Article Addendum

Microtubule amplification in the assembly of mitotic spindle and the maturation of kinetochore fibers

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Key words: Plk1, FAM29A, NEDD1, mitotic spindle assembly, microtubule nucleation, microtubule amplification, k-fiber maturation

Efficient assembly of a mitotic spindle and stable attachment of microtubules (k-fibers) to kinetochores are essential for the high fidelity of chromosome segregation. Both spindle assembly and k-fiber formation require robust nucleation and polymerization of microtubules mediated by the γ-tubulin ring complex (γ-TuRC). It has been well established that centrosomes and chromatin are the two centers for microtubule nucleation. We recently demonstrate a third mechanism for microtubule nucleation and polymerization, in which the existing microtubules in the spindle act as templates to promote the formation of new microtubules. We showed that a novel spindle-associated protein, FAM29A, plays a critical role in this microtubule-dependent microtubule amplification. FAM29A associates with spindle microtubules and directly interacts with and recruits NEDD1, the targeting subunit of γ-TuRC. Spindle-associated γ-TuRC then promotes microtubule nucleation required for spindle assembly and k-fiber formation. This novel microtubule amplification pathway provides a powerful mechanism to control the local cytoskeleton structures independent of centrosomes and chromatin. We speculate that microtubule amplification not only functions in mitosis, but may also act in other physiological processes to re-enforce existing cytoskeleton structures.

Nucleation of microtubules (MTs) by the γ-tubulin ring complex (γ-TuRC) is the first step in the formation of the mitotic spindle.1 MT nucleation and polymerization is also required for the maintenance of the spindle structure. Loss-of-function in one of the γ-tubulin isoforms in mice generates highly disorganized spindle and abnormal spindle poles, resulting in a characteristic mitotic arrest.2 Similarly, knockdown of NEDD1, an accessory subunit of the γ-TuRC that targets γ-tubulin to various mitotic structures,3,4 reduces the density of spindle MTs and leads to a stable prometaphase arrest.4 Thus, efficient nucleation and proper regulation of MTs are essential for normal mitotic progression and for the high fidelity of chromosome segregation in mitosis.

γ-TuRC is localized to centrosomes, the main microtubule organizing center in eukaryotic cells.1,5 NEDD1 recruits γ-TuRC to centrosomes to initiate MT nucleation and polymerization.3,4 Although centrosomes increase the efficiency of MT nucleation, they are not required for mitosis, as a spindle can form in the absence of functional centrosomes.6 In mitosis, chromatin also promotes the polymerization of MTs through its associated guanine nucleotide exchange factor, RCC1, which generates a gradient of RanGTP around chromatin.7 The RanGTP gradient dissociates the spindle assembly factors (SAFs) from their interactions with inhibitory importins and free SAFs then promote MT growth around the chromatin.8-10 Lastly, both NEDD1 and γ-tubulin are localized to the mitotic spindle,4 suggesting that MTs may nucleate and polymerize from existing MTs in the spindle. We recently demonstrate that a novel spindle-associated protein, FAM29A, is required for targeting NEDD1 and γ-tubulin to the spindle and that FAM29A promotes the MT-dependent MT polymerization critical for the assembly of mitotic spindle and essential for the maturation of kinetochore MT fibers (k-fibers).11

We initially identified FAM29A as a protein interacting with the Polo-like kinase, Plk1, an essential mitotic kinase that controls spindle assembly and its bipolarity.12,13 We found that FAM29A is a MT-associated protein (MAP) and that its MAP activity is regulated in the cell cycle.11 It associates with MTs in vitro in the presence of mitotic extracts and colocalizes with MTs in vivo, but only during mitosis. In fact, in prometaphase and metaphase cells, FAM29A preferentially associates with cold-resistant stable k-fibers. FAM29A directly interacts with Plk1 and the active kinase of Plk1 targets FAM29A to the mitotic spindle. FAM29A also interacts with NEDD1, again only in mitosis. Through this direct interaction, spindle-associated FAM29A targets NEDD1 to the spindle and promotes the MT nucleation through NEDD1-associated γ-TuRC. Thus, FAM29A uses existing spindle MTs as templates to generate addition MTs, a phenomena we termed “MT amplification” (Fig. 1).

MT amplification is important to proper spindle assembly.11,14 Knockdown of FAM29A prevents the association of NEDD1 and γ-tubulin with the spindle and reduces the MT density by 60%.11 The weak bipolar spindle remained in the FAM29A-depleted cells
is not sufficient to efficiently capture the chromosomes and there is a delay in the progression through prometaphase. In mitotic cells that are released from an arrest by nocodazole, a MT destabilizing drug, the initial kinetics of MTs nucleation and polymerization from centrosomes and chromatin is independent of FAM29A. However, once the initial seeds of MTs are formed, the subsequent expansion of MTs is absent and the bipolar spindle fails to form in FAM29A-depleted cells. Thus, MT amplification provides sufficient amounts of MTs critical for spindle assembly.

MT amplification is also essential for the maturation of k-fibers. Upon entry into mitosis, individual MTs extend from centrosomes, search the three-dimensional cytoplasmic space for chromosomes, and capture the kinetochores. The dynamic turnover of kinetochore-attached MTs provides the driving force for chromosome congression and segregation. However, a single attached MT is not sufficient to drive the chromosome movement and a mature k-fiber usually consists of 25–30 MTs that act in concert. How a single attached MT quickly matures into a bundle of 25–30 MTs is a central mystery in cell biology. We showed that FAM29A-mediated MT amplification is essential for the maturation of k-fibers. Depletion of FAM29A in cell biology. We showed that FAM29A-mediated MT amplification is an evolutionarily conserved mechanism. For example, the Augmin complex recently identified in Drosophila has a subunit that shares a weak sequence homology to FAM29A and processes a similar biological activity. Furthermore, MT amplification is not unique to mitosis. Plant cells lack a MT organizing center, and \( \gamma \)-tubulin is associated with cortical MTs to promote the polymerization of MTs that branch from existing cortical MTs. Similarly, \( \gamma \)-tubulin in yeast is associated with interphase MTs and promotes the generation of new MTs to form MT bundles. We speculate that MT amplification also acts beyond the cell cycle in physiological processes in mammalian cells, such as in neurons, where MT amplification promotes the local control of cytoskeleton structure independent of centrosomes.

**Figure 1** FAM29A mediates MT amplification in spindle assembly and in k-fiber maturation. Active kinase of Plk1 targets FAM29A to spindle microtubules and FAM29A directly interacts with and recruits NEDD1, the targeting subunit of \( \gamma \)TuRC. \( \gamma \)TuRC then promotes microtubule nucleation required for spindle assembly and k-fiber formation. Drawings for various cellular structures are representative, but not to the scale.

**Figure 2** FAM29A controls the partition of NEDD1 between spindle MTs and centrosomes. The interaction between Plk1 and NEDD1 is responsible for Plk1-mediated recruitment of NEDD1 to centrosomes, which controls microtubule nucleation and polymerization from centrosomes (I). Recruitment of NEDD1 to the spindle depends on the interaction between Plk1 and FAM29A as well as the interaction between FAM29A and NEDD1, which mediate microtubule nucleation and polymerization from the mitotic spindle (II). The levels of FAM29A determine the partition of NEDD1 between centrosomes and the spindle, which, in turn, controls the relative contributions of MT nucleation and polymerization between these two mitotic structures (III).
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