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Received 20 Sep 2012 | Accepted 4 Apr 2013 | Published 14 May 2013

DOI: 10.1038/ncomms2831

A protein phosphatase 2A complex spatially controls plant cell division

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In the absence of cell migration, the orientation of cell divisions is crucial for body plan determination in plants. The position of the division plane in plant cells is set up premitotically via a transient cytoskeletal array, the preprophase band, which precisely delineates the cortical plane of division. Here we describe a protein complex that targets protein phosphatase 2A activity to microtubules, regulating the transition from the interphase to the premitotic microtubule array. This complex, which comprises TONNEAU1 and a PP2A heterotrimeric holoenzyme with FASS as regulatory subunit, is recruited to the cytoskeleton via the TONNEAU1-recruiting motif family of proteins. Despite the acentrosomal nature of plant cells, all members of this complex share similarity with animal centrosomal proteins involved in ciliary and centriolar/centrosomal functions, revealing an evolutionary link between the cortical cytoskeleton of plant cells and microtubule organizers in other eukaryotes.

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Fine-tuning of the spatial positioning of division planes is crucial for coordinating multicellular development in plants^{1,2}. Cells of land plants are glued together by a pecto-cellulosic cell wall, and do not have the ability to migrate. Their location is set up irreversibly by oriented cell divisions occurring in specialized histogenic tissues, the meristems. Consequently, the three-dimensional cellular organization of a plant tissue closely reflects the pattern of cell division and cell expansion that has occurred during its development.

In contrast to animal cells, plant cells establish their plane of division early during the cell cycle, and one of the first conspicuous signs of commitment to division is the transition of the cortical microtubular array into a dense preprophase band (PPB) encircling the nucleus³. The PPB precisely forecasts the cortical region reached by the growing cell plate during cytokinesis. It is thus supposed to leave a positional cue at the cell cortex, which persists there until cytokinesis^{2,4}. At the end of prophase, the PPB disappears concomitantly with nuclear envelope breakdown and spindle formation. In telophase, the new cell plate separating the daughter cells is deposited through vesicle transport guided by a mitotic cytoskeletal array composed of actin filaments and microtubules, the phragmoplast, which initiates at the centre of the cell, and grows centrifugally until it connects with the plasma membrane at the cortical site previously occupied by the PPB. Thereafter, microtubules re-colonize the cell cortex and organize into parallel arrays transverse to the cell elongation axis, guiding deposition of cellulose microfibrils on the outer side of the cell^{5,6}. Unlike many eukaryotes, cells of land plants lack a conspicuous microtubule organizing centre (MTOC) such as a centrosome, with the exception of basal bodies formed briefly in flagellate motile sperm cells of land plants that rely on aqueous fertilization (mosses and ferns). Siphonogamous Angiosperms and Pinophytes are devoid of centrosomes and centrioles at any stage.

Few genes involved in PPB formation have been identified⁷. Among them, *TON1* and *FASS/TON2* are the only ones whose disruption leads to a complete absence of PPB formation. *Arabidopsis* mutants in either *TON1* or *FASS* display identical phenotypes^{8–12}: plants are dwarf and misshapen, display abnormal cell elongation and random positioning of cell division planes^{8,9}. In addition to absence of PPBs in premitotic cells, interphase microtubule arrays are also affected, and lack the typical parallel organization of wild-type cells^{9–11}. In maize, simultaneous loss of function of the two *FASS* homologues is embryo lethal, while *TON1* disruption in moss phenocopies the developmental syndrome of *Arabidopsis ton1* mutants^{13,14}.

The *TON1* N-terminus shares sequence similarity with the animal centrosomal proteins FOP and OFD1 and interact with *Arabidopsis* CENTRIN¹¹, which in animal cells is a major constituent of MTOCs. *TON1* has recently been shown to interact with *TON1*-recruiting motif proteins (TRMs)¹⁵, which possess several motifs also present in the human centrosomal protein CAP350, responsible for FOP recruitment at the centrosome¹⁶. In addition, TRM1 was shown to target *TON1* to the cortical microtubules¹⁵. *FASS* encodes a putative protein phosphatase 2A (PP2A) regulatory B' subunit^{10,12} similar to RSA-1, a *Caenorhabditis elegans* centrosomal protein controlling microtubule outgrowth during spindle formation¹⁷. PP2As are major eukaryotic phosphatases involved in nearly every cellular process¹⁸. They comprise a catalytic C subunit, a scaffolding A subunit and a regulatory B subunit, which controls substrate selectivity and subcellular localization of the enzyme¹⁸. *FASS* directly interacts with the *Arabidopsis* PP2AA1/RCN1 subunit¹⁰, suggesting that it is a genuine subunit of a PP2A complex.

Here, we identify a regulatory complex composed of *TON1*, TRM and a PP2A holoenzyme containing *FASS* as regulatory

subunit using a proteomics approach. We show that the activity of this complex, named TTP (for *TON1*/TRM/PP2A), is required for PPB formation and spatial control of cell division in land plants.

Results

TON1 and FASS form a phosphatase 2A complex in *Arabidopsis*. To gain insight into the protein network involving *TON1* and *FASS*, we performed a series of tandem affinity purifications (TAP) from *Arabidopsis* suspension cells, using either *TON1* (isoforms *TON1a* and *TON1b*) or *FASS* as baits, fused to the GS-tag^{19,20}. On one hand, complex purifications were performed on cell extracts made in homogenization buffer (HB) with a standard low detergent concentration (0.1% (v/v) Nonidet P-40 (NP-40)) for extraction of water-soluble proteins. On the other hand, complex purifications were performed on extracts made in HB with high detergent concentration (1% (v/v) digitonin) for extraction of both water-soluble and membrane-associated proteins. Both extraction procedures gave comparable results. Moreover, in a total of 30 TAP purifications, only 17 different proteins were specifically and repeatedly co-purified (Fig. 1 and Table 1), indicating the consistency of the TAP approach. Using *FASS* as bait, we repeatedly identified not only *TON1a* and *TON1b* but also PP2A catalytic subunits (PP2AC3/C4), and all three PP2A-scaffolding subunits present in *Arabidopsis* (PP2AA1/RCN1, PP2AA2 and PP2AA3) (Fig. 1, Table 1, Supplementary Fig. S1 and Supplementary Tables S1–S3), showing that *FASS* is part of a PP2A complex *in vivo*. Conversely, these PP2A subunits, including *FASS*, were retrieved with both *TON1a* and *TON1b* as baits, confirming that all these proteins are core components of a multi-protein complex (Fig. 1 and Table 1). Interestingly, TAP experiments with *TON1a* identified *TON1b* and *vice versa*, indicating that these proteins likely form heterodimers, presumably via their LisH dimerization domain^{11,21}.

Furthermore, six members from the recently identified TRM protein superfamily¹⁵ were retrieved using *TON1a*, *TON1b* and *FASS* as baits (Fig. 1 and Table 1). Reverse-TAP using the TRM19 GS-tagged protein confirmed co-purification with *TON1a* and *b*, *FASS* and other PP2A subunits (Fig. 1 and Table 1), thereby linking the *TON1*/*FASS*-PP2A complex to the TRM family of proteins.

