 activity counteracts this phosphorylation (D’Aquino et al., 2005; Pereira and • Bertazzi et al. 1037
Figure 3. Kin4 catalytic kinase activity is required for Kin4–Lte1 interaction. (A) Representative frames and fluorescence intensity line traces of cells expressing KIN4-GFP LTE1-3Cherry. Line profiles above the images represent the fluorescence intensities (FI) in arbitrary units (A.U.), measured for the indicated rectangular area, for Kin4-GFP (green lines) and Lte1-3Cherry (red lines). Note that FI is not comparable between cells. Cell boundaries in each graph are indicated as M and D, and bud neck region as BN. Bar, 5 µm. (B) LTE1 and LTE1-9Myc strains carrying KIN4-6HA (WT), KIN4-6HA elm1Δ, and KIN4-T209A-6HA were subjected to immunoprecipitation using anti-Myc beads. (C) Interaction between Lte1-9Myc and Kin4-6HA was investigated in RTS1 (WT, wild type) and rts1Δ cells upon immunoprecipitation of Lte1-9Myc with anti-Myc beads. (D) Localization of Kin4-GFP and Kin4-T209A-GFP in strains carrying LTE1-3Cherry (no overexpression) and Gal1-LTE1-3Cherry (overexpression), growing in galactose-containing medium. Bars, 5 µm.
Lte1 regulation of Kin4

Lte1 promotes Kin4 hyperphosphorylation

To explore the mechanisms by which Lte1 influences Kin4 activity in vivo, we asked whether Lte1 affected Kin4 phosphorylation and/or localization. Because Kin4 hyperphosphorylation increased upon SPOC activation (Chan and Amon, 2009; Caydasi et al., 2010b), we analyzed Kin4-6HA in wild-type and lte1Δ cells that had been synchronized in G1 phase and subsequently treated with nocodazole to activate the SPOC (Fig. 6 A). Slower migrating, hyperphosphorylated Kin4 forms were observed in wild-type but not in lte1Δ cells (Fig. 6 A, asterisk). The lack of Kin4 hyperphosphorylation in lte1Δ was even more apparent when nocodazole-treated cells reassumed cell cycle progression after nocodazole wash-out (Fig. 6 B). The effect of Lte1 upon Kin4 phosphorylation was not restricted to SPOC activation via nocodazole, as cells held in metaphase by depletion of the APC activator Cdc20 (Gal1-CDC20) also failed to accumulate hyper-phosphorylated forms of Kin4 in the absence of LTE1 (Fig. 6 C).

The kinase Elm1 phosphorylates Kin4 at threonine 209 (T209) within its activation loop (Caydasi et al., 2010b). Kin4-T209 phosphorylation is essential to promote Kin4 catalytic activity and appearance of the hyperphosphorylated forms of Kin4 whose migration is retarded on SDS-PAGE gels (Caydasi et al., 2010b; Moore et al., 2010). We established that lack of Kin4 hyperphosphorylation was not due to inappropriate phosphorylation of Kin4 by Elm1 within the activation loop, as the levels of T209 phosphorylation were comparable to wild-type and lte1Δ cells (Fig. 6, D and E). Thus, Lte1 promotes hyperphosphorylation of active Kin4 in vivo.

The PP2A phosphatase subunit, Rts1, is important for the dephosphorylation and localization of Kin4. In rts1Δ cells, hyperphosphorylated Kin4, while retaining kinase activity, is unable to function in SPOC signaling with Bfa1. This is most likely due its inability to bind to SPBs (Chan and Amon, 2009; Caydasi et al., 2010b). We therefore asked whether hypophosphorylation of Kin4 in lte1Δ cells arose as a consequence of an influence over the function of PP2A-Rts1. If this were the case, deletion of RTS1...
in \textit{lte1}\textsuperscript{Δ} cells should restore Kin4 hyperphosphorylation. Whereas Kin4-6HA was mainly hypophosphorylated in \textit{lte1}\textsuperscript{Δ} cells, phosphorylation of Kin4-6HA increased slightly in \textit{lte1}\textsuperscript{Δ} \textit{rts1}\textsuperscript{Δ} cells (Fig. 6 F). Re-introduction of \textit{LTE1} in the \textit{lte1}\textsuperscript{Δ} \textit{rts1}\textsuperscript{Δ} background restored the full accumulation of hyperphosphorylated Kin4 (Fig. 6 F). Thus, the mechanism by which Lte1 promotes Kin4 phosphorylation is in part counteracted by PP2A-Rts1.

