Misato Controls Mitotic Microtubule Generation by Stabilizing the TCP-1 Tubulin Chaperone Complex

Highlights

- Misato interacts biochemically with the Tubulin chaperone complex, TCP-1/CCT
- Misato stabilizes the TCP-1 complex, possibly by filling its Tubulin-folding cavity
- Loss of Misato or TCP-1 complex subunits leads to similar mitotic phenotypes
- In the absence of Misato, Tubulin is unstable and unable to efficiently polymerize

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In Brief

Palumbo et al. show the conserved protein Misato interacts with the Tubulin chaperone complex, TCP-1/CCT, in Drosophila. In the absence of Misato, TCP-1 subunit levels are reduced, functional Tubulin is unstable, and mitotic microtubule generation is compromised. The predicted structure of Misato suggests it acts as a molecular placeholder in the absence of Tubulin, stabilizing the TCP-1 complex.
Misato Controls Mitotic Microtubule Generation by Stabilizing the TCP-1 Tubulin Chaperone Complex

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SUMMARY

Mitotic spindles are primarily composed of microtubules (MTs), generated by polymerization of α- and β-Tubulin hetero-dimers [1, 2]. Tubulins undergo a series of protein folding and post-translational modifications in order to fulfill their functions [3, 4]. Defects in Tubulin polymerization dramatically affect spindle formation and disrupt chromosome segregation. We recently described a role for the product of the conserved misato (mst) gene in regulating mitotic MT generation in flies [5], but the molecular function of Mst remains unknown. Here, we use affinity purification mass spectrometry (AP-MS) to identify interacting partners of Mst in the Drosophila embryo. We demonstrate that Mst associates stoichiometrically with the Tubulin chaperone complex, TCP-1/CCT, with the hetero-hexameric Tubulin Prefoldin complex, and with proteins having conserved roles in generating MT-competent Tubulin. We show that RNAi-mediated in vivo depletion of any TCP-1 subunit phenocopies the effects of mutations in mst or the Prefoldin-encoding gene merry-go-round (mgr), leading to monopolar and disorganized mitotic spindles containing few MTs. Crucially, we demonstrate that Mst, but not Mgr, is required for TCP-1 complex stability and that both the efficiency of Tubulin polymerization and Tubulin stability are drastically compromised in mst mutants. Moreover, our structural bioinformatic analyses indicate that Mst resembles the three-dimensional structure of Tubulin monomers and might therefore occupy the TCP-1 complex central cavity. Collectively, our results suggest that Mst acts as a co-factor of the TCP-1 complex, playing an essential role in the Tubulin-folding processes required for proper assembly of spindle MTs.

RESULTS AND DISCUSSION

Misato Biochemically Interacts with the Tubulin Chaperone Complex, TCP-1/CCT

We previously demonstrated that mutations in misato (mst) lead to frequent monopolar spindles (~70%) with low microtubule (MT) density in Drosophila larval brains and that, upon MT regrowth after cold exposure, mst cells are primarily defective in kinetochore-driven MT generation [5].

To address the molecular function of Mst, we expressed a GFP-tagged variant in the Drosophila syncytial blastoderm embryo, which undergoes a series of rapid, synchronous nuclear divisions (Figure 1A; Movies S1 and S2). Expression of Mst-GFP fully rescued the lethality associated with the mst1 mutation and, under the control of the V32-GAL4 female germline-specific driver, Mst-GFP was expressed in embryos at similar levels to endogenous Mst (Figure 1B). Time-lapse experiments (Figure 1A; Movies S1 and S2) and immunolocalization studies on fixed larval brains (Figure 2F) failed to reveal Mst enrichment on the spindle. Thus, Mst is specifically associated with spindles of syncytial embryos.

