Drosophila melanogaster γ-TuRC is dispensable for targeting γ-tubulin to the centrosome and microtubule nucleation

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In metazoans, γ-tubulin acts within two main complexes, γ-tubulin small complexes (γ-TuSCs) and γ-tubulin ring complexes (γ-TuRCs). In higher eukaryotes, it is assumed that microtubule nucleation at the centrosome depends on γ-TuRCs, but the role of γ-TuRC components remains undefined.

For the first time, we analyzed the function of all four γ-TuRC-specific subunits in Drosophila melanogaster: Dgrip75, Dgrip128, Dgrip163, and Dgp71WD. Grip motif proteins, but not Dgp71WD, appear to be required for γ-TuRC assembly. Individual depletion of γ-TuRC components, in cultured cells and in vivo, induces mitotic delay and abnormal spindles. Surprisingly, γ-TuSCs are recruited to the centrosomes. These defects are less severe than those resulting from the inhibition of γ-TuSC components and do not appear critical for viability. Simultaneous cosilencing of all γ-TuRC proteins leads to stronger phenotypes and partial recruitment of γ-TuSC. In conclusion, γ-TuRCs are required for assembly of fully functional spindles, but we suggest that γ-TuSC could be targeted to the centrosomes, which is where basic microtubule assembly activities are maintained.

Introduction

In metazoans, the centrosome organizes the microtubule cytoskeleton. The molecular mechanisms responsible for the initiation and the regulation of microtubule assembly remain unclear, although γ-tubulin appears critical to these processes. In addition to a centrosomal fraction, γ-tubulin is present in cytosolic high-order protein structures (Akashi et al., 1997; Murphy et al., 1998; Oegema et al., 1999; Fujita et al., 2002). In Drosophila melanogaster, two main complexes have been characterized. The simplest ones, called γ-tubulin small complexes (γ-TuSCs), are salt-stable tetramers of ~10S that are composed of two γ-tubulin molecules and two associated proteins, Dgrip84 and Dgrip91 (Oegema et al., 1999). They represent the basic components of the larger complexes, the γ-tubulin ring complexes (γ-TuRCs). γ-TuRCs, whose sedimentation coefficients range from 25 to 35S, contain at least four other proteins (Dgrip75, Dgrip128, Dgrip163, and Dgp71WD) in addition to the γ-TuSC subunits, in a yet unknown stoichiometry. Dgrip75, Dgrip128, and Dgrip163 exhibit sequence homologies called grip motifs, with the two γ-tubulin–associated proteins of the γ-TuSC (Fava et al., 1999; Gunawardane et al., 2000; Murphy et al., 2001). In contrast, Dgp71WD does not posses any grip motifs, but contains seven WD (tryptophan–aspartic acid) repeats (Haren et al., 2006). In vitro, this protein directly interacts with the grip motif containing γ-TuRC subunits, suggesting that it may play a scaffolding role in γ-TuRC organization (Gunawardane et al., 2003). Current models suggest that γ-TuRCs that have been previously assembled in the cytoplasm are recruited to the centrosomes, where they play a role in microtubule nucleation and stabilization (Stearns and Kirschner, 1994; Zheng et al., 1995; Fava et al., 1999; Zhang et al., 2000).

The functions of γ-tubulin and its associated proteins in the γ-TuSC have been extensively studied. Deletion of either corresponding gene is lethal, resulting in an accumulation of cells in mitosis. This is usually correlated with the appearance of strong mitotic defects such as monopolar structures or bipolar spindles exhibiting unfocused poles, impairment of pole

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maturation, and increase in aneuploidy (Oakley et al., 1990; Sunkel et al., 1995; Geissler et al., 1996; Spang et al., 1996; Knop et al., 1997; Barbosa et al., 2000; Paluh et al., 2000; Vardy and Toda, 2000; Hannak et al., 2002; Colombie et al., 2006). In contrast, our knowledge about the function of γ-TuRC–specific components is limited. In *D. melanogaster*, *Dgrip75* is essential for fertility, but not for viability. *Dgrip75* loss-of-function mutants specifically affect localization of the maternal determinant *bicoid* during oogenesis, suggesting a role in the organization or in the dynamics of a subset of microtubules (Schnorrer et al., 2002). In *Schizosaccharomyces pombe*, mutants in *Gfh1* (a *Dgrip75* orthologue) and in *Alp16* (a *Dgrip163* orthologue) exhibit defects associated with altered microtubule function, but without any effect on cell viability (Fujita et al., 2002; Venkatram et al., 2004). Current models cannot explain these data. Instead, they raise questions not only about the redundancy or the specificity of the γ-TuRC–specific proteins but also about the respective functions of γ-TuSCs and γ-TuRCs. In this work, we have tested whether γ-tubulin is only recruited to the centrosome in the form of γ-TuRCs or if γ-TuSCs can be recruited and subsequently matured into functional γ-tubulin complexes by attracting additional components. To this aim, we have developed two strategies: (1) RNA interference (RNAi) in cultured cells involving individual or concomitant depletion of γ-TuRC components, and (2) genetic analyses by taking advantage of the availability of mutant strains (*Dgrip75*, *Dgrip163*, and *Dgp71WD*). We demonstrate that the γ-TuRC–specific subunits display functional specificities and that the γ-TuSCs could be targeted to the centrosome where basic microtubule assembly functions are maintained.