The hyperphosphorylated Kin4 species that arise in the absence of \textit{RTS1} fail to bind to the mother cell cortex and the SPB (Chan and Amon, 2009; Caydasi et al., 2010b). To explore whether this was also the case in \textit{lte1}\textsuperscript{Δ} \textit{rts1}\textsuperscript{Δ} cells, we analyzed the localization of Kin4-GFP in cells carrying the SPB marker Spc42 fused to the red fluorescent protein eqFP611 (Spc42-eqFP; Wiedenmann et al., 2002). Cells were treated with nocodazole to increase the proportion of Kin4 associated with the SPB (Pereira and Schiebel, 2005). Kin4-GFP localized at the mother cell cortex and SPBs in both wild-type and \textit{lte1}\textsuperscript{Δ} cells (Fig. 6, G and H). In agreement with previous
Figure 6. \textit{Lte1} regulates the phosphorylation status of Kin4. (A and B) Phosphorylation of Kin4-6HA in the presence or absence of \textit{LTE1}. (A) Cells carrying \textit{KIN4-6HA} were synchronized in G1 with α-factor (\(t = 0\)) and released into media containing nocodazole to induce metaphase arrest and Kin4 phosphorylation (asterisk). (B) Nocodazole-arrested cells were released in nocodazole-free medium. Clb2 levels and percentage of metaphase cells were monitored over time. (C) Kin4-6HA of wild-type and \textit{lte1Δ} cells arrested in metaphase upon \textit{CDC20} depletion in the presence of solvent control (DMSO) or nocodazole. Asterisks mark Kin4-phosphorylated forms. Tubulin in A–C served as loading control. (D) \textit{KIN4-6HA}, \textit{KIN4-6HA lte1Δ}, and \textit{kin4-T209A-6HA} strains, arrested in metaphase (\textit{CDC20} depletion), were subjected to immunoprecipitation using anti-HA antibodies. The levels of Kin4-6HA and phosphorylation at T209 are shown. (E) Quantification of D. (F) Strains expressing \textit{KIN4-6HA} were arrested in G1 with α-factor and released into medium containing nocodazole for 2 h at 30°C. CEN-LTE1 indicates a centromeric plasmid carrying \textit{LTE1}. Asterisk points to the Kin4-6HA-hyperphosphorylated form. (G) Kin4-GFP localization was determined by fluorescence microscopy in nocodazole-treated metaphase-arrested cells. Spc42-eqFP served as SPB marker (arrowheads). Bar, 5 µm. (H) Quantification of G. (I) SPOC proficiency of the indicated strains grown at 23°C and shifted to 30°C for 4 h before inspection. G and I show one representative experiment of three. 100–150 cells were scored per sample.
Interaction between Kin4 and Lte1 requires Cla4

The Pak kinase Cla4 contributes to mitotic exit by phosphorylating Lte1 and promoting Lte1’s localization at the bud cell cortex (Höfken and Schiebel, 2002; Jensen et al., 2002; Seshan et al., 2002; Yoshida et al., 2003). In the absence of CLA4, Lte1 is no longer able to promote mitotic exit, showing that Cla4-dependent phosphorylation of Lte1 is essential for its mitotic function (Höfken and Schiebel, 2002; Seshan et al., 2002; Yoshida et al., 2003; Nelson and Cooper, 2007). Furthermore, we found that the growth defect of cla4Δ cells at lower temperatures (Höfken and Schiebel, 2002) was rescued by deletion of KIN4 (Fig. 8A), suggesting that increased Kin4 activity might contribute to the growth defect of cla4Δ cells. We therefore asked whether Cla4 influenced the ability of Lte1 to interact with Kin4. In cla4Δ cells, less Kin4 immunoprecipitated with Lte1 (Fig. 8B) (although it was comparable to control cells (Fig. S3D)). Thus, Cla4 regulates Lte1–Kin4 complex formation.