To establish the biological process in which Mst functions, we sought to identify interacting partners. Syncytial embryos, which undergo 13 mitotic divisions predominantly in the absence of cytokinesis, contain large amounts of mitotic proteins. We developed a pipeline based on GFP-TRAP-A affinity purification and mass spectrometry (AP-MS) combined with bioinformatic-based removal of non-specific contaminants, which allows the robust identification of GFP-bait protein interacting partners (see Supplemental Experimental Procedures). We undertook this procedure in triplicate for embryos expressing Mst-GFP.
Incubation of 0–3 hr clarified embryo extracts with GFP-TRAP-A consistently depleted ~95% of Mst-GFP, without affecting levels of untagged Mst, demonstrating that Mst is a monomer in the embryo (Figure 1C). Bioinformatics-based analysis of AP precipitates identified interactors, which suggested a core functional relationship with Mst. Table 1 shows a list of Mst-GFP interacting proteins from one of these experiments, after stringent filtering (for an extended list based on less stringent filtering, see http://www.thewakefieldlab.com/ms.html). A set of eight “top hit” proteins had similar MS scores and peptide coverages as Mst itself (~1,300–3,000 and 70%–90%, respectively). These constitute all the subunits of the Tubulin chaperone complex, TCP-1 (Table 1). Also known as the Chaperonin Containing TCP-1 (CCT) or Tcp-1 Ring Complex (TRiC), the TCP-1 complex is an integral component of the Tubulin-folding pathway [3, 6, 7]. In this pathway, a complex termed Prefoldin [8, 9] interacts with newly synthesized α- and β-Tubulin, delivering them to the “donut”-shaped TCP-1 complex, which provides a suitable environment in which Tubulin can be correctly folded [3, 6–9]. A further set of conserved TCP-1 complex modulating proteins are additionally required, ensuring newly translated α- and β-Tubulin can be modified and incorporated into MTs [10]. The same pathway controls the proper folding of Actin and γ-Tubulin [11, 12].

All six subunits of the Prefoldin complex were also present in Mst-GFP AP precipitates (Table 1). Furthermore, two of the five remaining significant hits (Viaf and PDCD-5) are TCP-1 interactors. Viaf, a member of the Phosducin family, associates with the TCP-1 complex in flies [13], and its closest human homolog (PhLP-3) binds TCP-1, working antagonistically to Prefoldin [14]. Similarly, the human homolog of PDCD-5 binds to Phosducin and TCP-1β, interfering with the Tubulin-TCP-1 complex interaction [15].

The interaction between Mst and TCP-1α, the only subunit of the TCP-1 complex for which antibodies are available, was verified through reciprocal immunoprecipitation and western blotting in wild-type and Mst-GFP-expressing embryos (Figures 1D and 1E). Thus, the interaction between the TCP-1 complex and Mst reflects a normal in vivo interaction. Moreover, quantitative analysis of the MS data confirmed that similar quantities of all eight TCP-1 complex subunits and Mst were present in AP precipitates (Table 1). From this we infer that one molecule of Mst has the ability to interact with a single TCP-1 hetero-octameric complex. Notably, the Prefoldin subunits and the additional Tubulin-folding interactors were precipitated at ~10- to 100-fold-lower amounts than the TCP-1 subunits. This most likely reflects transient ternary complexes that are formed between these proteins and the TCP-1 complex [8–10, 13–15]. In summary, we demonstrate an in vivo biochemical relationship between Mst, the TCP-1 complex, and other Tubulin-folding pathway components.