Reverse transcriptase PCR (RT-PCR) analyses and available microarray data showed that all five PP2A catalytic subunits are expressed at comparable levels in our suspension cells (Supplementary Fig. S1a,b), yet we identified only the closely related isoforms PP2AC3 and C4 in TAP experiments. This likely reflects their preferential incorporation into the TTP complex. On the contrary, over-representation of TRM19 with respect to other TRMs coincides with a high expression level in our cell culture (Supplementary Fig. S1b). However, we also retrieved much lower expressors, such as TRM4 or TRM6, confirming that co-purification of a specific isoform does not merely reflect its abundance in our cell system.

These results suggest the existence of a high combinatorial diversity of TTP complexes: apart from *FASS*, all other identified components come into several isoforms, including *TON1* (a and b), PP2AC (C3 and C4), PP2AA (A1, A2, A3), as well as a large variety of TRM family members.

Several PP2AA and PP2AC isoforms contribute to TTP activity. The *Arabidopsis* genome encodes three PP2AA-scaffolding subunits. PP2AA1 is considered to have a cardinal role in seedlings. Although morphologically similar to the wild type, *PP2AA1* mutants display abnormal ethylene and ABA

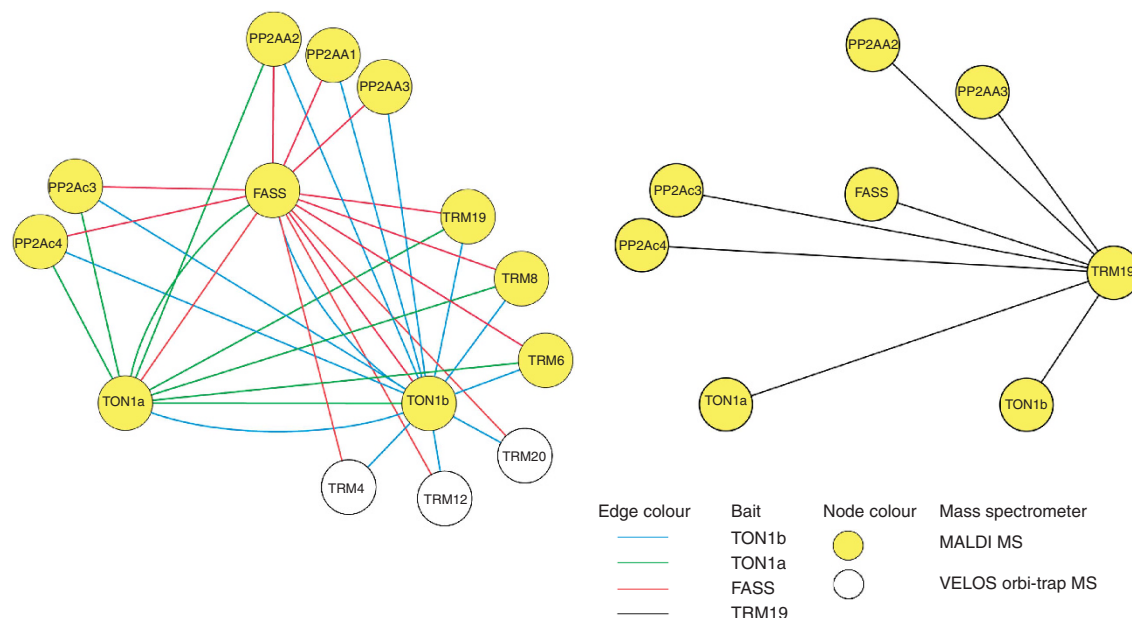


Figure 1 | TON1 and FASS form a phosphatase 2A complex. TTP protein-protein interaction networks on the basis of the TAP experiments. The left part corresponds to TAP results using TON1a, TON1b or FASS as bait, and the right network corresponds to reverse-TAP experiments using TRM19 as bait protein. Each line represents an interaction in TAP and colours indicate the tagged protein used as bait. White nodes represent the additional interactions identified using the more sensitive VELOS orbitrap analysis.

responses, as well as perturbed polar auxin transport^{22–25}. *PP2AA2* and *PP2AA3* functions are unmasked only in a *pp2aa1* mutant background²⁵ and neither *pp2aa2* and *pp2aa3* mutant nor *pp2aa2-a3* double mutant plants have altered morphological phenotypes *per se* (see ref. 25 and Fig. 2a). To analyse genetic interactions between *PP2AA* genes and *FASS*, we crossed a weak allele of *fass* (*fass-14*) with various *pp2aa* mutants. Introducing a *pp2aa1* mutation in a weak *fass* background resulted in a much stronger *fass* phenotype, with a clear gene dosage effect: in a *fass-14* background, a heterozygous *pp2aa1* mutation significantly enhanced the severity of the *fass* phenotype, and at the homozygous state the weak *fass* phenotype was fully converted to a loss-of-function *fass* phenotype (Fig. 2b). Although less marked, a similar enhancer effect was noted when *pp2aa2* and *pp2aa3* mutations were introduced in a *fass-14* background, confirming that all three *PP2AA* subunits can associate with *FASS* *in vivo* (Fig. 2c).

Likewise, to investigate genetic interactions between *FASS* and catalytic subunits isolated in our TAP experiments, we introduced *pp2ac3* and *pp2ac4* mutations into a weak *fass-15* mutant background (Fig. 2d). The *PP2AC3* mutation is expected to be leaky since residual *PP2AC3* transcripts are still detectable. Single *pp2ac3* or *pp2ac4* mutants do not display significant morphological alterations, but *pp2ac3-c4* double mutant are dwarfed with thick leaves, flowering stems and siliques (Fig. 2a and Supplementary Fig. S2). When introduced into a *fass-15* background, the *pp2ac4* mutation induced a strong reduction in root growth as compared with the *fass-15* parental seedlings (Fig. 2d). *pp2ac3-c4 fass-15* triple mutants are even more severely affected and strongly resemble a *fass* null allele.

Depleting scaffolding and catalytic *PP2A* subunits thus enhance a weak *fass* phenotype, confirming our physical interaction data, and providing genetic evidence that *FASS* is part of a *bona fide* *PP2A* complex *in planta*, requiring A1 A2 A3, C3 C4 for full activity.

A network of binary interactions establishes the TTP complex.