**Results**

*RNAi*-directed depletion of *Dgrip75* impairs cytosolic γ-TuRC assembly or stability

First, we characterized the consequences of the depletion of a γ-TuRC–specific protein on the assembly of cytosolic γ-tubulin complexes. Cultured *D. melanogaster* S2 cells were treated by RNAi to deplete *Dgrip75*, a grip protein specifically present in the γ-TuRC (Fava et al., 1999). The treatment led to a strong decrease of the protein level (Fig. 1 A; >95% of the control level, as judged by Western blot analysis). This effect was specific, as determined by examining the amount of the three γ-TuSC proteins (γ-tubulin, *Dgrip84*, and *Dgrip91*) and actin (Fig. 1, A and D). Immunofluorescence analysis of control cells showed that although *Dgrip75* was undetectable at the interphase centrosome, it localized to the poles at the onset of mitosis, where it was maintained throughout cell division...
In marked contrast, the protein was absent from the mitotic centrosomes in Dgrip75-depleted cells, consistent with Western blot quantification (Fig. 1 B, d; and Table I). When extracts from treated cells were submitted to sucrose gradient sedimentation, γ-TuRCs were severely reduced, as indicated by immunoblotting of soluble fractions with antibodies against γ-tubulin, Dgrip84, Dgrip91, Dgrip128, Dgrip163, and Dgp71WD (Fig. 1 C). The main remaining complexes appeared to be γ-TuSCs, as judged by their protein content and sedimentation coefficient. In addition, the total level, as well as the soluble and cytoskeletal fractions of the three γ-TuSC proteins, are unchanged in control and Dgrip75-depleted cells (Fig. 1 D), suggesting a redistribution of these proteins in the different complexes rather than a change in quantity. In contrast, after Dgrip75-RNAi treatment, we noticed a decrease in the total level of the two other grip-motif proteins of the γ-TuRC, Dgrip128 and Dgrip163. The remaining Dgrip128 protein was distributed on the gradient in the form of heterogeneous and uncharacterized complexes with apparent masses equal or slightly higher than the mass of the γ-TuSC. Dgrip163 protein migrated mainly in light fractions (<10S). One hypothesis could be that this protein was present as a monomeric or dimeric form. Thus, Dgrip75 appears to be required for efficient assembly or stability of cytoplasmic γ-TuRCs.

**Table I. Accumulation of centrosomal components in mitotic cells after Dgrip75 depletion in cultured cells**

<table>
<thead>
<tr>
<th>Staining</th>
<th>Control</th>
<th>Dgrip75RNAi</th>
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<tr>
<td></td>
<td>Bipolar</td>
<td>Monopolar</td>
</tr>
<tr>
<td></td>
<td>%</td>
<td>%</td>
</tr>
<tr>
<td>γ-Tubulin</td>
<td>100 [n = 100]</td>
<td>100 [n = 124]</td>
</tr>
<tr>
<td>Dgrip84</td>
<td>99 [n = 186]</td>
<td>98 [n = 170]</td>
</tr>
<tr>
<td>Dgrip91</td>
<td>91 [n = 103]</td>
<td>100 [n = 55]</td>
</tr>
<tr>
<td>Dgrip75</td>
<td>100 [n = 95]</td>
<td>6 [n = 70]</td>
</tr>
<tr>
<td>Dgrip128</td>
<td>78 [n = 71]</td>
<td>3 [n = 93]</td>
</tr>
<tr>
<td>Dgrip163</td>
<td>98 [n = 132]</td>
<td>24 [n = 119]</td>
</tr>
<tr>
<td>Dgp71WD</td>
<td>95 [n = 109]</td>
<td>100 [n = 65]</td>
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</tbody>
</table>

Microtubules were stained with antibodies against α-tubulin and chromosomes with DAPI, whereas the spindle poles were immunostained with antibodies against γ-tubulin, Dgrip84, Dgrip91, Dgrip75, Dgrip128, Dgrip163, and Dgp71WD. For a correct interpretation, we need to keep in mind a decrease in the total amount of Dgrip128 and Dgrip163 in Dgrip75-depleted cells. n, number of prometaphases/metaphases analyzed. nd, not determined.

Impairment of γ-TuRC assembly induces moderate mitotic defects, but does not preclude γ-TuSC centrosomal recruitment

We next investigated whether Dgrip75 depletion, and thus the subsequent decrease of γ-TuRCs, affected mitotic progression. The mitotic index was significantly increased by ~2.6-fold (2.9%; n = 4,766; probability of 95% [P95] = 2.5–3.3 in treated cells, compared with 1.1%; n = 9,235; P95 = 0.9–1.3 in control cells). This mitotic accumulation coincided with the maintenance of an active mitotic checkpoint, as judged by a strong BubR1 signal at kinetochores (not depicted), which was indicative of a transient block in prometaphase. α-Tubulin immunostaining analysis confirmed an accumulation of cells in prometaphase stages (increased 1.8-fold in frequency), whereas postmetaphase figures exhibited a 2.6-fold decrease relative to controls (Table II). Aberrant mitotic figures were observed, mainly monopolar spindles and bipolar spindles that were either elongated or with lagging chromosomes (Fig. 2 and Table II). Approximately one third of prometaphases and metaphases were still organized as bipolar structures, albeit with a longer interpolar distance (average increase of 50%; 7.9 ± 0.9 in treated cells vs. 5.3 ± 0.7 in controls) and a poor microtubule density. However, treated bipolar spindles exhibited astral microtubules (Fig. 2 A, d, h, and l). Their poles were normally focused, consistent with polar localization of Asp (Fig. 2 B, a), a marker of the spindle microtubules minus ends (Wakefield et al., 2001). They seemed to properly separate their centromeres, as 97% (n = 112; P95 = 92–99) exhibited centromerin (Cnn) labeling at both poles (Fig. 2 B, b; Megraw et al., 1999). Surprisingly, in Dgrip75-depleted cells, γ-tubulin was still recruited to the poles at all stages of mitosis (Fig. 2 A, a–d; and Table I). The maximal fluorescence values raised by γ-tubulin antibodies at the two poles of symmetrical spindles did not differ between Dgrip75-treated and control cells (Fig. 2 C). The two γ-tubulin partners in the γ-TuSC, Dgrip84, and Dgrip91 were also targeted to the poles in an efficient manner (Fig. 2 A, e–i; and Table I). At the same time, pole localization of γ-TuRC-specific proteins (Dgrip128, Dgrip163, and Dgp71WD) was significantly inhibited, but not completely abolished (Fig. 2 D and Table I). The reductions in Dgrip128 or Dgrip163 stainings (Fig. 2 D, a–f, Table I) may result, at least partly, in the overall decrease in the levels of these proteins after Dgrip75 depletion. Dgp71WD staining was present, but reduced in intensity in most of the cases (Fig. 2 D, g–i, insets). To illustrate these changes, we performed double-labeling immunofluorescence on cells depleted of Dgrip75, showing colocalization of γ-tubulin with γ-TuSC-associated proteins (Dgrip84 or Dgrip91) or with Dgp71WD, whereas the γ-TuRC proteins Dgrip128 or Dgrip163 were no longer detectable (Fig. S1, available at http://www.jcb.org/cgi/content/full/jcb.200511071/DC1). The localization of γ-tubulin along spindle microtubules, observed in 97% (n = 138; P95 = 93–99) of control cells, was detected in