Lte1 inhibits the binding of Kin4 to the dSPB in anaphase

To further explore the role of Lte1 in regulating Kin4, we asked whether Lte1 would influence Kin4 localization in an unperturbed cell cycle (Fig. 7). In LTE1 wild-type cells, Kin4 localizes preferentially at the mother cell cortex and from mid- to late anaphase onwards to the mSPB. Kin4-GFP was barely detected at the dSPB (Pereira and Schiebel, 2005). Deletion of LTE1 profoundly affected localization of Kin4. In 5–7% of lte1Δ cells in anaphase, enhanced binding of Kin4-GFP to the daughter cell cortex was observed in comparison to 1–2% in LTE1 wild-type cells (Fig. 7A, lte1Δ, bottom panel; and unpublished data). The number of lte1Δ cells in which a Kin4-GFP signal could be detected at the dSPB during anaphase was greatly increased compared with wild-type cells at 30°C (Fig. 7A and B; from <1% in LTE1 to 62% in lte1Δ cells). This was even more pronounced at 14°C (<1% in LTE1 in comparison to 80% in lte1Δ cells). Thus, Lte1 impinges upon the recruitment of Kin4 to the dSPB.

Interaction between Kin4 and Lte1 requires Cla4

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Considering that mother cell–located Lte1 promotes mitotic exit of cells with misaligned spindles, we asked whether Lte1-GFP caused SPOC deficiency in SFK1-GBP kar9Δ cells in a Cla4-dependent manner. The binding of Lte1-GFP to Sfk1-GBP was Cla4 independent (Fig. 8D). However, the bud cortex signal of Cla4-3Cherry was decreased in LTE1-GFP SFK1-GBP cells (unpublished data) and the phosphorylated forms of Lte1, which are observed in SFK1-GBP cells, were lost upon deletion of CLA4 (Fig. 8E). This suggested that Cla4-dependent Lte1 phosphorylation, which activates the mitotic function of Lte1, is in place in SFK1-GBP cells. In agreement with this
Figure 8. Lte1–Kin4 interaction requires Cla4. [A] Growth of the indicated strains in YPD plates for 2 d (30°C) and 14 d (11°C). [B] Interaction between Lte1-9Myc and Kin4-6HA was investigated in wild-type (WT) and cla4Δ cells upon immunoprecipitation of Lte1-9Myc with anti-Myc beads. Note that this experiment was done with the experiment shown in Fig. 3 C; i.e., the blots of the WT strain are identical. [C] Quantification of B. One representative experiment of two is shown. [D] Localization of Lte1-GFP. [E] Phosphorylation profile of Lte1-GFP in the indicated strains. The brackets indicate Lte1-phosphorylated forms. [F] Strains were arrested with α-factor and released into nocodazole-containing media. Kin4-6HA and Clb2 levels were determined by immunoblotting at the indicated time points. Tubulin served as a loading control. [G] Localization of Kin4-GFP. Spc42-eqFP served as SPB marker. [H] Quantification of G. One representative experiment of three is shown. Bar, 5 µm.
toward Bfa1 in vivo and in vitro. We propose that Lte1 contributes to mitotic exit working upstream of Tem1 through its inhibitory influence on the activity of Kin4 kinase (Fig. 9).