The co-purification of *FASS*, *TON1*, *PP2A* and *TRM* indicated that these proteins are bridged *in vivo* by physical contacts strong enough to survive the purification procedure. We set up yeast two-hybrid and/or Bimolecular Fluorescence complementation (BiFC) experiments to clarify binary interactions occurring within this large protein complex. *TON1* was previously shown to interact with *TRM* proteins *in vivo* via the M2 C-terminal motif of the *TRMs*¹⁵. BiFC assays revealed that *TON1* and *FASS* interact in *Nicotiana* cells (Fig. 3b). Yeast two-hybrid assays localized this interaction to N-terminal parts of both proteins (Fig. 3a). Furthermore, *FASS* and some *TRMs* are also capable of interaction: from a two-hybrid screen using *FASS* as bait, we indeed recovered two *TRM* clones. One encoded the full-length *TRM29*, while the other corresponded to the C-terminal part of *TRM3* (Fig. 3c). To identify the regions responsible for interaction with *FASS*, we performed two-hybrid assays between *FASS* and a series of *TRM1* deletion fragments. These experiments identified a 110-residue fragment (587–698) sufficient for interaction with *FASS* (Fig. 3c). As *TRM1*^{587–698}, *TRM29* and *TRM3*^{645–978} all have the M3 motif in common, the latter likely represents the *TRM*–*FASS* interaction region (Fig. 3c).

Together with previously published data¹⁵, this demonstrates that triangular interactions between *TRM*, *FASS* and *TON1* occur within the TTP complex (Fig. 3d), the M2 motif being responsible for the *TRM*–*TON1* interaction, whereas the M3 motif is required for the *TRM*–*FASS* interaction. *FASS* itself is able to form a heterotrimeric *PP2A* enzyme with A- and C-type subunits. The core TTP is thus composed of at least five protein components.

TRMs recruit TTP proteins to microtubule arrays. *TRMs* are a large and diverse superfamily of proteins, in which about half of the members are expected to localize to microtubule arrays¹⁵. We showed previously that *TRM1* localized to cortical microtubules

Table 1 | Summary of tandem affinity purification experiments.

Prey	MALDI														VELOS				Grand total (30)			
	TRM19		TON1a				TON1b				FASS				TON1a	TON1b	FASS					
	No dig	TRM19	No dig		With dig		TON1a	No dig		With dig		TON1b	No dig		With dig		FASS	MALDI		No dig		
	C (2)	total (2)	C (2)	N (2)	C (2)	N (2)	total (8)	C (3)	N (2)	C (2)	N (2)	total (9)	C (2)	N (2)	C (2)	N (2)	total (8)	total (27)	C (1)	C (1)		
TON1a (At3g55000)	2	2	Bait	Bait	Bait	Bait	Bait	3	2	2	2	9	2	2	2	2	8	19	Bait	1	1	21
TON1b (At3g55005)	2	2	2	2	2	2	8	Bait	Bait	Bait	Bait	Bait	2	2	2	2	8	18	1	Bait	1	20
TON2/FASS (At5g18580)	2	2	2	2	2	2	8	3	2	2	2	9	Bait	Bait	Bait	Bait	Bait	19	1	1	Bait	21
PP2AA1 (At1g25490)															1		1	1		1		2
PP2AA2 (At3g25800)	2	2	1			2	3	2	2	2	1	7			1		1	13	1	1	1	16
PP2AA3 (At1g13320)	1	1						1				1			1		1	3		1	1	5
PP2AC3/PP2AC4 (At3g58500/At2g42500)	2	2	1			2	3	3		1		4	1	2	2	2	7	16	1	1	1	19
TRM4 (At1g74160)																				1	1	2
TRM6 (At3g05750)															2		2	2	1	1	1	5
TRM8 (At5g26910)															1		1	1	1	1	1	4
TRM12 (At1g63670)																				1	1	2
TRM19 (At3g53540)	Bait	Bait								2		2	1	2	2	5	7		1	1	1	10
TRM20 (At4g28760)																				1	1	2
HAP 13 (At1g60780)										2	2							2				2
MAP1A (At2g45240)								1				1	1		2	2	5	6	1		1	8
IBR3 (At3g06810)						1	1	1				1						2		1		3
At5g57580																			1	1		2
TAP experiments were performed using TON1a, TON1b, FASS and TRM19 as bait proteins expressed in <i>Arabidopsis</i> suspension cell cultures. Baits were expressed either as N-terminal (N) or C-terminal fusion (C) with the GS-tag. The proteins identified in TAP eluates are indicated in the left column. Complex purification conditions (with or without digitonin in buffers) and the mass spectrometer used (MALDI MS/MS or VELOS orbitrap) are indicated. The number of independent TAP experiments analysed is given for each bait protein (between brackets). The VELOS orbitrap experiments concern additional analysis performed on eluate fractions obtained from either TON1a, TON1b or FASS TAP experiments. The numbers in the table correspond to the number of times the protein was co-purified with the bait protein. Note that PP2AC3 and C4 differ by only six residues and give undistinguishable mass profiles. The four last proteins of the prey list, although identified multiple times in our TAP experiments, were not further characterized in this study.																						

and is able to recruit TON1 to microtubules¹⁵. We therefore tested whether TRM1, an archetypal TRM member, could recruit other TTP components to microtubule arrays by transient co-expression of TTP proteins in *Nicotiana* leaf cells. The cytosolic GFP-FASS fluorescence (Fig. 3b) relocalized to the cortical microtubules (Fig. 4d) upon co-expression with RFP-TRM1. In addition, the BiFC signal of YFP^N-FASS and YFP^C-TON1a constructs also relocalized from the cytoplasm (Fig. 4a) to the cortical microtubule array where it co-localized with RFP-TRM1 (Fig. 4e) upon co-expression of the latter. TRM1 was also able to recruit the PP2AA1 subunit complexed with FASS to the microtubules (Fig. 4f). Hence, TRM proteins have the ability to recruit not only FASS and TON1 but also other PP2A subunits to microtubule arrays through their interaction with FASS. TRM8 also had the ability to recruit the TON1-FASS BiFC fluorescence to microtubule arrays (Fig. 4g), suggesting that microtubule recruitment of TTP proteins is a general function of microtubule-associated TRM proteins.