**Table II. Distribution of the different mitotic stages after Dgrip75 depletion in cultured cells**

<table>
<thead>
<tr>
<th>Mitotic figures</th>
<th>Control n = 100</th>
<th>Dgrip75RNAi n = 244</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>%</td>
<td>%</td>
</tr>
<tr>
<td>Unfocuseda</td>
<td>21 ± 8b</td>
<td>24 ± 6</td>
</tr>
<tr>
<td>Prometaphases/metaphases</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Symmetrical bipolar</td>
<td>28 ± 9</td>
<td>20 ± 5</td>
</tr>
<tr>
<td>Abnormal bipolar</td>
<td>4 ± 4</td>
<td>22 ± 6</td>
</tr>
<tr>
<td>Monopolar</td>
<td>0</td>
<td>16 ± 4</td>
</tr>
<tr>
<td>Postmetaphase stages</td>
<td>47 ± 10</td>
<td>18 ± 4</td>
</tr>
</tbody>
</table>

Mitotic stages were determined after chromosome staining (DAPI) and immunolabeling of microtubules (α-tubulin antibodies). n, number of mitotic cells analyzed. These results are representative of three independent experiments. *Spindles with no evident polarity, which represent mainly prophase stages or highly abnormal spindles. **Confidence intervals calculated for P95.

Role of γ-TuRC-specific proteins during mitosis • Vérollet et al. 519
only 10% \((n = 146; \text{P95} = 5–15)\) of the depleted cells (Fig. 2 A, a–d). Similarly, the two \(\gamma\)-TuSC proteins, Dgrip84 and Dgrip91, were no longer detected along spindle microtubules (Fig. 2 A, e–l). Moreover, staining of \(\gamma\)-TuSC proteins at the midbody was impaired in treated cells (not depicted). Hence, the assembly of cytoplasmic \(\gamma\)-TuRCs does not appear as a pre-requisite for centrosomal recruitment of the \(\gamma\)-TuSCs, but seems essential for its localization along spindle microtubules and at the midbody.

**Dgrip75 loss-of-function mutant is delayed in mitosis, but still recruits \(\gamma\)-tubulin to the centrosome**

To examine the functional significance of Dgrip75 in vivo, we took advantage of the availability of the mutant allele 175–14 (Dgrip75175), which was either a null or a strong allele (Schnorrer et al., 2002). Dgrip75175 mutant was viable, although adults of both sexes exhibited a slight increase in lethality some days after hatching. Moreover, they showed abnormalities in the abdominal cuticle segmentation and the thoracic macrochaete pattern (unpublished data), which were common in mutations affecting mitosis. Finally, mutant females were sterile because of a failure to undergo normal oogenesis.

Although Dgrip75 did not appear essential for viability, the phenotypes observed after RNAi treatment of cultured cells prompted us to study the mitotic processes in mutant L3 larval brains. Western blot analysis showed that Dgrip75 levels were reduced below the threshold of detection in Dgrip75175 brain extracts (Fig. 3 A). The mitotic index was elevated approximately three-tos fourfold (2.5%; \(n = 12,410; \text{P95} = 2.2–2.8\)) compared with wild-type cells (0.7%; \(n = 16,134; \text{P95} = 0.6–0.8\)). Mutant cells accumulated in prometaphase stages, whereas post-metaphase figures were reduced (unpublished data). More than half of the mutant cells exhibited overcondensed chromosomes.

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**Figure 2. Immunofluorescence analysis of phenotypes induced by Dgrip75 depletion in S2 cells.** (A) Localization of \(\gamma\)-TuSC components in mitotic cells. Control or Dgrip75-treated cells (Dgrip75RNAi) were analyzed by staining with antibodies against \(\gamma\)-tubulin (a and c), Dgrip84 (e and g), and Dgrip91 (i and k). Merge (b, d, f, h, j, and l): blue, chromosomes; red, microtubules; green, \(\gamma\)-TuSC components. (B) Spindle pole focusing and localization of centrosomal markers. Dgrip75-depleted cells were stained with antibodies against Asp (a, green) or Cnn (b, green). Red, \(\alpha\)-tubulin; blue, chromosomes. (C) Quantification of polar \(\gamma\)-tubulin by immunofluorescence. The maximal signal obtained after \(\gamma\)-tubulin staining was scored on each pole of bipolar spindles for control (closed bars) and Dgrip75-depleted cells (open bars). Correlations of 0.73 \((n = 67; \text{P95} = 0.62–0.84)\) and 0.72 \((n = 69; \text{P95} = 0.61–0.83)\), respectively, were observed between the maximal fluorescence measured at the two poles of control or treated cells. (D) Localization of \(\gamma\)-TuRC–specific components (Dgrip128, Dgrip163, and Dgp71WD) in mitotic cells. Blue, chromosomes; red, microtubules; green, \(\gamma\)-TuRC–specific components. Insets represent a hemispindle stained with antibodies against Dgrip128, Dgrip163, and Dgp71WD. For A and D, a quantitative view is shown in Table I. Bars, 5 \(\mu\)m.
ROLE OF γ-TURC–SPECIFIC PROTEINS DURING MITOSIS • VÉROLLET ET AL. 521