**Depletion of the TCP-1 Complex Phenocopies the Loss of Mst and Merry-Go-Round**

To investigate the functional relationship between Mst and its principal interactors, we compared the mitotic phenotypes elicited by depletion of either Mst, the *Drosophila* Prefoldin 3 subunit encoded by *merry-go-round* (*mgr*; [16, 17]), and the TCP-1 complex subunits. Available mutations in the genes encoding the TCP-1 complex subunits caused early embryonic lethality, preventing cytological analysis in larval brains. Similarly, in vivo RNAi, using UAS-RNAi lines and ubiquitous drivers, produced an early lethal phenotype. We thus generated flies carrying a suitable UAS-RNAi construct and the conditional tubGAL4–tubGAL80ts driver, permitting time-restricted expression of this construct.
Brain preparations from larvae grown for 72 hr at 29°C, expressing UAS-RNAi constructs against the TCP-1 complex subunits, were immunostained for Tubulin and the centrosomal marker DSpl-2 [18] and compared with mst and mgr mutant brain preparations. Consistent with previous observations, mst and mgr mutant brains displayed a metaphase arrest phenotype and frequent polyploid cells [5, 16, 17, 19]. In addition, most mst and mgr prometaphase/metaphase figures showed monopolar spindles with reduced MT density (Figures 2A and S2). Importantly, RNAi against Tcp-1 caused an ~80% reduction of the protein level and a mitotic phenotype indistinguishable from that elicited by mst mutations (Figures 2A–2C and S2). Moreover, RNAi against each of the other seven TCP-1 subunits also resulted in a very similar phenotype; we consistently observed high frequencies of monopolar spindles (ranging from 39.4% to 76.7%; n = 200 per each TCP-1 subunit) with strongly reduced MT density (Figures 2B, 2C, and S2).

To further investigate the functional relationship between Mst and the TCP-1 complex in spindle assembly, we analyzed MT regrowth after cold exposure in mitotic cells of TCP-1α-depleted brains. After cold-induced MT depolymerization, wild-type cells rapidly form new MTs, first from chromosomes and then from both chromosomes and centrosomes. The two MT populations merge within 5 min, giving rise to morphologically regular spindles (S3; Figure S3). We previously showed that mutations in mst strongly reduce chromosome-driven MT regrowth, having little effect on MT regrowth from centrosomes [6]. Similarly, in TCP-1α-depleted brains exposed to cold and then returned at room temperature (RT), the majority of mitotic cells showed initial MT regrowth exclusively from the centrosomes; only after 10 min recovery did these cells show monopolar and bipolar spindles similar to those of untreated cells (Figure S3). Thus, the absence/reduction of Mst and TCP-1α subunits similarly affect spindle MT generation in larval neuroblasts, suggesting a functional link between Mst and the Tubulin-folding machinery.

Mst Is Required for the Stability of the TCP-1 Complex and Efficient Tubulin Polymerization

We next sought to distinguish whether Mst is a substrate for the TCP-1 and Prefoldin complexes or whether it acts as a co-factor for the Tubulin-folding pathway. We first assessed the amount of Mst, Mgr, and TCP-1α in larval brain extracts depleted of each of these proteins. In extracts from RNAi larvae depleted of any TCP-1 subunit, the level of TCP-1α was always dramatically reduced (Figure 2C). These findings indicate that all TCP-1 complex components are required for TCP-1α stability and, consistent with previous results in mammalian cells [20], suggest that all TCP-1 subunits are mutually required for complex stability. However, their reduction did not affect the levels of either Mgr or Mst (Figure 2D). Remarkably, while levels of both Mst and TCP-1α were unaffected in brain extracts from mgr mutants, mutations in mst, although not affecting Mgr, caused a strong reduction in the levels of TCP-1α (Figure 2E). These findings were corroborated by immunolocalization experiments. Similar to Mst [5], TCP-1α specifically accumulated in the cytoplasm of mitotic cells of wild-type brains but was almost undetectable in dividing cells of mst mutant brains (Figure 2F).

To assess whether the low level of TCP-1α observed in mst mutants reflects a reduction of the entire TCP-1 complex, we subjected wild-type and mst mutant larval extracts to size exclusion chromatography, probing separated fractions for TCP-1α. The TCP-1α of mst mutants, although reduced in amount, eluted at ~500–550 kD (corresponding to the predicted size of the Drosophila TCP-1 hetero-octameric complex) with a profile identical to wild-type TCP-1α (Figure 2G), indicating that Mst is required to stabilize the entire complex.