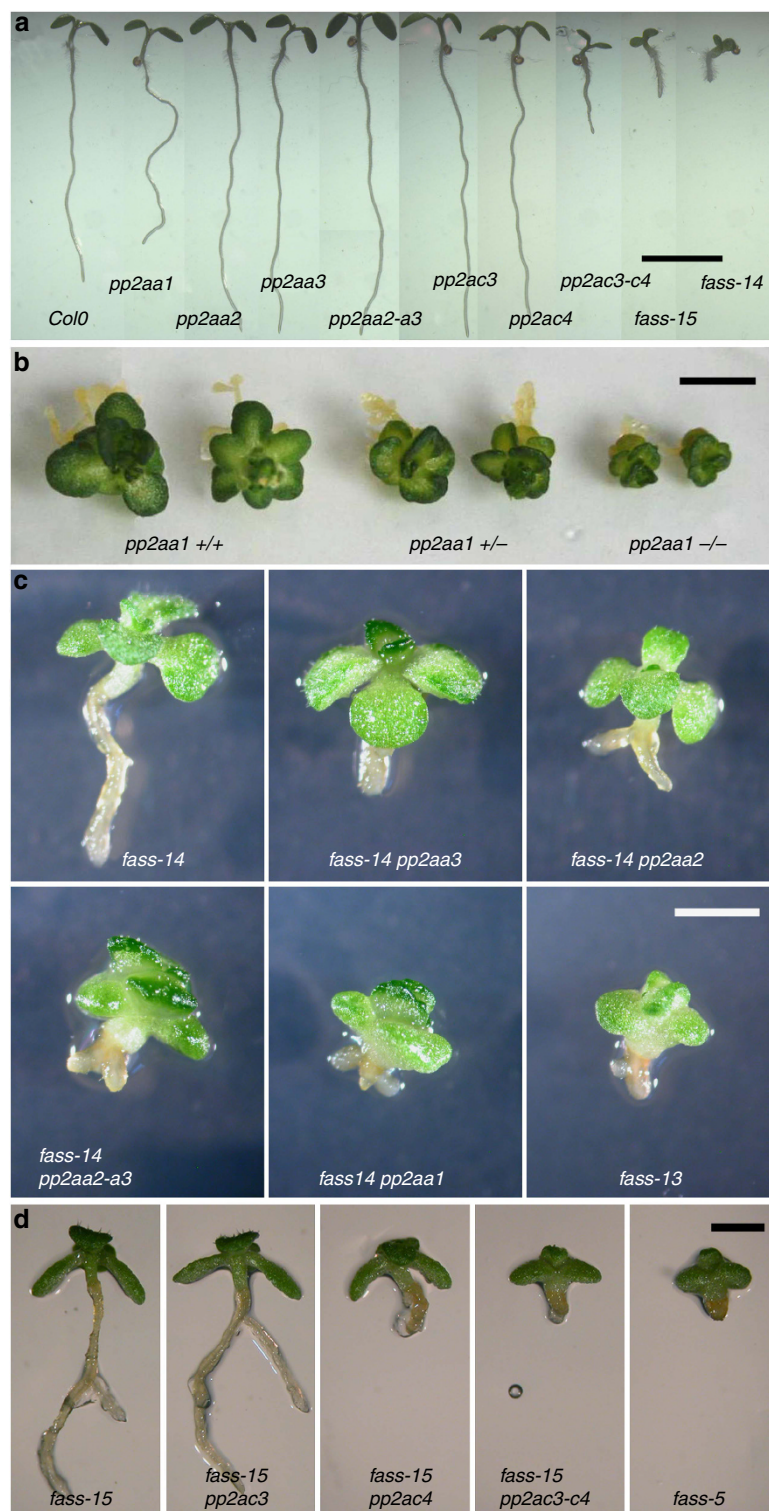


Figure 2 | Mutations in PP2AA and PP2AC subunits enhance the phenotype of weak *fass* alleles. (a) Photographs of all crossing partners 6 days after germination. From left to right: wild type (Col0), *pp2aa* (*pp2aa1*, *pp2aa2*, *pp2aa3*, *pp2aa2-a3*), *pp2ac* (*pp2ac3*, *pp2ac4*, *pp2ac3-c4*) and *fass* (*fass-14*, *fass-15*) mutants. (b) Dose-dependent enhancement of the *fass* phenotype upon introducing a PP2AA1 mutation into a weak *fass-14* mutant background: *fass-14* seedlings heterozygous for the PP2AA1 mutation (middle) already exhibit an enhanced *fass* phenotype compared with the *fass-14* line (left), whereas *pp2aa1* homozygotes (right) show an even more severe *fass* phenotype. (c) Genetic interaction between FASS and PP2AA subunits. Combining PP2AA mutations with the *fass-14* allele enhanced the *fass* phenotype, independent of the A subunit considered. The strongest enhancer effect is obtained with the *pp2aa1* mutation, as *pp2aa1 fass-14* double mutant plantlets are indistinguishable from *fass* null ones (*fass-13*). Mutating either *pp2aa2*, *pp2aa3* or both significantly increased the severity of the phenotype, although no full conversion to a loss-of-function phenotype was seen in this case. (d) Genetic interaction between FASS and PP2AC subunits. Introducing either the *pp2ac3*, *pp2ac4* mutations or both into the *fass-15* weak allele enhanced the *fass* phenotype as well, the *pp2ac3-c4 fass-15* triple mutant being indistinguishable from the *fass-5* null allele. Scale bars, 1 cm (a); 2 mm (b-d).