(60%; \( n = 186; P95 = 53–67 \)) compared with wild type (2%; \( n = 211; P95 = 0–4; \) Fig. 3 B). These results were consistent with prolonged prometaphase and metaphase stages. However, only very few mutant cells showed an aneuploid phenotype (6%; \( n = 112; P95 = 3–9 \)) compared with wild type (1%; \( n = 116; P95 = 0–3 \)), and this low frequency was consistent with \( Dgrip75^{175} \) adult viability.

Transient prometaphase delays observed in the \( Dgrip75 \) mutant could be a consequence of defects in the mitotic apparatus. Nevertheless, immunofluorescence analysis revealed neither a clear difference in spindle morphology nor a significant decrease of cells stained positively for γ-tubulin or Dgrip84 (Fig. 3 C). Given the limitations of immunofluorescence in neuroblasts, we cannot exclude defects in microtubule organization, such as changes in microtubule density or interpolar distance.

These data suggest that in \( Dgrip75^{175} \) brains, most of the cells complete cell division even though mitosis might be slowed down, consistent with our observations in RNAi-treated cultured cells. γ-TuSC components are still recruited to the centrosomes and most of the mitotic apparatus are able to ensure correct chromosome segregation.

Figure 3. In vivo phenotypes observed in \( Dgrip75 \) mutant brains. (A) Characterization of \( Dgrip75 \) depletion by Western blot. Total protein extracts from wild-type (WT) or \( Dgrip75 \) mutant \( (Dgrip75^{-/-}) \) L3 larval brains were analyzed with affinity-purified \( Dgrip75 \) antibodies. Actin was used as an internal loading control. (B) Analysis of chromosomal figures. Wild-type or mutant L3 larval brains were stained with DAPI. Five brains were scored for each genotype. (C) Immunofluorescence analysis of the mitotic figures in \( Dgrip75 \) mutant neuroblasts. Green, spindle; blue, chromosomes; red, γ-tubulin (a and b) and Dgrip84 (c and d). Bars, 5 \( \mu m \).

Removal of \( Dgrip128 \) or \( Dgrip163 \) is not essential for centrosomal γ-TuSC recruitment

To evaluate whether the phenotypes induced by \( Dgrip75 \) depletion were specific of this grip-motif protein, we performed an RNAi treatment against \( Dgrip128 \) or \( Dgrip163 \), the two other grip-motif γ-TuRC–specific components. Depletion was efficient in either case: after \( Dgrip128 \) RNAi, no poles (\( n = 142 \)) were labeled, and after \( Dgrip163 \) RNAi, <4% of the poles (\( n = 150 \)) were labeled with respective \( Dgrip128 \) and \( Dgrip163 \) antibodies (Fig. 4 A). Western blot analysis confirmed this view (not depicted). Sucrose gradient analysis of extracts depleted for \( Dgrip128 \) or \( Dgrip163 \) showed a marked decrease in γ-TuRC content (Fig. 4 B). These experiments show that these proteins, like \( Dgrip75 \), appear involved in the assembly or stability of γ-TuRCs. Moreover, silencing of either protein led to accumulation of cells in mitosis (approximately three- to fourfold), which resulted from a higher frequency of prometaphases concomitant with a decrease in postmetaphase stages (Fig. 4 C). Mitotic phenotypes were mainly characterized by elongated bipolar spindles and less frequently by monopolar structures. γ-Tubulin (Fig. 4 D) as well as Dgrip84 and Dgrip91 (not depicted) were efficiently recruited to the centrosomes, even in mitotic cells that exhibited severe phenotypes (i.e., monopolar spindles). However, γ-tubulin labeling along spindle microtubules was no longer observed (Fig. 4 D). It was striking that these phenotypes were similar to the defects obtained after depletion of \( Dgrip75 \).

Furthermore, we next performed an in vivo study using a \( Dgrip163 \) P insertion mutant (\( Dgrip163^{GE27807} \)) in which the \( P \) transposable element was inserted into exon 4 (out of 5) between codons 822 and 823. This mutant behaved genetically as a null or strong allele (unpublished data). \( Dgrip163 \) mutants exhibited reduced viability. The emerging homozygous or hemizygous \( Dgrip163^{GE27807} \) adults displayed abdominal abnormalities, and mutant females were sterile. Altogether, these studies reveal that inhibition of each individual grip-motif protein specific of the γ-TuRC leads to similar phenotypes in cultured cells and in vivo.