**Requirements for Lte1–Kin4 interaction**

Our data show that the in vivo interaction between Kin4 and Lte1 requires Kin4 kinase activity and the Pak kinase Cla4, but not the PP2A subunit Rts1. The requirement of Cla4 is explained by the regulation of Lte1 by Cla4. Only Lte1 that is phosphorylated by Cla4 is correctly localized to the bud cortex and has the ability to promote mitotic exit (Höfken and Schiebel, 2002; Jensen et al., 2002; Seshan et al., 2002; Nelson and Cooper, 2007). The kinase-dependent interaction between Kin4 and Lte1 is much less understood. Kin4 could phosphorylate Lte1 and thereby create a binding site for Kin4. However, there is presently no evidence supporting a role for Kin4 in phosphorylating Lte1. Alternatively, Kin4 catalytic activity might be required to dislodge a protein that inhibits Lte1–Kin4 complex formation. Detailed biochemical and biophysical analysis using purified components will be necessary to clarify the molecular requirements of Kin4 activity for the Kin4–Lte1 interaction.

**Lte1-dependent Kin4 regulation**

Our in vitro and in vivo studies suggested that Lte1 is an inhibitor of Kin4 catalytic activity. However, whereas in vitro analysis suggested that Lte1 might work as a competitive inhibitor of...
Kin4 catalytic activity, analysis of Kin4 behavior in \textit{lte1Δ} cells showed that the in vivo situation is more complex. Deletion of \textit{LTE1} had a striking influence upon Kin4 phosphorylation. Interestingly, deletion of \textit{LTE1} inhibited the appearance of hyperphosphorylated forms of Kin4; however, the activating phosphorylation of Kin4 at T209 by Elm1 was as in wild-type cells. Lte1 might control Kin4 by restricting the activity of a phosphatase or by acting as an adapter protein for a kinase that preferentially facilitates the hyperphosphorylation of catalytically active Kin4.

The phosphatase PP2A-Rts1 is involved in Kin4 dephosphorylation (Chan and Amon, 2009). The fact that deletion of \textit{RTS1} only partially restored Kin4 hyperphosphorylation in the \textit{lte1Δ} background implies that Lte1 might promote Kin4 hyperphosphorylation by activating a kinase rather than simply restricting PP2A-Rts1 activity over Kin4.

In addition to influencing Kin4's phosphorylation, Lte1 also affects Kin4 localization (Fig. 9). In the absence of Lte1, Kin4 accumulated at the dSPB in anaphase. This accumulation contrasts the wild-type situation, in which Kin4 is excluded from the daughter cell compartment. We therefore suggest that Lte1 makes a partial but important contribution to Kin4 asymmetry by excluding it from binding to the dSPB. At present, the relationship between Kin4 function, phosphorylation, and localization is not fully understood. Rts1 is crucial for the proper localization of Kin4 to the mother cortex and SPBs (Chan and Amon, 2009). Deletion of \textit{RTS1} leads to the accumulation of hyperphosphorylated and mislocalized, catalytically active Kin4 species (Chan and Amon, 2009; Caydasi et al., 2010b). Given that Kin4 needs to bind to SPBs to phosphorylate Bfa1 upon SPOC activation (Maekawa et al., 2007), the current understanding is that an Rts1-dependent dephosphorylation of Kin4 is required for Kin4 SPB association and hence SPOC function (Caydasi and Pereira, 2009; Caydasi et al., 2010b). Deletion of \textit{LTE1} rescued Kin4 SPB localization in the \textit{rts1Δ} background, despite the fact that Kin4 was partially hyperphosphorylated. The phosphorylation profile of Kin4 in \textit{rts1Δ} and \textit{rts1Δ lte1Δ} cells might differ in key amino acid residues.

How would deletion of \textit{LTE1} restore Kin4 localization at mother and daughter SPBs in \textit{rts1Δ} cells? One possibility is that Kin4 is locked in an intermediate phosphorylated state in \textit{lie1Δ} cells, rendering it insensitive to PP2A-Rts1 control over Kin4 localization. Alternatively, Rts1 might regulate Kin4 localization working upstream of Lte1. In this respect, deletion of \textit{LTE1} not only rescues the localization but also the SPOC deficiency of \textit{rts1Δ} cells. This functional link might indicate that Rts1 could restrict the activity of Lte1 over Kin4; an exciting possibility that awaits detailed analysis. Interestingly, a cross talk between the fission yeast homologue of Lte1, Etd1, and the B-type regulatory phosphatase subunit of PP2A, Pab1, was recently described (García-Cortés and McCollum, 2009; Lahoz et al., 2010). A functional homologue of Kin4 in fission yeast has however not yet been identified.