The TCP-1 complex in other experimental systems is essential for the correct folding of both Tubulin and Actin. However, while RNAi-mediated depletion of human TCP-1 complex subunits leads to a dramatic reduction in Tubulin levels, it has little effect on Actin [21]. To determine whether Mst depletion differentially affects Tubulin and Actin in Drosophila, we analyzed levels of α- and β-Tubulin, γ-Tubulin, and Actin in wild-type and mst mutant extracts from either brains or whole larvae. Although Tubulin levels were similar in wild-type and mst mutant brains, mst larvae displayed substantially reduced levels of α- and β-Tubulin compared to wild-type larvae (Figure 2H); this difference likely reflects variable susceptibilities of different tissues to loss of components of the Tubulin-folding pathway, as previously reported for Drosophila Mgr [17]. In contrast, both wild-type and mst mutant brains and larvae showed very similar levels of both γ-Tubulin and Actin. Moreover, size exclusion chromatography of wild-type, mst mutant extracts, or extracts of Tcp1-α RNAi larvae failed to reveal any effect on Actin level or size distribution profile, while confirming the reduction of α-Tubulin (Figure 2G).

To further investigate the specific loss and functionality of α- and β-Tubulin in mst mutant larvae, we performed Tubulin stability and MT sedimentation assays; we could not perform these experiments with isolated brains due to the difficulty in collecting enough tissue. First, by monitoring the amounts of Tubulin and Actin in extracts incubated at 27°C for varying times, we found that, while α-Tubulin levels remained unchanged in wild-type extracts over the course of 1 hr, α-Tubulin in mst mutant extracts was rapidly lost (Figure 2I). Next, we subjected extracts from wild-type, mst, and Tcp1-α RNAi larvae to in vivo MT and Actin sedimentation assays. We found that Actin polymerization was unaffected in mst or Tcp1-α RNAI extracts, although a consistent band-shift, possibly reflective of post-translational modification, was observed (Figure 2J). To control for the reduced concentration of α-Tubulin in mst and Tcp1-α RNAI larval extracts (Figure 2G), we performed the MT sedimentation assay with various diluted wild-type extracts. Even in wild-type extracts diluted 1:16, a proportion of α-Tubulin was able to pellet under polymerization conditions. In contrast, α-Tubulin in mst mutant larval extracts, similar to that present in Tcp1-α RNAI larvae, remained in the supernatant, suggesting an inability to polymerize (Figure 2K).

The Role of Mst in Spindle MT Assembly

Our data demonstrate that the functions of Mst and the TCP-1 complex are integrally linked. Mst binds the complex and is required for its stability. Moreover, Mst loss causes a mutant phenotype indistinguishable from that elicited by loss of any TCP-1 subunit, and the biochemical properties of extracts depleted of either Mst or TCP1-α are identical. Together, these
Figure 2. Mst Is Required for TCP-1 Complex Stability

(A) mst and TCP-1α-depleted (Tcp-1α) neuroblast metaphases stained for Dspd-2 (red), α-Tubulin (green), and DNA (blue) exhibit monopolar spindles with low MT density. The scale bar represents 5 μm.

(B) Frequencies of monopolar spindles observed in larval brains after RNAi-mediated depletion of TCP-1 subunits (indicated with Greek letters from α to θ).

(C) Western blots of brain extracts depleted of TCP-1 subunits probed with anti-TCP-1α and anti-Lamin (Lam, loading control) antibodies. TCP-1α is reduced in all extracts.

(D) Western blot of brain extracts depleted of TCP-1 subunits, probed for Mst, Mgr, and Lam. Mst and Mgr levels are similar to wild-type (WT).

(E) Western blot from null mst and mgr mutant brains probed for Mst, TCP-1α, Mgr, and Giotto (Gio, loading control). TCP-1α levels are strongly reduced in mst extracts.

(F) Localization of TCP-1α and Mst in larval brains. In WT brains, TCP-1α and Mst specifically accumulate in mitotic cells; in mst mutants, the TCP-1α signal is almost undetectable. Arrows and arrowheads point to metaphases and anaphases, respectively. The scale bar represents 5 μm.