Dgp71WD is neither essential for γ-TuRC assembly nor for its recruitment to the centrosome

In contrast to the other proteins specific of the γ-TuRCs, Dgp71WD does not belong to the same structural family (Gunawardane et al., 2003). RNAi treatment induced a strong inhibition, as judged by Western blot analysis (Fig. 5 A). In agreement with this quantification, Dgp71WD centrosomal
recruitment was severely impaired in treated cells (1%; \( n = 385; \) \( P95 = 0–2 \)) compared with control cells (95%; \( n = 296; \) \( P95 = 93–97 \); Fig. 5 B). In treated cells, we noticed the systematic loss of Dgp71WD staining on other mitotic structures, such as spindle microtubules (Fig. 5 B, inset) and midbodies (not depicted). As Dgp71WD contains WD repeats and interacts \( \gamma \)-TuRC assembly. When control extracts were subjected to sucrose gradient sedimentation, this protein appeared not only as part of \( \gamma \)-TuRCs but also of smaller uncharacterized complexes (Fig. 5 C). In Dgp71WD-depleted conditions (Fig. 5 C), no significant modification in the quantity of \( \sim 30S \) complexes is observed, as judged by \( \gamma \)-tubulin or Dgrip75 labeling. This result suggests that Dgp71WD, in contrast to Dgrip75, appears to be dispensable for the assembly or the stability of large \( \gamma \)-tubulin complexes. However, we cannot exclude the possibility that the assembly of these complexes is mediated by residual amounts of Dgp71WD.

Studies of the mitotic phenotypes observed after Dgp71WD down-regulation revealed similarities to the defects recorded after the removal of Dgrip75, Dgrip128, or Dgrip163; an increase of the mitotic index (2.7%; \( n = 7,970; \) \( P95 = 2.3–3.1 \)) compared with control cells (0.9%; \( n = 19,650; \) \( P95 = 0.8–1 \); and an accumulation of cells in the prometaphase and metaphase stages (Fig. S2 A, available at http://www.jcb.org/cgi/content/full/jcb.200511071/DC1). Dgp71WD-depleted cells showed defects in spindle morphology, such as bipolar spindles with unfocused poles or monopolar structures (Fig. 5, B and D; and Fig. S2 B) with unseparated poles, as determined by Cnn staining (not depicted). \( \gamma \)-TuSC components (\( \gamma \)-tubulin, Dgrip84, and Dgrip91), as well as \( \gamma \)-TuRC–specific subunits (Dgrip75 and Dgrip163), were still targeted to the mitotic centrosomes, but often the stainings were weak (Fig. 5 D and Fig. S2 B). These proteins were no longer detected at the spindles. These results suggest that Dgp71WD is dispensable for \( \gamma \)-TuRC assembly and recruitment, but is required for the efficient function of this complex.
A genetic approach was performed to test this hypothesis in vivo. We used a mutant strain that carried a P element insertion (GE30807) in the 5′ untranslated mRNA region at position 49 from the translation initiator ATG. As shown by Western blot, the level of Dgp71WD was strongly reduced in a mutant larval brain extract compared with a wild-type extract (Fig. 6 A). An immunofluorescence analysis performed in larval brains sustained that Dgp71WD was no longer detectable in mutants (n = 120), whereas 96% of wild-type poles (n = 110; P95 = 92–100) exhibited a clear staining (Fig. 6 B). These results indicate that Dgp71WD GE30807 is at least a strong allele. This was confirmed by analysis of hemizygous flies, using a chromosomal deficiency that covers Dgp71WD.

Most of the homozygous or hemizygous mutants reached the adult stage. However, they showed a shorter life, morphological abnormalities, and female sterility. Mitotic phenotypes had been analyzed in L3 larval brains confirming RNAi-mediated phenotypes. We observed a fourfold increase in the mitotic index, an accumulation in prometaphase stages associated with a significant hypercondensation of chromosomes and a high incidence of disorganized or monopolar spindles (Fig. S3, available at http://www.jcb.org/cgi/content/full/jcb.200511071/DC1). However, γ-tubulin is detected at all the centrosomes of Dgp71WDGE30807 neuroblasts (n = 130; Fig. 6 C). Moreover, mutant neuroblasts exhibited an increase in aneuploidy (12%; n = 139; P95 = 7–17) versus wild-type brains (1%; n = 125; P95 = 0–5), suggesting that spindles without Dgp71WD were not fully functional.

**γ-TuSC components are targeted to the poles even after cosilencing of the four γ-TuRC subunits**

Whatever their role in the assembly and recruitment of cytosolic γ-tubulin complexes, the inhibition of each individual γ-TuRC subunit allowed efficient γ-TuSC anchoring to the centrosomes. It was possible that some grip-motif proteins were in part redundant and could complement each other to some extent. Alternatively, the residual γ-TuRC–specific proteins could account for the formation of a γ-TuRC–like structure that was unstable in the cytoplasm, but stabilized upon assembly with the pericentriolar matrix. Hence, we performed cosilencing of the four proteins Dgrip75, Dgrip128, Dgrip163, and Dgp71WD. The levels of the four proteins were dramatically reduced as judged by Western blot (Fig. 7 A) and immunofluorescence analyses (Fig. 7 B and Table III). The mitotic index was increased in treated cells (3.9%; n = 2,552; P95 = 3.4–4.4) compared with control cells (1.1%; n = 8,567; P95 = 0.8–1.4). Most of the mitotic structures consisted of monopolar prometaphases/metaphases (70%; n = 80; P95 = 60–80) compared with control cells (1%; n = 104; P95 = 0–3; Fig. 7, B and C). A striking feature of this treatment was the significant accumulation of polyploid cells and the appearance of very large interphase cells.
be directly targeted to the poles. γ-TuSC assembled in the cytoplasm can, at least to some extent, type (unpublished data). Altogether, these data confirm that the m. γ-tubulin (C). Bars, 5 μm.

Dgp71WD (B) or μ-tubulin to the spindle poles of mutant neuroblasts. green, microtubules; blue, chromosomes; red, Dgp71WD (B) or γ-tubulin (C). Bars, 5 μm.

Discussion

The molecular mechanisms responsible for γ-tubulin recruitment to nucleation centers and microtubule nucleation itself remain poorly understood. Several models have been proposed, and all imply that the cytosolic assembly of γ-TuRCs is a critical step for γ-tubulin targeting to the centrosome. For the first time, we analyzed the function of all four γ-TuRC–specific subunits during mitosis, using either RNAi treatment in cultured cells or genetic analyses in D. melanogaster.