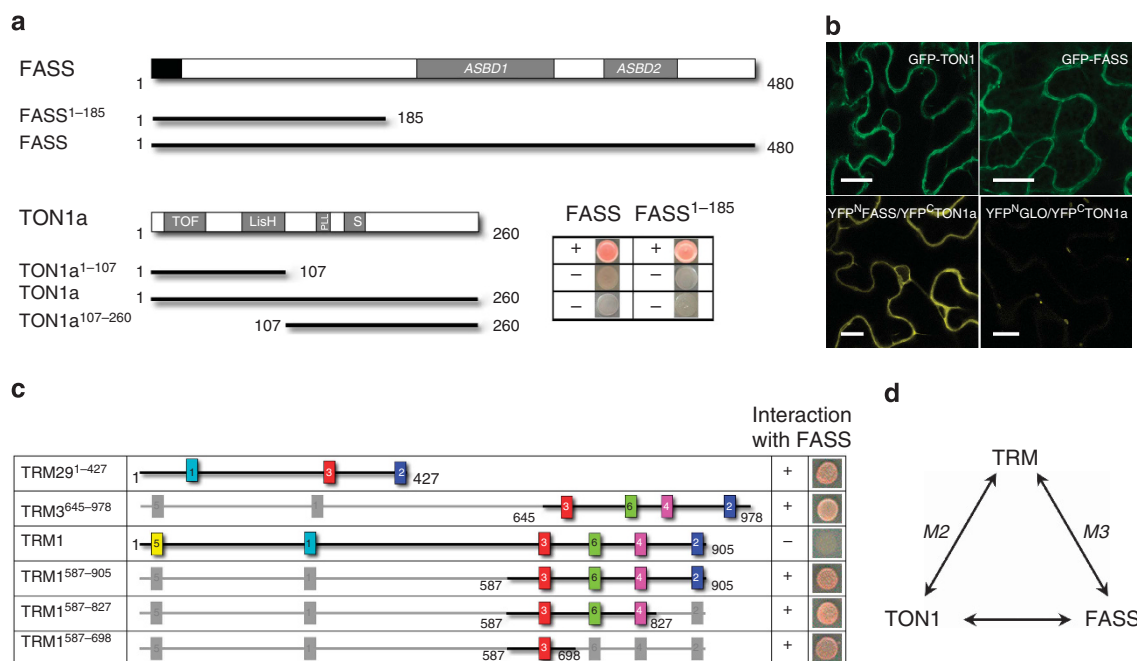


Figure 3 | FASS interacts with both TON1 and TRMs. (a,b) Interaction between TON1 and FASS. FASS interacts with TON1 in yeast two-hybrid experiments **(a)** and BiFC experiments in *Nicotiana benthamiana* abaxial epidermal leaf cells **(b)**. **(a)** Full-length, N- or C-terminal fragments of FASS and TON1 were tested for interaction in yeast two-hybrid. (+) and (-) indicates positive and negative results. Absence of interaction between TON1 full-length protein and FASS might result from improper folding in yeast or from the full-length protein being locked into a 'closed' conformation, masking potential interaction domains. Interactions with FASS¹⁸⁶⁻⁴⁸⁰ were meaningless due to strong self-activation of this construct. The position of the two A subunit-binding domains of FASS (ASBD1 and ASBD2) and the TOF, LisH-PLL, and the serine-rich (S) domains of TON1 are indicated. TON1 fragments were fused to the GAL4 activation domain and FASS to the LEX-binding domain. **(b)** Transient expression assays in *Nicotiana benthamiana* leaf epidermal cells. When overexpressed in this system, GFP-TON1 and GFP-FASS accumulate in the cytoplasm, which is restricted to the cell's periphery in these epidermal jigsaw puzzle cells. Co-expression of BiFC constructs YFP^N-FASS and YFP^C-TON1 gave a clear cytoplasmic YFP fluorescence signal, revealing interaction between FASS and TON1. Negative controls correspond to co-expression of YFP^C-TON1 with the unrelated YFP^N-GLOBOSA protein¹¹ (YFP^N-GLO). Scale bars, 20µm. **(c)** The M3 motif of TRMs is involved in direct interaction between TRMs and FASS. A schematic representation of TRMs fragments tested for interaction with FASS in yeast two-hybrid is shown. Coloured boxes correspond to the position of the motifs present in the TRM proteins. (+) and (-) indicates positive and negative results. **(d)** Schematic view of the interactions between TON1, FASS and TRM proteins. The M2 and M3 motifs of TRMs are involved in interaction with TON1 and FASS, respectively.

TON1 and FASS together with PP2AA1 decorate the PPB. *Arabidopsis* TON1 and FASS proteins have been shown previously to localize to the PPB^{11,12}. In addition to a PPB localization, DCD1 and ADD1, the two maize homologues of FASS, persist at the cortical division site after PPB disassembly¹³. To see whether this occurs in *Arabidopsis* for TON1 and FASS, we studied the dynamic localization of TON1 and FASS during mitosis, using functional GFP-TON1 (ref. 11) and GFP-FASS fusions (Supplementary Fig. S3) in root tip cells co-expressing an H2AX-RFP histone marker allowing to precisely define the mitotic phases²⁶. Time-lapse imaging of dividing cells showed that both GFP-TON1 and GFP-FASS labelled the PPB and persisted at the cortex up to metaphase, well after PPB disassembly (Fig. 5a,b). We could also show the presence of PP2AA subunits at the PPB by introducing a YFP-PP2AA1 construct²⁷ in an mCherry-β-tubulin6 background²⁸. In root tip cells, YFP-PP2AA1 displayed a cytoplasmic and perinuclear localization as reported before²⁷. In addition to this ubiquitous subcellular localization, YFP-PP2AA1 fluorescence distribution along the cortex in cells at the preprophase stage, showed a twofold enrichment at the PPB site (Fig. 5c), suggesting association of the PP2AA1 subunit with this structure.

TTP mutants are impaired in division plane positioning. Young plantlets of TTP mutant lines exhibit a gradation of a common developmental syndrome, from the strong *ton1/fass*

loss-of-function phenotype to less-affected *fass* weak mutants, *pp2aa1-a2* or *pp2aa1-a3* double mutants (Fig. 6a), and to a lesser extent *pp2ac3-c4* seedlings, likely due to leakiness of the available *pp2ac3* allele (Supplementary Fig. S2).

To trace back the origin of these alterations, we followed the cell division pattern in early embryos of TTP mutants. Early embryogenesis in *Arabidopsis* involves a well-characterized and invariant pattern of cell divisions (Fig. 6b). *ton1* and *fass* embryos displayed abnormal division planes in the suspensor and in the embryo proper as early as the two-celled stage, as reported earlier for *fass*⁸ (Fig. 6c). Imaging of double *pp2aa1-a2*, *pp2aa1-a3* or *pp2ac3-c4* mutant embryos also revealed notable defects in division plane orientation (Fig. 6c). Even *pp2aa1* single mutant embryos displayed abnormal division planes (Fig. 6c and Supplementary Fig. S4). This contrasts with the overall normal morphology of *pp2aa1* plants post germination (Supplementary Fig. S5) and suggests that the two other PP2AA subunits might compensate for *PP2AA1* disruption at later stages of development. Abnormal division plane positioning is thus a shared syndrome of *ton1*, *fass*, *pp2aa* and *pp2ac* mutants, further linking these genes in a functional network required for division plane positioning in plant cells.

TTP activity is required for preprophase band formation. *ton1* and *fass* mutants are defective in PPB formation⁹⁻¹¹. To search for similar defects in *pp2aa* or *pp2ac* mutant cells, we performed

