Mitosis-specific kinesins in Arabidopsis

Marleen Vanstraelen¹, Dirk Inzé¹ and Danny Geelen²

¹Department of Plant Systems Biology, Flanders Interuniversity Institute for Biotechnology (VIB), Ghent University, Technologiepark 927, B-9052 Gent, Belgium
²Department of Plant Production, Faculty of Bioscience and Bio-engineering, Ghent University, Coupure links 653, B-9000 Gent, Belgium

Kinesins are a class of microtubule-associated proteins that possess a motor domain for binding to microtubules and, in general, allows movement along microtubules. In animal mitosis, they function in spindle formation, chromosome movement and in cytokinesis. In addition to the spindle, plants develop a preprophase band and a phragmoplast that might require multiple kinesins for construction and functioning. Indeed, several kinesins play a role in phragmoplast and cell plate dynamics. Surprisingly few kinesins have been associated with the spindle and the preprophase band. Analysis of expression datasets from synchronized cell cultures indicate that at least 23 kinesins are in some way implicated in mitosis-related processes. In this review, the function of kinesins in animal and plant mitoses are compared, and the divergence that originates from plant-specific aspects is highlighted.

Specialized kinesins for mitotic plant cells

During cell division, microtubules (MTs) arrange into dynamic structures that drive the different stages of mitosis. The dynamic organization of MTs requires the functional cooperation of different MT-associated proteins (MAPs) [1]. Kinesins use a motor to convert the energy of ATP into movement along the MT or to control the (de)polymerization of MTs [2]. In animal cells, the role of different kinesins in distinct mitotic stages is well documented. Even though kinesins are strongly represented in plant genomes, with the exception of the class kinesin 2 and 3 for which there are no homologs, only a few members have been implicated in spindle organization, chromosome movement and cytokinesis [3]. Plants have additional, strongly diverged kinesins that do not classify within the subfamilies that have been defined to date [4]. These kinesins could be involved in the organization of the unique mitotic MT arrays, the preprophase band (PPB) and the phragmoplast that occur in plants but are absent in other eukaryotes [1]. The completion of sequencing the Arabidopsis thaliana genome and the classification of kinesins into different functional subfamilies [5,6], along with genome-wide expression analyses [7,8], allows us to make predictions about kinesin functioning in dividing plant cells.

Arabidopsis kinesins up-regulated during M-phase

Progression through the cell cycle involves several hundreds of genes, which are regulated both at the transcriptional and post-translational level. Transcript profiling of synchronized Arabidopsis tissue culture cells has identified 23 kinesins that are up-regulated during mitosis [7,8] (Figure 1, Tables 1 and 2). The promoter sequence of the mitosis activation motif (MSA) is a plant-specific cis-acting element that drives periodic activation of gene expression at the start of mitosis or at the M phase of the cell cycle [9,10]. This cis element is strongly represented in the promoter sequences of the mitosis up-regulated kinesins (Tables 1 and 2). The kinesin NACK1 of tobacco (Nicotiana tabacum) contains two MSA sequences in the promoter region that are responsible for mitosis-dependent expression [11]. Accordingly, the NACK1 protein accumulates during M-phase, which is consistent with its function in cell plate expansion [12]. The MSA element is absent in the immediate upstream region of kinesins At4g21270, At4g05190 and At5g27550, suggesting that either the MSA sequence has degenerated beyond computer-assisted recognition or another mitosis-specific element exists for the expression of these particular kinesins.