γ-TuRC-specific proteins play a distinct role in γ-TuRC assembly

Individual depletion of Dgrip75, Dgrip128, or Dgrip163 induced a strong decrease in cytoplasmic γ-TuRCs in cultured S2 cells. This is consistent with previous data obtained in vitro showing that immunodepletion of Xgrip210, the Xenopus laevis orthologue of Dgrip163, blocks the reassembly of purified and salt-treated γ-TuRCs (Zhang et al., 2000). In contrast to the strong decrease of γ-TuRCs observed after the depletion of grip-motif proteins, the down-regulation of Dgp71WD did not produce a significant effect on the ratio and the sedimentation coefficients of the γ-tubulin complexes. However, we cannot exclude the possibility that residual amounts of this protein are sufficient to maintain the overall structure of these complexes. Dgp71WD is probably associated with the γ-TuRCs at a low stoichiometry, explaining why no change in the sedimentation coefficient could be detected after its depletion. Our data suggest that although the grip-motif proteins are essential for the assembly and stability of γ-TuRCs, Dgp71WD appears dispensable for these processes.

The complete γ-TuRC is necessary for an efficient progression through mitosis

In Dgp71WD-depleted cells, we observed no obvious effect on the level of cytosolic γ-TuRCs and on the recruitment of the different γ-TuRC components to the mitotic centrosomes, suggesting that γ-TuRCs depleted of Dgp71WD had not completely lost their ability to be targeted to the poles. However, γ-tubulin localization along spindle microtubules was impaired and monopolar figures with poorly separated poles appeared with a high occurrence. These monopolar phenotypes are similar to those observed after depletion of Nedd1, the human Dgp71WD orthologue (Haren et al., 2006). The mitotic defects could result from a partial functionality of the γ-tubulin complexes recruited to the centrosomes or from modification in the properties of microtubule arrays that were no longer decorated by γ-tubulin. Thus, it appears that the complete γ-TuRC is required for normal mitotic progression.

γ-TuRC assembly is not a prerequisite for the centrosomal targeting of γ-tubulin

If one assumes that γ-tubulin is recruited to the centrosome in the form of γ-TuRCs, depletion of γ-TuRCs and γ-tubulin would be expected to lead to similar phenotypes. In fact, depletion of grip-motif components of the γ-TuRCs, such as Dgrip75, did not fully mimic the effects of depletion of γ-TuSC subunits on mitotic progress (Sunkel et al., 1995; Barbosa et al., 2000; Paluh et al., 2000; Vardy and Toda, 2000; Fujita et al., 2002; Venkatram et al., 2004; Colombie et al., 2006). Less severe defects were observed as indicated by the lower occurrence of monopolar spindles, the efficient distribution of centrosomes to the two poles, the maintenance of bipolar spindles with astral microtubules and focused poles. These phenotypes had been confirmed in vivo, using mutant larval neuroblasts. These moderate defects were consistent with the viability of Dgrip75 mutants and the weak percentage of aneuploidy. Surprisingly, despite a significant reduction of the cytosolic pool of γ-TuRCs, the

Figure 6. In vivo phenotypes observed in Dgp71WD mutant brains. (A) Analysis of Dgp71WD depletion in mutant brains by Western blot. Wild-type or Dgp71WD mutant (Dgp71WD−/−) brains were analyzed with Dgp71WD antibodies or actin (internal loading control). (B) Organization of spindles in mutant neuroblasts. green, microtubules; blue, chromosomes; red, Dgp71WD (B) or γ-tubulin (C). Bars, 5 μm.
tethering of the three γ-TuSC proteins to the centrosomes was not impaired, as judged by immunofluorescence analyses both in mutant neuroblasts and in cultured cells. Similar phenotypes were observed after individual silencing of the two other grip-motif proteins, Dgrip128 and Dgrip163. It was noteworthy that after an RNAi treatment against any one of the three grip-motif proteins specific of the γ-TuRCs, a decrease in the level of the two others was noticed (unpublished data), suggesting some co-regulation between this set of proteins. Altogether, these results show that the complete γ-TuRC is not a prerequisite for the centrosomal accumulation of γ-TuSC proteins.

γ-Tubulin can be recruited to the centrosomes as γ-TuSC

Because γ-tubulin could be recruited to the centrosomes independently of the γ-TuRCs, we wanted to characterize the complex that mediates its targeting. Sucrose gradient and immunofluorescence analyses after depletion of individual γ-TuRC components strongly suggest that γ-tubulin is recruited to the centrosomes as γ-TuSC. Codepletion of the four γ-TuRC–specific proteins still allowed the polar accumulation of the γ-TuSC components in most of the cells. This latter experiment supports the hypothesis that the γ-TuSC is a vector competent for targeting γ-tubulin to the centrosome. However, as RNAi treatments do not completely remove all protein, it is possible that the residual γ-TuRC–specific proteins participate in the formation of a γ-TuRC–like structure that is unstable in the cytoplasm, but stabilized upon assembly within the pericentriolar matrix. The idea that γ-TuSC could be recruited to the poles is consistent with data reported in *Saccharomyces cerevisiae*, in which no orthologues of the γ-TuRC–specific proteins have been reported. In this organism, the main interactions of γ-tubulin with the microtubule-organizing centers are mediated by the two γ-tubulin–associated proteins in the γ-TuSC (Knop and Schiebel, 1997, 1998; Nguyen et al., 1998; Vinh et al., 2002). Similarly, in mammals, the γ-TuRCs can be tethered to the centrosomes via interactions of the orthologues of Dgrip84 and Dgrip91 with the centrosomal anchoring proteins kendrin and centrosome- and Golgi-localized protein kinase N–associated protein (Takahashi et al., 2002; Zimmerman et al., 2004). In *D. melanogaster*, the calmodulin-binding protein CP309 has been proposed to anchor γ-TuRCs to the centrosome by direct binding to the γ-TuSC (Kawaguchi and Zheng, 2004). However, γ-tubulin recruitment to the centrosomes is probably a complex process, as additional proteins, including ninein (Delgehyr et al., 2005) and centrosomin (Terada et al., 2003), provide alternative sites for γ-tubulin anchorage. Although evidence suggests that γ-tubulin can be targeted to the centrosomes in the form of γ-TuSCs, we cannot rule out that in *D. melanogaster* other proteins at low abundance, previously uncharacterized proteins such as the Dgrip79 and Dgrip223 identified by in silico analyses, or different combinations of known γ-tubulin...
partners could be involved in this recruitment (Gunawardane et al., 2003). Moreover, after concomitant depletion of all γ-TuRC–specific proteins, spindles were nonfunctional and the amount of γ-TuSC recruited to the mitotic centrosomes was reduced. Because of the observation that the small complexes are less active than γ-TuRCs in promoting nucleation (Oegema et al., 1999; Gunawardane et al., 2000), we propose that the decrease in the amount of γ-TuSCs under a threshold can be critical for spindle functionality.