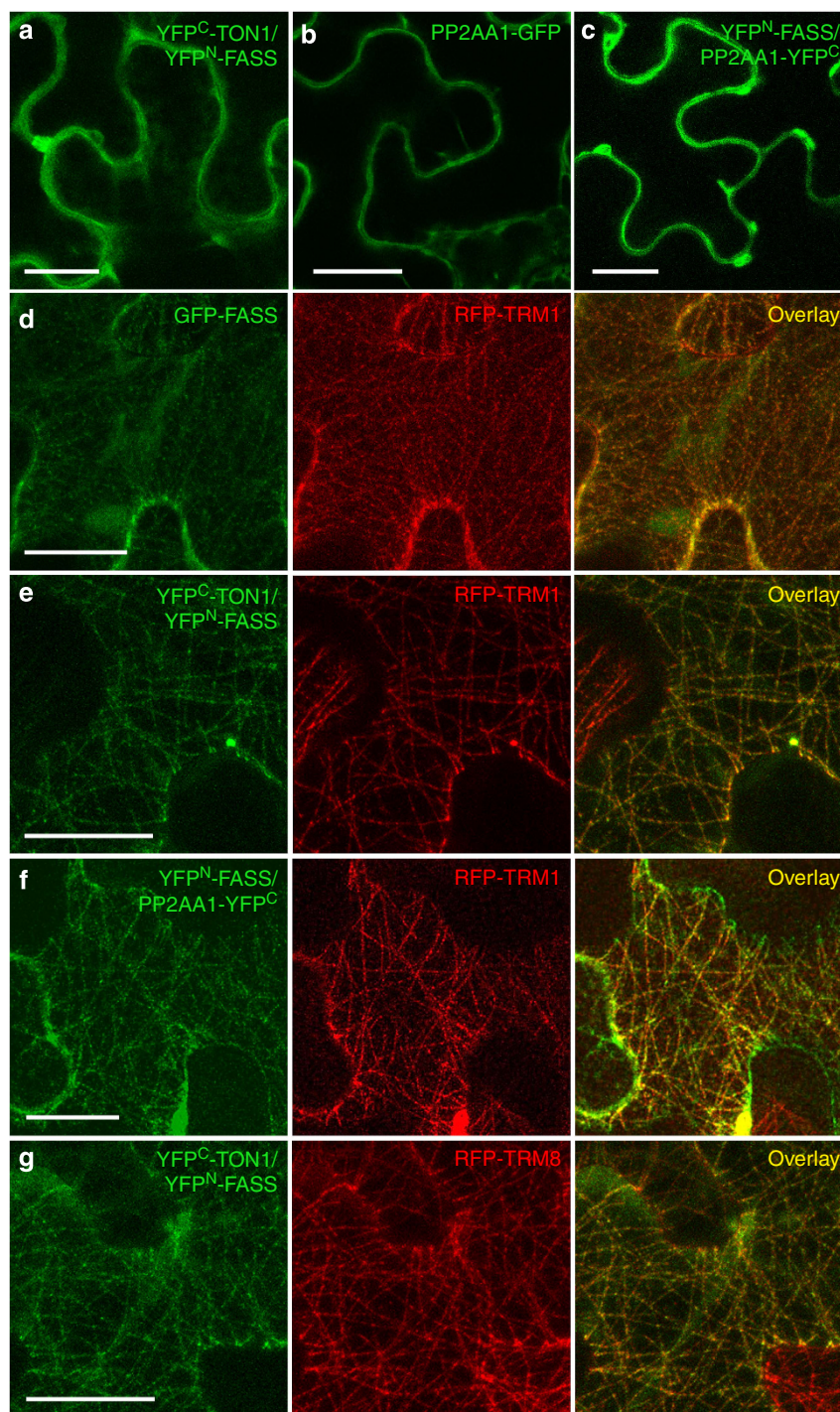


Figure 4 | TRM proteins target TTP proteins to the microtubular cytoskeleton. (a–c) TON1-FASS (a) and FASS-PP2AA1 (c) BiFC signals, as well as PP2AA1-GFP (b) fluorescence all localized to the cytoplasm when expressed in *N. benthamiana* leaf epidermal cells. (d–f) Upon co-expression with RFP-TRM1, which labels the cortical microtubular cytoskeletal array (middle panels, red), the GFP-FASS signal (d), the BiFC TON1-FASS fluorescence (e) or the BiFC FASS-PP2AA1 fluorescence (f) relocalized to microtubule arrays where it co-localized as a punctuate staining with TRM1. (g) The BiFC TON1-FASS fluorescence also relocalized to microtubule arrays upon co-expression with RFP-TRM8. All micrographs are projections of Z-stack confocal images. Scale bars, 20 μ m (a–g).

immunolocalization of microtubules in dividing root cells. Counter-staining with an anti-KNOLLE antibody allowed to identify cells in early mitosis, as this mitotic marker accumulates at the trans-Golgi network at G2/M^{29,30} (Fig. 7). At the preprophase stage, in wild type, 96% of KNOLLE-positive cells displayed a PPB, whereas only 4% did not ($n = 71$). Strikingly, in *pp2aa1-a2* or *pp2aa1-a3* double mutant cells, no PPBs were observed. In *pp2ac3-ac4* double

mutants, a majority of cells at the G2/M transition had no discernable PPB (79%, $n = 34$). At the premitotic stage, *pp2aa* or *pp2ac* double mutants showed an increased accumulation of perinuclear microtubules, as already noted for *ton1* and *fass* mutants^{9–11}. In contrast, all tested mutants formed spindles and phragmoplasts with frequencies and appearance comparable to wild-type cells (Fig. 7e and Supplementary Fig. S6).

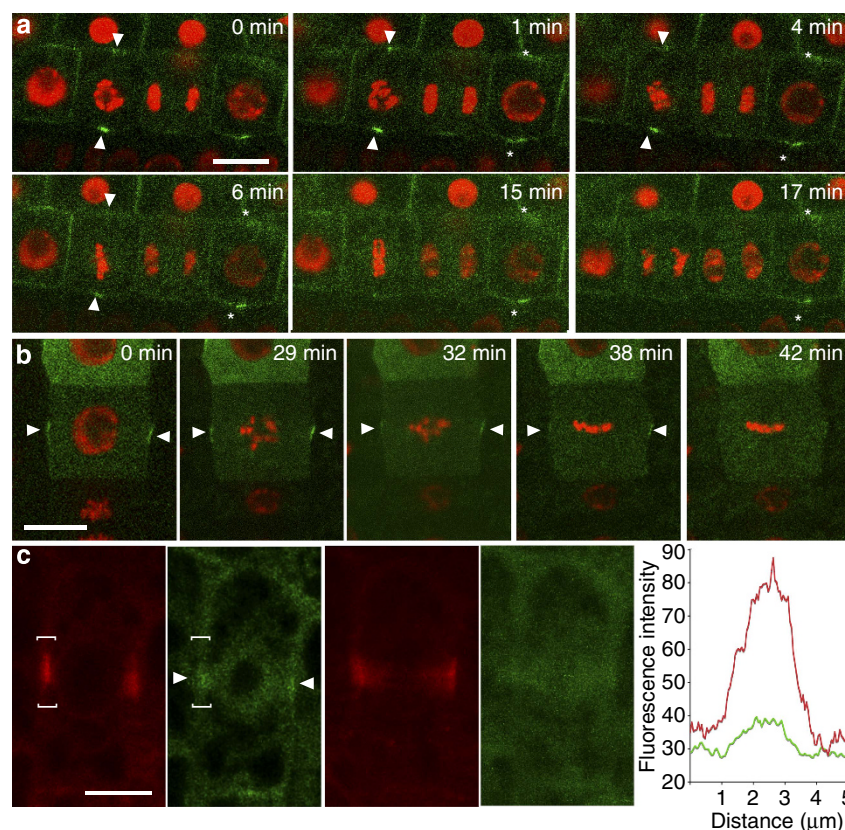


Figure 5 | TON1 and FASS and PP2AA1 all localize to the PPB. (a) Time-lapse analysis of root tip cells expressing both Pro35S:GFP-FASS and H2AX-RFP showed accumulation of FASS at the cortical division zone during PPB formation, and persistence of the signal beyond PPB degradation up to the metaphase stage. H2AX-RFP served as marker for the different cell-cycle phases. Arrowheads indicate PPB and cortical division zone localization of FASS up to metaphase in cell #1. Note persistence of the signal in cell #2 at $t = 17$ min (asterisks), excluding photobleaching-based disappearance of GFP-FASS signal in cell #1. (b) Time-lapse analysis of a root tip cell expressing both Pro35S:GFP-TON1 and H2AX-RFP, showing accumulation of TON1 at the PPB and persistence of the signal at the cortical division zone up to metaphase well after PPB disassembly. (c) Transverse (left) and tangential (right) views of a root tip cell at the preprophase stage expressing ProPP2AA1:YFP-PP2AA1 (green)²⁷ and mCherry- β -tubulin6 (red). YFP-PP2AA1 is enriched at the PPB (arrowheads) as also evidenced by the measurement of the fluorescence intensity along the plasma membrane indicated by brackets in transverse views (green curve corresponds to the YFP-PP2AA1 fluorescence and red curve to the mCherry- β -tubulin6 signal). Scale bars, 10 μ m (a–c).