Cell cycle-controlled phosphorylation of kinesins

Cyclin-dependent kinases (CDKs) regulate progression through the cell cycle by reversible phosphorylation of downstream effectors [13]. In animal cells, CDKA phos- phorylation regulates the MT-binding activity of kinesin members of the Kinesin-4, Kinesin-5, Kinesin-6 and Kinesin-7 subfamilies that are required for spindle functioning during mitosis [14–16]. CDKA1 phosphorylation sites are present in 14 out of the 23 kinesins that are mitotically up-regulated (Tables 1 and 2). The phosphorylation of Kinesin-5 (also known as BimC-type kinesins) Eg5 and KLP61F at a consensus CDKA phosphorylation site in the C-terminal BimC box is necessary to localize these proteins to the mitotic spindle in Xenopus laevis egg cells and Drosophila melanogaster embryos, respectively [17,18]. There are four Kinesin-5 homologs in Arabidopsis (i.e. AtKRP125a or At2g37420, AtKRP125b or At2g36200, AtKRP125c or At2g28620, and At3g45850) that, together with tobacco TKRP125, contain the conserved phosphorylation site embedded in their BimC box. AtKRP125a, AtKRP125b and AtKRP125c are up-regulated during mitosis whereas At3g45850 is not. By contrast, the Arabidopsis Kinesin-4 and Kinesin-7 family members lack a classic CDKA1 phosphorylation site (Table 1), suggesting that unlike their animal counterparts, they are not regulated by CDK phosphorylation. In addition to control by CDK-type kinases, Aurora kinases have also
been shown to play important roles in the regulation of the subcellular localization of Kinesin-6 kinesins through phosphorylation. Aurora kinases are present in plants and localize to the spindle and cell plate [19]. Because Kinesin-6 kinesins are absent in Arabidopsis, it remains to be seen whether Aurora kinases have a role in kinesin regulation. The recent finding that Aurora B also phosphorylates the Kinesin-5 BMK-1 in Caenorhabditis elegans has opened new avenues for investigation in plants [20]. Hitherto, there has been no direct evidence for CDK- or Aurora-dependent phosphorylation of kinesins. KCA, belonging to the Kinesin-14 group, has been analyzed in the most detail and has been shown to bind CDKA in two-hybrid assays and in vitro precipitation tests [21,22]. Furthermore, KCA is phosphorylated by an olomoucine-sensitive kinase, possibly a cell cycle-dependent kinase, in Sf9 insect cells [21]. Mutagenesis of two adjacent, putative CDK phosphorylation sites in the tail domain of KCA1 to glutamate codons, such that serine phosphorylation is mimicked, results in a failure to target the GFP-tagged protein correctly to the cell plate and the plasma membrane [23]. KCA–GFP with the same serines changed to non-phosphorable alanines accumulates in the cytoplasm, indicating that dephosphorylation is required for membrane localization. The KCA-related data are the first evidence for phosphorylation-controlled cellular distribution of kinesins in plants.

**Cell cycle control of the kinesin protein level**

For progression through the different steps of the mitotic phase, successive phosphorylation events are needed as well as the removal of proteins by proteolytic degradation. A well-studied event of collective removal of proteins takes place at anaphase during which key cell cycle regulators, such as cyclins and structural proteins, are rapidly degraded via a multistep process involving ubiquitination and subsequent targeting to the 26S proteasome. The targeting to the proteolytic complex relies on conserved domains, also known as destruction boxes, present in proteins that are targeted for proteolytic degradation [24]. The MAP Ase1 of yeast (Saccharomyces cerevisiae) carries a D-type destruction box that, when mutated, prevents proteolytic degradation and spindle disassembly [25]. A conserved D-box is also present in some members of the MAP65 protein family that show similarity to Ase1. For instance, the AtMAP65-4 disappears at the end of anaphase, suggesting that this protein might also be removed via the anaphase promoting complex (APC) [26]. One or more conserved D-boxes occur in 22 out of the 23 kinesins with mitosis-dependent expression (Tables 1 and 2). The strong conservation of the D-box supports a general role for protein degradation in controlling plant kinesin function. However, D-box-dependent or other proteolytic degradation of plant kinesins has not yet been demonstrated directly. The 26S proteasome inhibitor MG132 arrests mitotic cells at metaphase because of the activation of the spindle checkpoint mechanism [27]. MG132 also blocks the M phase, G1 phase transition whereby it protects the TKRP125 from degradation and provokes the formation of extra phragmoplasts [28].

For several plant kinesins, protein levels have been investigated throughout the cell cycle in synchronized Bright Yellow-2 (BY-2) cells [12,29–31]. The protein levels
of the kinesins ATK2, ATK3, TKRP125, NACK1 and KCBP of tobacco rise during M-phase and decline at mitotic exit. The *Arabidopsis* counterparts of ATK2, ATK3 and KCBP are not significantly up-regulated in microarrays of synchronized cells \[7,8\], suggesting that control over transcript and protein production is not a conserved mechanism in plant kinesin functioning.