Interestingly, the transient association of γ-tubulin to the spindle, as well as to the midbody (Julian et al., 1993; Lajoie-Mazenc et al., 1994; Khodjakov and Rieder, 1999; Raynaud-Messina et al., 2000, 2004), was no longer detectable after individual or concomitant RNAi treatment against Dgrip75, Dgrip128, Dgrip163, and Dgp71WD. Hence, mitotic γ-tubulin localization to structures outside the pericentriolar material requires the fully assembled γ-TuRCs, and Dgp71WD could play an active role in this process. The absence of γ-tubulin recruitment along the spindle microtubules and at the midbody can also be explained by distinct properties of docking proteins at centrosomes and at noncentrosomal sites. Actually, novel proteins in charge of the recruitment of γ-tubulin complexes along spindle microtubules have recently been identified in fission yeast (Sawin et al., 2004; Janson et al., 2005).

**Poles lacking complete γ-TuRC fulfill basic microtubule organization**

In contrast with the impairment of microtubule assembly from the pericentriolar material after removal of γ-TuSC components (Barbosa et al., 2000; Raynaud-Messina et al., 2004; Colombie et al., 2006), the pericentriolar material appears to remain active as a microtubule-nucleating center after down-regulation of specific γ-TuRC proteins, as shown by the presence of astral microtubules identified both by immunofluorescence and electron microscopy (unpublished data). Moreover, microtubules containing 13 protofilaments were still nucleated, challenging the “template model” (Fig. S4, available at http://www.jcb.org/cgi/content/full/jcb.200511071/DC1; Moritz and Agard, 2001). This indicates that preassembly of cytosolic γ-TuRCs does not seem required for the formation of canonical microtubules. It is not excluded that a ringlike structure could be assembled from γ-TuSCs alone or in combination with small amounts of γ-TuRC–specific proteins at the pericentriolar level, although the purified γ-TuSC does not assemble into larger structures in vitro (Oegema et al., 1999). In that case, such ringlike complexes would exhibit stoichiometries and protein compositions different from γ-TuRCs.

Our observations could reconcile the findings of microtubule nucleation in animal cells and in S. cerevisiae. Moreover, after depletion of γ-TuRC–specific components, recruitment of γ-TuSC appeared sufficient, in most of the cases, to control the formation of spindles competent for chromosome segregation. However, mitotic processes were partly disrupted in cells lacking γ-TuRCs, leading to a transient prometaphase accumulation and a poor density of spindle microtubules. Several possibilities could account for these defects, such as a lower efficiency of γ-TuSCs compared with γ-TuRCs in microtubule nucleation (Oegema et al., 1999; Gunawardane et al., 2000), the presence of distinct binding sites for γ-TuRCs in microtubule nucleation. Moreover, after concomitant depletion of γ-TuRC–specific proteins, recruitment of γ-TuSC appeared sufficient, in most of the cases, to control the formation of spindles competent for chromosome segregation. However, mitotic processes were partly disrupted in cells lacking γ-TuRCs, leading to a transient prometaphase accumulation and a poor density of spindle microtubules. Several possibilities could account for these defects, such as a lower efficiency of γ-TuSCs compared with γ-TuRCs in microtubule nucleation (Oegema et al., 1999; Gunawardane et al., 2000), the presence of distinct binding sites for γ-TuSCs and γ-TuRCs, or the loss of γ-TuRC recruitment along spindle microtubules, which would affect the organization or the dynamics of specific microtubule arrays.

Collectively, our results strongly suggest that the assembly of γ-TuRCs is not essential for γ-tubulin–dependent microtubule nucleation at the centrosome, but instead is required for other, noncentrosomal localization of γ-tubulin. In D. melanogaster, γ-TuRCs may target a fully organized “nucleation machinery” to the sites of nucleation; γ-TuSC would act as a functional unit, whereas γ-TuRC–specific proteins would rather play a more refined role in regulating or optimizing of microtubule arrays during mitosis. γ-Tubulin can be targeted to the poles by the direct docking of γ-TuSCs that exert basic nucleation activity (Fig. 8). This γ-TuSC recruitment could be a “salvage pathway,” involved only when the dominant microtubule
assembly mechanism mediated by γ-TuRCs is impaired or as a physiological pathway acting in parallel to γ-TuRC nucleation activity. Altogether, our data should prompt the reexamination of the current nucleation models.