**Coordination of mitotic exit and SPOC by Lte1**

Our data are consistent with the compartmentalization model that was proposed based on cellular phenotypes upon mislocalization of Kin4 and Lte1 (Bardin et al., 2000; Castillon et al., 2003; D’Aquino et al., 2005; Maekawa et al., 2007; Geymonat et al., 2009a; Chan and Amon, 2010). Keeping Lte1 in the daughter cell is essential for SPOC function, whereas restricting Kin4 activity to the mother cell compartment is important for mitotic exit. We now provide a molecular understanding for the differential distribution of the Kin4 and Lte1 to mother and daughter cell compartments. Lte1 contributes to the exclusion of Kin4 from the dSPB in anaphase, which allows accumulation of Bub2–Bfa1 at the dSPB and subsequent Cdc5-dependent inactivation of the Bub2–Bfa1 GAP complex (Hu et al., 2001; Geymonat et al., 2003). This function of Lte1 is important during an unperturbed cell cycle most likely to inactivate any Kin4 that accidently enters the daughter cell compartment.

It is however puzzling that deletion of \textit{LTE1} also influenced the phosphorylation status of Kin4 in metaphase and upon SPOC activation, two conditions in which Lte1 and Kin4 should be confined to different cellular compartments. A fraction of Lte1 and Kin4 might however colocalize throughout the cell cycle, as supported by the fluorescent profiles of strains carrying Kin4-GFP and Lte1-3Cherry (Fig. 3). It is thus feasible that compartment-specific regulation of Lte1–Kin4 complexes might represent an important level of mitotic regulation, in addition to the control of Lte1 and Kin4 localization.

Previous reports established that forcing Kin4 onto the dSPB delayed mitotic exit and increased the symmetric binding of Bfa1 to SPBs in cells with normally aligned spindles (Maekawa et al., 2007; Caydasi and Pereira, 2009; Geymonat et al., 2009a; Chan and Amon, 2010). At present it is unclear whether the delay in mitotic exit observed in \textit{lte1Δ} at lower temperatures, which can be rescued by deletion of \textit{BFA1} or \textit{KIN4}, arises from the persistent binding of Kin4 to the dSPBs of anaphase cells. Nevertheless, recruitment of Kin4 to the dSPB would explain the increased symmetric localization of Bfa1 observed in \textit{lte1Δ} cells (Geymonat et al., 2009a); a consequence of the increased Bfa1 dynamics in response to Kin4 activity at SPBs (Caydasi and Pereira, 2009).

Cell polarity proteins function in parallel to Lte1 to promote mitotic exit (Höfken and Schiebel, 2002; Seshan et al., 2002; Chirol et al., 2003). The Cdc42 effectors Gic1 and Gic2 were shown to contribute to mitotic exit (Höfken and Schiebel, 2004), possibly by inhibiting the activity of the Bub2–Bfa1 GAP complex. Lte1 and cell polarity determinants all localize to the bud tip, the site where microtubules are anchored by the dynein-dependent pathway (Yeh et al., 1995; Carminati and Stearns, 1997). Interestingly, Kar9 and the cyclin Clb4, which are proteins involved in cytoplasmic microtubule interaction with the cell cortex (Miller and Rose, 1998; Liakopoulos et al., 2003; Maekawa et al., 2003; Maekawa and Schiebel, 2004), also prevent binding of Kin4 to the dSPB by an unknown mechanism (Chan and Amon, 2010). It is thus tempting to speculate that proper cytoplasmic microtubule attachment negatively regulates Kin4 activity via Lte1 at the dSPB and/or alternative mechanisms to allow mitotic exit. This would be in line with a function for the SPBs as a sensor for spindle alignment, as previously suggested (Bardin et al., 2000; Gruenberg et al., 2000; Pereira et al., 2001; D’Aquino et al., 2005; Maekawa et al., 2007; Chan and Amon, 2010).
Centrosomes of higher eukaryotic cells play an important role in the orientation of the mitotic spindle in respect to the cell polarity axis, which is particularly important in cells undergoing asymmetric divisions (Yamashita and Fuller, 2008; Siller and Doe, 2009). It will be interesting to determine whether centrosomes also function as sensors to control cell cycle progression in respect to the cell polarity axis among eukaryotes, as recently suggested for Drosophila germ line stem cells (Cheng et al., 2008; Inaba et al., 2010).