### M-phase-specific kinesins related to animal kinesins

In animal cells, minus-end-directed kinesins are of major importance for the organization of the spindle poles (Figure 2). The founding member of this class, Ned, localizes to the spindle poles and cooperates with dynein to bundle MTs that are attached to the centrosome at the poles \[32\]. As a general rule, kinesins with a motor domain

<table>
<thead>
<tr>
<th>Kinesin subfamily</th>
<th>MiPs code</th>
<th>MSA elements</th>
<th>CDK sites</th>
<th>Destruction boxes</th>
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</table>

*Kinesins are organized per subfamily according to the nomenclature of Lawrence et al. \[6\]. Sequences of the mitosis activation motif (MSA) elements in a 1000-bp region upstream of the ATG start codon, CDKA phosphorylation sites (S/T-P-x-K/R) and destruction boxes (R-x2-L-x4-N) are shown.*
near the C-terminal end exhibit minus-end directionality. The Arabidopsis genome contains 21 C-terminal kinesins now classified as Kinesin-14, of which seven are transcriptional and up-regulated at mitosis [6,8]. The Ncd-related ATK1 and ATK5 have been shown to move in the minus-end direction of MTs and to associate with spindle MTs where they are required for focusing the MTs at the poles, which is similar to Ncd activity (Figure 2) [33–36]. Compared with other organisms, the Kinesin-14 group is relatively large. It has been postulated that these kinesins fulfill functions normally performed by cytoplasmic dynein [37]. However, some of the C-terminal kinesins have distinct properties by which they differentiate themselves from the dynein characteristics. For instance, MT-binding by kinesin KCBP is inhibited upon interaction with calcium-calmodulin [38–40]. Calcium levels rise during mitosis [41]. Yet, KCBP localizes to the spindle poles in Haemanthus endosperm, possibly through the interaction with other proteins that are components of the spindle pole complexes. Because of this localization at the poles, it has been proposed that KCBP bundles MT minus-ends in a similar way to Ncd (Figure 2) [42,43].

Kinesin-5 members antagonize the forces of the Kinesin-14 subfamily members to establish spindle bipolarity [2]. Kinesin-5 KLP61F of Drosophila and Eg5 of human are plus-end-directed bipolar homotetramers that cross-link and slide antiparallel MTs of the spindle midzone, thereby generating poleward forces. Failure of Kinesin-5 members to associate with the spindle results in the formation of monopolar spindles [17,44]. This phenotype can be partially rescued by mutations in the C-terminal kinesin Ncd, suggesting that a balance between forces directed towards plus-ends and minus-ends establishes the bipolar spindle [44]. A mechanism of balancing forces seems conserved because similar results were obtained in yeast and fungi [45,46]. Because the Arabidopsis AtKRP125 is predominantly concentrated at the spindle midzone, this Kinesin-5 probably acts in a similar way to BimC and antagonizes the force of the minus-end-directed kinesin ATK1 (Figure 2).

The plant spindle is formed in the absence of centrosomes. Consistently, the spindle poles are broader than those in animal cells [47]. Centrosomes are also absent in Drosophila oocytes [32,48]. Here, acentrosomal

Table 2. Mitotic kinesins specific to plants*

<table>
<thead>
<tr>
<th>Kinesin subfamily</th>
<th>MiPs code</th>
<th>MSA elements</th>
<th>CDK sites</th>
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</table>