Materials and methods

Cell culture and RNA-mediated interference

RNAi was performed in S2 D. melanogaster cells (Schneider, 1972) according to Clemens et al. (2000). Cells were treated twice with RNAi at days one and five and harvested on day seven for immunoblotting and immunofluorescence staining. To perform coincubation, cells were incubated in the same way, with the four double-stranded RNAs (dsRNAs) against Dgrip75, Dgrip128, and Dgrip163, and Dgp71WD together (20 mM each). The dsRNAs used correspond to positions 717–1,577, relative to the start position for Dgrip75; 717–1,577, relative to the start position for Dgrip128; 717–1,577, relative to the start position for Dgrip163; and 717–1,577, relative to the start position for Dgp71WD (clone RE59956). These dsRNAs were generated from the cDNA plasmid clones as described in Raynaud-Messina et al. (2004).

Fly strains

Strains w1118 or w118 were used as control strains, and mutant strains Dgrip75/75 (Schroemer et al., 2002), Dgrip128/128, and Dgrip163/163 (GenExel, Inc.) were used for the study. Each mutant chromosome was balanced over CyO P[Kr:Gal4], TM3,Sb,P[Kr:Gal4], and TM3,Sb,P[Kr:Gal4], respectively. The following rabbit antibodies were raised: R62 against 23C-tubulin (Raynaud-Messina et al., 2004); R403 against the COOH-terminal peptide of γ-tubulin; and R367 against the recombinant COOH-terminal γ-tubulin; respectively, were used to produce hemizygous flies. Note that Dgp71WD is referred as Dgrip71 in the Flybase database (http://flybase.bio.indiana.edu).

Antibodies

The mouse monoclonal antibodies T5168, GTU88 (Sigma-Aldrich), and 1501 (CHEMICON International, Inc.) were used to stain γ-tubulin, γ-tubulin, and α-tubulin, and actin, respectively. The following rabbit antibodies were raised: R26 specifically directed against D. melanogaster 23C-γ-tubulin (Raynaud-Messina et al., 2004); R403 against the COOH-terminal peptide of Dgrip19 (1-554) (Raynaud-Messina et al., 2004); and R367 against the recombinant full-length Dgrip84. R300 was raised against recombinant full-length Dgrip75 and affinity purified on recombinant Dgrip75, which is produced in Escherichia coli, R267 was raised against the recombinant COOH-terminal Dgrip171 (486–647 aa), and R360 against the recombinant COOH-terminal Dgrip163 (1150–1352 aa) was used for Western blot. For γ-tubulin detection, we used GTU88 for Western blot analysis and R26 for immunofluorescence experiments, except for double labeling, where monoclonal GTU88 antibody was used instead of polyclonal R26 antibody. We also used Rb666 against Bub1 (Logarinho et al., 2004; gift from C. Sunkel, Universidad do Porto, Porto, Portugal), Rb3133 against Asp (Sauders et al., 1997; gift from D. Glover, University of Cambridge, Cambridge, UK), R19 against D. melanogaster (Harauz et al., 1995), and antibodies against Dgrip128 and Dgrip163 (Gunawardane et al., 2000; gift from Y. Zheng, Howard Hughes Institute, Carnegie Institution of Washington, Baltimore, MD).

Sucrose gradients

5–40% sucrose gradient was prepared as described previously (Oegema et al., 1999). Cells extract (500 µg of protein) was overlaid onto the gradient and centrifuged for 4 h and 30 min at 4°C in a swinging rotor (SW55.1; Beckman-Coulter) at 45,000 rpm (average 150,000 g). 10 µL/ml fractions were collected and 250 µL of each fraction were precipitated with cold methanol (−20°C). The nine fractions corresponding to the soluble part of the extracts were analyzed by immunoblotting. The calibration of the gradient was determined by running in parallel 0–3 h D. melanogaster embryonic extracts that contain γ-TuScs and γ-TuRcs.

Western blotting

Protein extracts from cultured cells (Raynaud-Messina et al., 2004) and from total larval brains (Colombie et al., 2006) were prepared and subjected to Western blot analyses (7.5% polyacrylamide gel and SDS [Sigma-Aldrich]). Apparent masses were determined by comparison with the SDS-PAGE molecular weight standards (broad range), which were obtained from Bio-Rad Laboratories.

Cytological analysis and microscopy techniques

For immunostaining, S2 cultured cells and semisquashed L3 larval brains were fixed and permeabilized as previously described (Colombie et al., 2006). DAPI staining of squashed third instar larvae was performed as described previously (Sunkel et al., 1995). For detection, secondary antibodies conjugated to Alexa Fluor 488 or 568 (Invitrogen) were used. Fluorescence microscopy was performed on a microscope (Axiovert; Carl Zeiss Microimaging, Inc.) equipped with a 2-motor, using 100×/1.4 NA objectives. z-series images were acquired with a camera and software (AxioCam MRm and Axiovision; Carl Zeiss Microimaging, Inc.). S2 cell or brain images were subsequently deconvolved using Axiovision, and Z-planes were projected onto a single view. The percentages of the different mitotic phenotypes were determined with confidence intervals calculated for a Poisson. Usually, image processing and quantification of fluorescence were done using Photoshop (Adobe). For quantification of polar γ-tubulin by immunofluorescence, we measured the maximal signal obtained after γ-tubulin staining. The signal of the camera was proportional to the fluorescence intensity, and the background and the maximal fluorescence values were adjusted to 0 and 25, respectively (Lajoie-Mazenc et al., 1994).

Online supplemental material

Fig. S1 shows colabeling of γ-tubulin with associated proteins in Dgrip75-depleted cells. Fig. S2 shows phenotypes observed after Dgp71WD depletion in S2 cells. Fig. S3 shows mitotic phenotypes in the Dgrip75 WD mutant. Fig. S4 shows the number of microtubule protofilaments in mitotic cells after Dgrip75 RNAi treatment. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.200511071/DC1.

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