Materials and methods

Growth conditions
Yeast growth conditions in solid and liquid media were as described previously (Sherman, 1991). Yeast strains were grown in yeast peptone dextrose medium with 0.1 mg/l adenine (YPAD). Temperature-sensitive strains were grown at 23°C and shifted to 37°C for phenotypic analysis. For live-cell imaging, yeast cells were grown in sterile-sterilized YPD. Yeast peptone medium containing adenosine and 3% raffinose (YPAR) was used to grow strains carrying genes under the Gal1 promoter (Gal1 promoter repressed). For induction of genes under the Gal1 promoter, galactose (2%) was added to cells growing in YPAR media.

Yeast strains and plasmids
Yeast strains and plasmids are listed in Table S1. Gene deletions and epitope tagging were performed using PCR-based methods (Knop et al., 1999; Janke et al., 2004). Gal1-Clb2(CDC20) (Perreira and Schiebel, 2005), Gal1-clb2ΔD8 (Surana et al., 1993), and Gal1-UPL-TEM1 (Shou et al., 1999) were constructed using integration plasmids. All strains harboring a KAR9 deletion were maintained with KAR9 on a centromeric URA3-based plasmid and analyzed for phenotypes shortly after inducing plasmid loss on S-fluoroorotic acid (S-FOA) containing plates. The GBP:KanMX4 cassette was constructed by replacing GFP from pYM12 (Knop et al., 1999) by the gene coding sequence of the GFP-binding protein (provided by Heinrich Leonhardt, Ludwig-Maximilians University, Munich, Germany).

Cell cycle analysis
For synchronization, yeast cells were incubated with 10 µg/ml α-factor (Sigma-Aldrich) for 2–2.5 h at 30°C until >95% of cells showed a mating projection. To arrest the cells with nocodazole, 15 µg/ml nocodazole (Sigma-Aldrich) was added to the culture medium and incubated for 2–3 h. S-phase arrest was induced by adding 200 µM hydroxyurea (Sigma-Aldrich) to log phase cultures and further incubated for 2–3 h. To induce metaphase arrest of Gal1-Clb2(CDC20) cells, cells grown in YPAR media containing 2% galactose were washed in YPAR media lacking galactose and further incubated in this media until >95% of the cells were large budded with one DAPI-staining region. Gal1-Clb2(CDC20) cells, arrested in metaphase, were released from the cell cycle block by addition of 2% galactose.

Protein methods and reagents
Yeast protein extracts and immunoblotting were performed as described previously (Janke et al., 2004). In brief, cell pellets were collected by centrifugation and resuspended in 1 ml TCA-solution (7.5% trichloroacetic acid and 250 mM NaOH) and kept on ice for 15 min. Samples were centrifuged at 10,000 g for 20 min at 4°C. Precipitated proteins were resuspended in HUUDT (200 µM Tris-HCl, pH 6.8, 8 M urea, 5% SDS, 0.1 mM EDTA, 0.005% Bromophenol blue, and 15 µl/ml DTT). Samples were heated up for 15 min at 65°C before loading onto SDS-PAGE gels. Coomassie Brilliant Blue G-250 (Invitrogen) was used to stain protein gels. For immunoblotting, proteins were transferred from SDS-PAGE gels onto nitrocellulose membrane and stained with Ponceau S (Sigma-Aldrich) before massie Brilliant Blue G-250 (Invitrogen) was used to stain protein gels. For Western blots, proteins were transferred from the nitrocellulose membrane onto a polyvinylidene fluoride (PVDF) membrane (Millipore). PVDF membranes were blocked with 5% nonfat dry milk and incubated with primary antibodies overnight. Antibody incubation was performed in a blocking buffer containing 5% nonfat dry milk and 0.1% Tween-20.