*Arabidopsis* kinesins carrying a motor domain that are clustered together but do not have a close relative in other non-plant species were considered as specific [5]. Sequence analysis identified mitosis activation motif (MSA) elements in a 1000-bp region upstream of the ATG start codon, CDKA phosphorylation sites (S/T-P-x-K/R) and destruction boxes (R-x2-L-x4-N).
Figure 2. Kinesin motor proteins in plant and animal cell divisions. Overview of the different mitotic stages and the kinesin subfamilies that play a role in key events in mitotic progression. Many of the animal kinesins that are known to function in mitosis have been shown to be involved in multiple roles in different mitotic processes (see, for example, the Kinesin-4 and the depolymerizing Kinesin-13 subfamily). (a) During prophase in animal cells, centrosomes migrate to opposite sides of the nucleus, thereby constructing the spindle. This migration is triggered by the outward pulling forces of dynein at the cell cortex and by the antagonistic forces of Kinesin-14A (Ncd) acting on non-kinetochore MTs [81]. In plant cells, the MTs of the PPB encircle the prophase spindle, which is built and nucleated at the nuclear surface. Motor proteins involved in PPB or prophase spindle formation have not been identified to date. (b) During metaphase, sister chromatids face opposite poles of the bipolar spindle. Antagonistic forces of Kinesin-14, such as Ncd [81] and Arabidopsis ATK1 [35] and Kinesin-5, such as DmKLP61F [81], HsEg5 or AtTKRP125 establish spindle bipolarity, whereas Kinesin-14 members Ncd, ATK1 and ATK5 focus on the minus-ends of the spindle MTs [32–36]. Furthermore, members of the depolymerizing Kinesin-13 family (DmKLP10A and HsKIF2A) function in mitotic spindle assembly and maintenance [62]. Chromosomes are aligned by numerous kinesins in animal cells. Kinesin-4 members, such as Drosophila DmKLP3A or human and Xenopus KID, act on the chromosomal arms, whereas Kinesin-7 members (DmCENPmeta and HsCENP-E) and Kinesin-13 members (HsMCAK) function at the centromere [55–57,60,82]. (c) During anaphase A, chromosomes are separated to the spindle poles. The combined action of depolymerizing kinesins belonging to the Kinesin-13 family, which chew up MTs from the centromere (DmKLP9C and HsMCAK) and the centrosome (DmKLP10A or HsKIF2A), results in chromosome segregation [62]. (d) Later, during anaphase B, spindle poles are further separated. The Kinesin-5 motor DmKLP61F and dynein at the cortex drive spindle elongation, assisted by Kinesin-4 members. During late anaphase in animal cells, the cleavage furrow, which contains a contractile ring of actin–myosin, contracts and bundles the MTs into a central spindle. Both Kinesin-5 motors and Kinesin-6 motors function in the organization of the central spindle [65]. (e) During cytokinesis, the cleavage furrow further ingresses, thereby concentrating the central spindle into a compact midbody. The furrow is the site of cell plate formation. Phragmoplast formation is driven by Kinesins-5 and -14. Kinesin-7 and -14A are involved in chromosome alignment. Cell plate formation is driven by Kinesin-14A.
spindle assembly is accomplished in two steps: first, the spontaneous nucleation of MTs around condensed chromatin, followed by the sorting of randomly oriented MTs into a bipolar spindle involving the antagonizing action of Kinesin-5 and Kinesin-14 members [48]. The structural organization of the spindle MTs in plant cells starts before the nuclear envelope is broken down; therefore, it cannot rely directly on MT nucleation from chromatin. Nucleation seems to take place depending on γ-tubulin and Spc98p at the nuclear surface [49,50]. How spindle bipolarity is accomplished is not known. In the absence of ATK1, the spindle poles are wider than normal but still bipolar, indicating that minus-end-directed kinesins are needed merely to focus the MTs at the poles [34,36].

Plant Kinesin-5 members, such as TKRP125, not only associate with the spindle but also with the phragmoplast [30]. The phragmoplast MTs show a poleward translocation that is inhibited in the presence of TKRP125 antibody, in line with a role for TKRP125 in sliding MTs that are arranged in an antiparallel fashion. Recently, the tomographic analysis of phragmoplasts in whole-plant tissues of Arabidopsis has cast doubt on the commonly accepted idea that MTs interdigitate at the phragmoplast center [51]. If the MT plus ends are not overlapping, the activity of Kinesin-5 kinesins cannot operate as a result of the opposing phragmoplast MTs sliding. How then does the TKRP125 antibody inhibit outward translocation of phragmoplast MTs in permeabilized tobacco BY-2 cells? Perhaps the phragmoplasts in BY-2 cultured cells are organized differently to those in Arabidopsis and the MTs might temporally overlap, in a similar way to what occurs in miniphragmoplasts in endosperm tissue [52,53]. Alternatively, short stretches of MTs arranged in an antiparallel fashion occur near the plus-ends of phragmoplast MTs, which could be targeted by the TKRP125 kinesin. By pushing these short MTs against the centrally located cell plate via antiparallel sliding, the longer phragmoplast MTs would show an outward motion or a tread milling-type of translocation [30].