(G) Western blots of larval extracts, separated using size exclusion chromatography and probed for TCP-1α, α-Tubulin (α-Tub), and Actin (Act). Mst or TCP-1α loss reduces Tubulin, but not Actin, levels. Asterisk (*) indicates overexposed WB to show Tubulin.

(H) Western blots of larval brain and whole-larvae extracts showing total levels of α- and β-Tubulin, γ-Tubulin, and Actin in WT and mst mutants; α- and β-Tubulin are reduced in whole larvae, but not in brains.

(legend continued on next page)
The general secondary and tertiary structural elements found within both Tubulins and FtsZ map coherently onto Mst, strongly supporting the notion that Mst is a distant member of the Tubulin superfamily (Figure 3A). However, Mst contains additional stretches of amino acids that are predicted to be structurally disordered and that possibly affect both the nucleotide binding and oligomerization activities elicited by Tubulins (Figures 3A and S4). Upon superimposition of the Mst model onto the model of bovine Tubulin:TCP-1 complex [23], we found that Mst is capable of filling the internal cavity of the TCP-1 complex (Figure 3B). Although its additional loops exceed the cavity boundary, we hypothesize that the flexibility of these disordered protein stretches allows additional interactions with the TCP-1 complex. Therefore, although further work will be required to determine whether Mst does indeed sit within the TCP-1 complex cavity in vivo, our modeling is consistent with a scenario in which Mst stabilizes the TCP-1 complex in the absence of a substrate, through a Tubulin-like interaction.

Our study also highlights the existence of tissue-specific requirements for Tubulin folding and MT polymerization. We have shown that Mst associates with embryonic spindles, consistent with the report that Drosophila TCP-1 subunits co-sediment with MTs from early embryos [24]. However, Mst does not appear to be enriched in larval brain spindles ([5]; Figures 2F and S1). A possible explanation for this difference is that the rapid assembly of syncytial embryonic spindles [25] requires a high local concentration of assembly-competent Tubulin. A spindle-anchored folding machinery would, however, not be necessary in cells surrounded by a plasma membrane (such as brain cells), where sufficient folded Tubulin could be provided by increasing the intracellular concentration of the folding complexes in anticipation of mitosis.

Finally, the in vivo MT regrowth experiments presented here and previously [5] demonstrate that, in either mst mutant or Tcp-1 RNAi background, centrosome-driven MT regrowth after cold treatment is less affected than chromosome-/kinetochore-induced regrowth. The simplest explanation is that, following cold-induced depolymerization, kinetochores and centrosomes nucleate MTs with different dynamics. Cold treatment removes the vast majority of cellular MTs, probably leaving only short, cold-resistant MT seeds at centrosomes. While rewarming of wild-type cells essentially “reboots” spindle assembly pathways, allowing both polymerization from centrosomes and ab-initio nucleation/polymerization from kinetochores, the reduced pool of assembly-competent Tubulin present in mst or Tcp-1 mutant cells would be preferentially incorporated into the existing MT stubs at the centrosomes. An alternative hypothesis may reflect that the chromosome- and centrosome-driven MT formation pathways governing Drosophila cell division are at least in part under separate genetic control [5, 25, 26].

In summary, our study identifies Mst as a factor required for the stability of the TCP-1 complex, ultimately controlling the stability and polymerizing competency of α- and β-Tubulin.