Yeast protein extracts and immunoblotting were performed as described previously (Knop et al., 1999; Janke et al., 2004). For induction of genes under the Gal1 promoter, galactose (2%) was added to cells growing in YPAR media. For the tandem affinity purifications (Puig et al., 2001), whole-cell extracts were prepared from 2 liters of exponentially growing yeast cells expressing KIN4-TAP or LTE1-TAP. Untagged, wild-type strains were used as a negative control. Cell pellets were lysed in aqueous glass beads (Sigma-Aldrich). Lysates were cleared by centrifugation at 10,000 g for 20 min at 4°C and incubated with Magnetic beads coupled to IgGs (Dynabeads; Invitrogen). Extensive pre-incubations were extensively washed with TAP buffer (50 mM Tris-HCl, pH 7.4, 1 mM EDTA, 10 mM DTT, 10% glycerol, and 400 mM NaCl). Kin4- and Lte1-interacting proteins were eluted with 0.5 M NaOH, 0.5 mM EDTA solution and identified by mass spectrometry (MS) upon digestion with trypsin.

Yeast strains and plasmids
Yeast cells harboring Kin4-GFP and Lte1-3Cherry were grown in YPD media or YPARGal plates and analyzed by fluorescence microscopy without washing or fixation. A Z-series of 0.3-µm steps were captured with a microscope (Axiovert 200M; Carl Zeiss) equipped with a 100X N.A. 1.45 Plan-Fluar oil immersion objective (Carl Zeiss), Cascade 1K CCD camera (Photometrics), and Fluorescence microscopy

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Fluorescence microscopy
Yeast cells harboring Kin4-GFP and Lte1-3Cherry were grown in YPD media or YPARGal plates and analyzed by fluorescence microscopy without washing or fixation. A Z-series of 0.3-µm steps were captured with a microscope (Axiovert 200M; Carl Zeiss) equipped with a 100X N.A. 1.45 Plan-Fluar oil immersion objective (Carl Zeiss), Cascade 1K CCD camera (Photometrics), and
SPOC proficiency analysis

For determination of SPOC proficiency kar9Δ cells carrying TUB1-GFP or TUB1-Cherry were grown at 23°C and shifted to 30°C for 3 h. Cells were inspected after fixation with paraformaldehyde for 10 min at room temperature. DNA was stained with DAPI. SPOC proficiency was calculated by dividing the number of SPOC-arrested cells (cells containing two separated DAPI-stained regions and an intact anaphase spindle inside the mother cell) by the sum of SPOC-deficient and arrested cells (cells with more than two DAPI-stained regions in the mother cell body and broken and/or short spindles), 150–200 anaphase cells were counted per sample. Each experiment was done in triplicate.

Online supplemental material

Fig. S1 shows the Kin4 and Lte1 interactors identified by MS analysis and the coimmunoprecipitation using Kin4Δ as bait. Fig. S2 shows the coimmunoprecipitation experiments using Lte1Δ or Kin4Δ as bait. Fig. S3 shows Kin4-specific kinase activity in wild-type and lte1Δ cells. Fig. S4 shows the Bfa1 phosphorylation profile in the absence or presence of lte1. Table S1 presents the list of strains and plasmids used in this manuscript. The Helmholtz association grant (HZ-NF-111) and Marie Curie fellowship (MEXTCT2006-042544) supported the work of G. Pereira; D.T. Bertazzi et al. provided tools; Marcio Lazzarini for help with statistics, Ayse Caydasi and B. Kurtulmus are Marie Curie fellows.

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