In conclusion, defects in PPB formation observed in *ton1*, *fass* and various combinations of PP2A structural and catalytic subunits demonstrate that the TTP-driven PP2A activity is required for PPB formation at the G2/M transition.

Discussion

Since the initial discovery of the PPB almost 50 years ago³¹, numerous studies have confirmed its key role in division plane determination in land plant cells^{3,7}. However, molecular events contributing to the formation of this fascinating structure are just beginning to be uncovered. TON1 and FASS are among the few identified players in this process, their depletion disorganizing cortical microtubular arrays and preventing PPB formation^{8–12}. The similar *fass* and *ton1* mutant phenotypes already for long suggested a common mode of action^{8–11}, recently corroborated by the genetic interaction between *TON1a* and *FASS*¹². In this work, we identified a multi-protein complex including TON1, a PP2A heterotrimeric holoenzyme containing FASS as a regulatory subunit, and the TRM proteins. Proteomics, genetics as well as *in vivo* interaction and recruitment studies contribute to providing a remarkably consistent picture of TTP composition and function in PPB formation. Within the complex, FASS establishes contacts with both TON1 and TRM, which themselves physically interact, in a triangular relationship. These five

partners robustly co-purify together and constitute the core of the TTP complex. Co-expression experiments strongly suggest that this complex is recruited to microtubules via the TRM proteins. On the basis of available data, a tentative model emerges, where the PP2A part would represent the enzymatic activity, TON1 an assembly helper and/or activator, and TRM a recruiting agent bringing the PP2A activity into the vicinity of its target(s). Several recent works describe such adaptor proteins targeting PP2A activity to a particular location in the cell (the centrosome¹⁷ and the centromere³²) and reveal that PP2A's specificity is not determined solely by its regulatory B subunits. Therefore, assembly and activation of the whole complex at the proper time and subcellular location establishes specific phosphatase activity playing on the phosphorylation status of protein(s) essential for cortical microtubule dynamics and cytoskeletal transitions.

The stoichiometry and dynamics of assembly of core TTP components remains to be established. Our current data show that the core TTP is composed of at least five protein components, for a total MW of ~ 300 kDa, as also indicated by Blue NativePAGE analysis on cell extracts from cultures expressing GS-tagged TON1a and TON1b (Supplementary Fig. S1d,e). However, the diversity of subunits detected in our experiments argues for a combinatorial diversity of TTP isoforms. We provide evidence that all three A-type, but presumably only

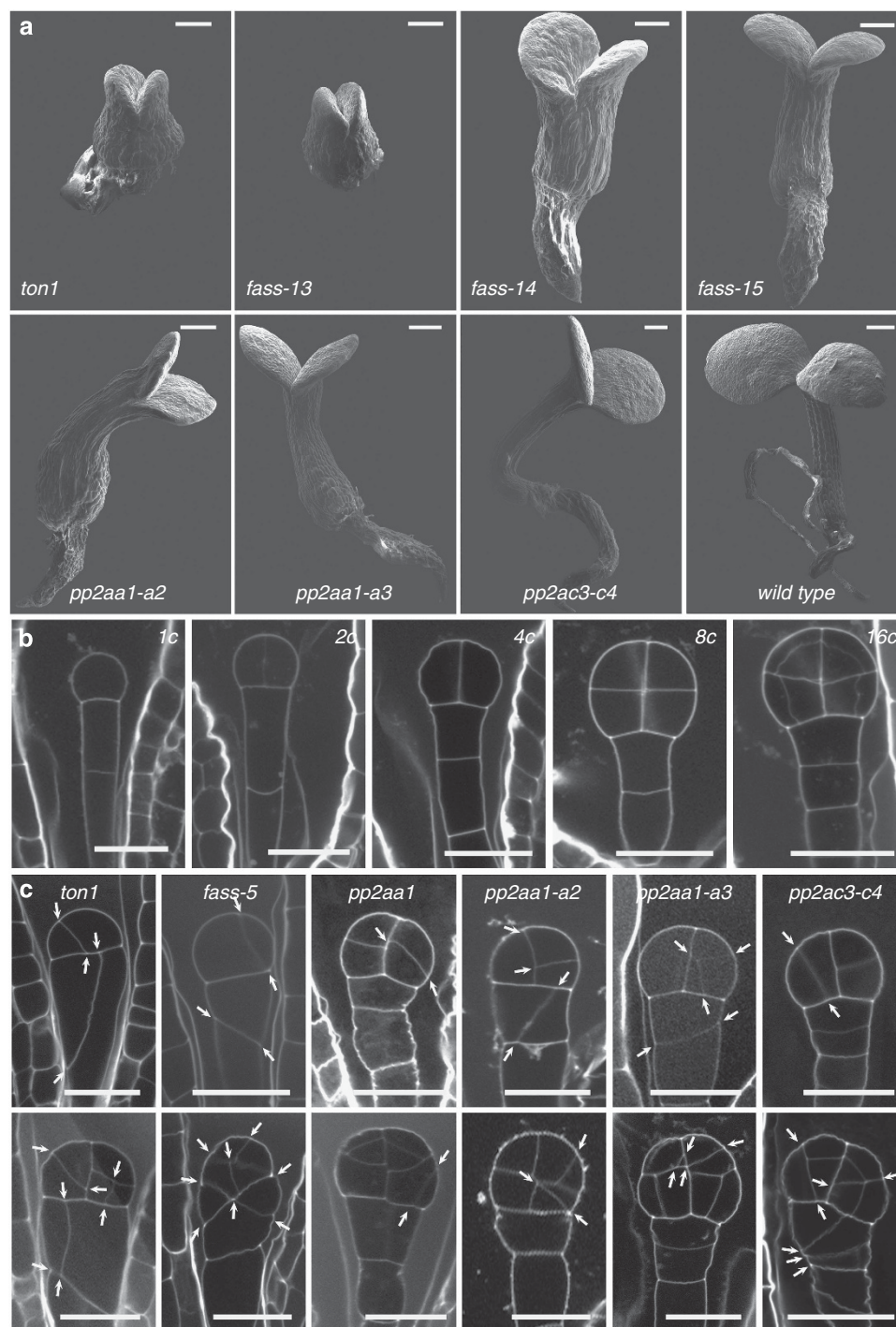


Figure 6 | TTP mutants share a common developmental syndrome. (a) Scanning electron microscopy images of 3-day-old seedlings of wild type and *ton1*, *fass* (*fass-13*, strong allele; *fass-14* and *fass-15* weak alleles), *pp2aa*, and *pp2ac* mutants. (b) Cleavage pattern in wild-type embryos from the single-celled (1c) to the 16-celled stage embryo (16c). The first division of the apical cell of the embryo is longitudinal, whereas the basal cell divides by transverse divisions leading to the formation of the suspensor. The two-celled embryo then divides longitudinally again, perpendicularly to the previous plane. The four resulting cells then divide by transverse divisions, giving rise to the eight-celled embryo. The next round of division establishes the protoderm cell layer by an asymmetric cleavage parallel to the surface, to form the 16-celled embryo. (c) Division plane defects in *ton1*, *fass-5*, *pp2aa1*, *pp2aa1-a2*, *pp2aa1-a3*, *pp2ac3-c4* mutant embryos. Two representative embryos are shown for each genotype. Abnormal division plane positioning in mutants occurs both in the suspensor and in the embryo proper (arrows). Scale bars, 250 μ m (a); 20 μ m (b,c).