Table 1. List of Mst-GFP Interacting Proteins

| Protein Name            | Percentage Coverage | Number of Peptides | MW (kDa) | Score   | Mean Area |
|-------------------------|---------------------|--------------------|----------|---------|-----------|
| TCP1-gamma              | 88.97               | 53                 | 59       | 2,924   | 2.47E10   |
| TCP1-zeta               | 82.74               | 46                 | 58       | 2,188   | 2.14E10   |
| TCP1-alpha              | 79.17               | 41                 | 60       | 2,146   | 1.76E10   |
| TCP1-beta               | 80.93               | 43                 | 58       | 2,017   | 1.58E10   |
| TCP1-eta                | 84.01               | 45                 | 59       | 1,877   | 2.03E10   |
| TCP1-delta              | 75.80               | 36                 | 57       | 1,818   | 1.55E10   |
| TCP1-Gamma              | 74.74               | 33                 | 65       | 1,598   | 2.26E10   |
| TCP1-theta              | 70.33               | 39                 | 59       | 1,367   | 1.93E10   |
| TCP1-epsilon            | 70.85               | 40                 | 59       | 1,345   | 1.18E10   |
| TCP1-theta              | 75.80               | 36                 | 57       | 1,818   | 1.55E10   |
| CG7770 (Prefoldin 6)    | 38.40               | 5                  | 14       | 77      | 6.60E8    |
| TCP1-Gamma              | 50.84               | 9                  | 16       | 75      | 5.14E8    |
| Prefoldin 2             | 20.29               | 2                  | 16       | 54      | 1.14E8    |
| Prefoldin 1             | 29.37               | 6                  | 15       | 52      | 5.26E8    |
| Prefoldin 1             | 48.33               | 11                 | 27       | 88      | 2.36E8    |
| Prefoldin 4             | 26.00               | 9                  | 67       | 67      | 1.51E8    |
| CHIP                    | 37.02               | 9                  | 34       | 65      | 1.43E8    |
| Prefoldin 5             | 57.89               | 6                  | 15       | 53      | 2.05E8    |
| Prefoldin 6             | 28.23               | 10                 | 51       | 51      | 1.38E8    |

Proteins identified via mass spectrometry isolated from 0–3 hr Mst-GFP-expressing Drosophila embryo extracts after stringent filtering (see Supplemental Experimental Procedures). The proteins shown have MS scores of >50 and coverage percentages of >20%, respectively. Mean area corresponds to Top 3 Protein Quantification (T3PQ), the mean of the three highest abundance peptides identified for each protein. Mst is identified with a similar score and mean area to all eight subunits of the TCP-1 complex. All subunits of the Prefoldin complex are also co-precipitated, albeit at approximately 10-fold-lower abundance. Similar profiles were obtained for the TCP-1 subunits in the other two experiments.

Results indicate that Mst directly regulates TCP-1 complex structure and function. Mst was initially described as having primary structural motifs similar to those found in the Tubulin superfamily members [19, 22]. We therefore sought to determine whether Mst could be structurally related to Tubulins at the tertiary level. To do this, we composed a three-dimensional model of Drosophila Mst. Briefly, we identified sequences homologous to Mst in the α-, β-, and γ-Tubulin and FtsZ family of proteins, structurally aligned the available homologous structures, and merged Mst to this alignment. This final alignment was then used for comparative modeling using the Rosetta software suite (Figures 3 and S4; Supplemental Experimental Procedures).

(1) Western blots of WT or mst larval extracts incubated at 27°C for 20, 40, or 60 min, probed for α-Tubulin and Actin. Tubulin is rapidly lost in mst extracts. 
(2) Western blot of an Actin sedimentation assay performed with WT, mst, and Tcp-1a RNAi larval extracts. Actin is able to polymerize in all extracts. 
(3) Western blot of a MT sedimentation assay. WT extracts were diluted 1:8 and 1:16 to control for reduced Tubulin levels in mst and Tcp-1a RNAi extracts. Tubulin is competent to polymerize in WT, but not in mst or Tcp-1a RNAi extracts.

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within the fly. Mst is a conserved protein; future work will clarify whether it is a TCP-1 complex cofactor with a mitotic role in humans.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, four figures, and three movies and can be found with this article online at http://dx.doi.org/10.1016/j.cub.2015.05.033.

AUTHOR CONTRIBUTIONS

V.P. performed most of the experiments. C.P. and K.J.H. performed the biochemical and MS analyses, respectively. V.M.-P. generated Mst-GFP. Modeling of Mst was undertaken by K.B.R. under the supervision of C.M.D. J.G.W. performed the size exclusion chromatography. V.P., S.B., M.G., and J.G.W. conceived the experiments, analyzed the data, and wrote the paper.

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