After nuclear envelope breakdown, pairs of sister chromatids associate with the spindle and oscillate until they are bi-oriented (attached to both spindle poles) and aligned at the metaphase plate [54]. For chromosome alignment, kinesins from different subfamilies must act cooperatively, generating poleward or antipoleward forces. Members of the Kinesin-4, Kinesin-7 and Kinesin-13 subfamily contribute to chromosome alignment [55–57]. Chromosome oscillations until metaphase alignment, which probably depend on the activity of kinesins, have been observed in plants [58].

Immunodepletion of Kinesin-4 Xkid, also known as chromokinesin, in frog egg extracts results in chromosome misalignment at metaphase [59]. Kinesin-4 members also function in chromosome condensation, segregation, spindle organization and cytokinesis as shown for Drosophila KLP3A (Figure 2) [60]. Arabidopsis has three kinesins with high levels of similarity to chromokinesins, one of which is up-regulated during mitosis (Table 1). The AtFRA1 homolog is not up-regulated during cell division and does not appear to be implicated in cell division. Indeed, its role is more consistent with the orientation of cellulose microfibrils in cell walls along with the MTs in the cortical array [61]. These findings underline the risk of drawing conclusions about the function of kinesins based on the extrapolation of amino acid conservation of the motor domain.

Kinesin-13 family members (also designated MCAK or Kin-1 in mammals) depolymerize MTs and contribute to multiple aspects of chromosome movement [62]: they localize to the centromeres, centrosomes and spindle midzone during mitosis; depletion or disruption of MCAK activity has shown that they function in spindle maintenance, chromosome alignment and segregation; and they control the proper attachment of MTs to sister chromatids [57]. Three Kinesin-13 members are present in mammals (Kif2A, Kif2B and MCAK) and in Drosophila (Klp10A, Klp59C and Klp59D) [6]. MTs from the centrosome and the centromere are depolymerized by Kif2A and Klp10A and by MCAK and Klp59C, respectively (Figure 2). Arabidopsis contains two members of the Kinesin-13 subfamily, AtKinesin-13A and AtKinesin-13B [3]. AtKinesin-13B is controlled by mitosis and Atkinesin-13A is not, but they both lack a Lys-rich neck motif that is commonly found in other Kinesin-13 members, implying functional divergence from their animal counterparts [3]. Indeed, Atkinesin-13A localizes to Golgi stacks and contributes to Golgi distribution in Arabidopsis trichomes [63].

Members of the Kinesin-7 subfamily control the interaction between the MT plus-ends of the spindle and the kinetochores of the chromosomes, both during metaphase alignment and anaphase segregation (Figure 2) [55,64]. Two Kinesin-7 kinesins are present in the Arabidopsis genome, the expression of one of which is controlled by mitosis (Table 1) [6].

The homology-based survey indicates that kinesins with putative functions related to chromosome movements are represented in Arabidopsis as multiple copies, of which a subset is mitotically expressed. This does not exclude the constitutively active copies from involvement in chromosomal movement because transcript levels do not always reflect protein accumulation, and kinesin function might also be controlled at the protein level.

M-phase-specific kinesins unique to plants

Animal cytokinesis is accomplished by an actin-myosin ring emerging at the cell equator to recruit components necessary for ingressing the plasma membrane. As a result, interpolar MTs from the anaphase spindle are bundled into a central spindle that is further compressed into a compact midbody at the end of telophase (Figure 2) [65]. Plants construct a phragmoplast consisting of antiparallel MTs and perform cytokinesis without constriction. The MTs guide the Golgi-derived vesicles to the midline where they fuse to form a cell plate (Figure 2) [66]. Thus, animal cells depend on a mechanism requiring contraction whereas plant cells use a mechanism involving centrifugal expansion [65]. Despite this mechanistic difference, the central spindle in animal cells has properties similar to those of the phragmoplast in plant cells [67]. During furrow progression, membrane vesicles accumulate at the leading edge of the cleavage furrow, which is
blocked upon drug-stimulated MT depolymerization [68,69]. Furthermore, Brefeldin A, an inhibitor of vesicle traffic machinery, suppresses the terminal phase of cytokinesis in C. elegans and is a potent blocker of cell plate formation [70,71]. These findings support possible analogies in the roles that MT-based motor proteins play during animal and plant cytokinesis.

Human Kinesin-6 Rab6-KIFL, which is related to MKLP1, accumulates at the central spindle and becomes highly concentrated in the midbody where it might deliver vesicles from the Golgi apparatus to the site of membrane fusion [72,73]. Arabidopsis does not possess Kinesin-6 family members, suggesting that other kinesins execute the transport of Golgi-derived vesicles to the equator. Of the 29 kinesins that do not belong to any of the established kinesin subfamilies and are unique to plants [6], nine are up-regulated during mitosis and are probably implicated in plant-specific MT-related processes associated with the PPB or the phragmoplast. The plant-specific kinesins that are known to function in phragmoplast dynamics are indeed induced mitotically (Figure 1, Tables 1 and 2).

AtPAKRP1 and the related AtPAKRP1L associate with the MTs of the cylinder-like and ring-like phragmoplasts and might maintain the bipolar structure of the phragmoplast once Kinesin-5 motors have established bipolarity [74,75]. Single mutants do not reveal any phenotype, perhaps because of mutual compensation for their respective loss of function [75]. AtPAKRP2, which is present in the Arabidopsis genome as a single gene, is not related to any other kinesin; it associates with the phragmoplast in a punctuate pattern and strongly accumulates at the cell plate in a Brefeldin A-sensitive manner [76]. Therefore, to date, AtPAKRP2 is the best candidate to deliver Golgi-derived vesicles to the phragmoplast midline. NACK1 (also named HIK [77]) and NACK2 (also named STUD [78] or TES [79]) play a role in phragmoplast MT dynamics [80]. NACK1 is essential for the completion of somatic cytokinesis and functions in the reorganization of MTs during the lateral expansion of the cell plate [12,77]; NACK2 is the closest homolog of NACK1 and takes part in the assembly of the radial MT array in male meiotic cytokinesis [78,79]. Both NACK1 and NACK2 interact with NPK1, a MAPKKK that controls phragmoplast expansion and cytokinesis [12]. Four mitotic up-regulated plant-specific kinesins that do not belong to the classic kinesin subfamilies are as yet uncharacterized (At3g44050, At3g17360, At3g20150, At5g23910; Table 2). Investigation of the function of these kinesins would be interesting to unravel plant-specific aspects of spindle and phragmoplast formation.

**Concluding remarks**

The combined effort of genome sequencing and genomewide transcription profiling in Arabidopsis has identified 23 mitotic kinesins that are categorized into different structural and functional subfamilies. These kinesins are the prime candidate motors to function in spindle organization and dynamics, chromosome movement, and cytokinesis. In addition to the transcriptionally activated kinesins, other kinesins contribute to mitosis without cell cycle-modulated RNA or protein levels. The scarceness of functional evidence for kinesins in spindle formation is striking. Indeed, most mitosis up-regulated kinesins have been assigned functions related to cytokinesis. No kinesins that have a role in chromosome movement have yet been identified and limited information is available on the role of kinesins in spindle dynamics. Sequence comparison suggests that the role of kinesins is only partially conserved in different aspects of chromosome alignment and segregation. Moreover, some homologs of animal kinesins with functions in chromosome movement are unrelated to plant cells and even seem to be important in processes that take place during interphase. Further functional divergence is expected because plant mitotic cytoskeletal structures display unique features. For instance, plants lack centrosomes and might use specially adapted kinesins to facilitate stabilization and condensation of MTs at the spindle poles. In addition, land plants form a PPB for which no counterpart exists in other eukaryotes. Until now, there is no evidence that kinesins contribute to the formation or dynamics of the PPB. Yet, in view of the significant number of plant-specific kinesins that have not been studied yet, they should be at the center of future research.

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