two (PP2AC3 and PP2AC4 subunits) of the five *Arabidopsis* C-type PP2A subunits take part in the complex, the three other PP2AC subunits indeed representing a distinct clade among the five PP2AC subunits³³. This is further corroborated by the recently published analysis of a complete knock-out double

pp2ac3-c4 mutant whose phenotype is strikingly reminiscent of *ton1* or *fass* mutants³⁴. TON1 also comes into two isoforms, and recent data suggest that TON1a and TON1b are not strictly functionally redundant¹². The most diverse TTP component is the TRM family, counting 34 members in *Arabidopsis*. TRMs are

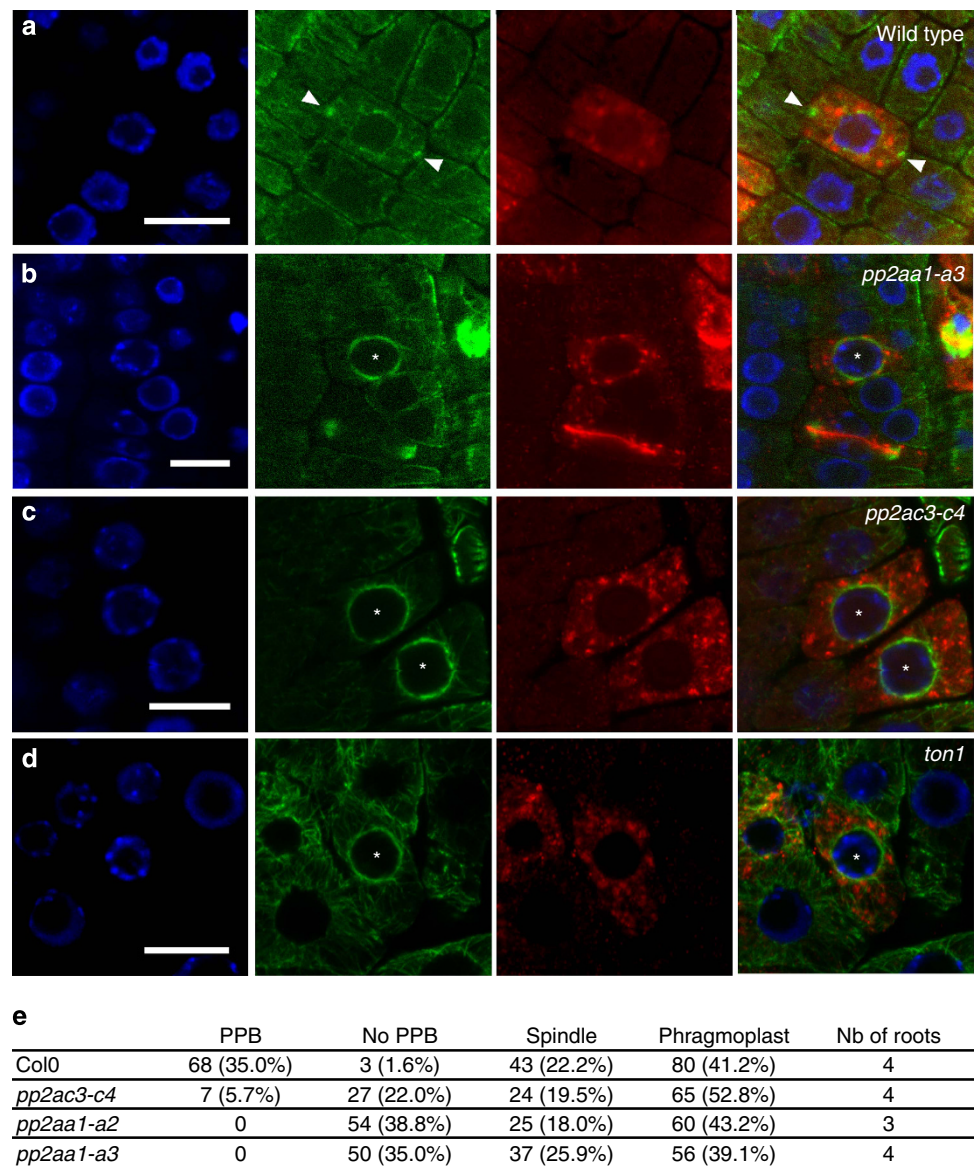


Figure 7 | *pp2aa* and *pp2ac* mutants are impaired in PPB formation. (a–c) Immunolocalizations on root tip cells of 4-day-old seedlings using anti-tubulin (green), DAPI staining (blue) and anti-KNOLLE (red). (a) Wild-type cell at the preprophase stage (PPB marked by the arrowheads) showing KNOLLE accumulation at the trans-Golgi network. (b–d) Absence of PPBs in *pp2aa1-a3* (b) and *pp2ac3-c4* (c) double mutants, and in the *ton1* mutant (d). Instead, dense perinuclear microtubules are visible at this stage (asterisks point out the nuclei). Scale bars, 10 μm (a–d). (e) PPB defects in *pp2aa* and *pp2ac* mutants. Mitotic stages in KNOLLE-positive cells were assessed using the anti-tubulin antibody. The number and percentage of KNOLLE-positive cells with spindles, phragmoplasts, PPBs or without PPB are indicated as well as the number of roots observed.

able to interact with both TON1 and FASS, thanks to their M2 and M3 motif, respectively. All TRM are predicted to possess a M2 motif, but 11 TRMs do not possess a predicted M3 motif, including TRM6 and TRM8. Yet, both are retrieved in TAP experiments using FASS as bait, suggesting that interaction with TON1 alone might be sufficient for incorporation into the TTP complex, or that the TTP complex can simultaneously contain several TRMs. In total, our TAP and yeast two-hybrid experiments identified half of the TRM family members (17 out of 34) as interactors of TON1 and/or FASS (Supplementary Fig. S7). This makes it likely that all TRM proteins are to some extent capable of contributing to a TON1–TRM–PP2A(FASS) complex. Given their variability in sequence, expression profiles and localization¹⁵, TRMs are likely a key element for targeting TTP activity to specific cell types and/or subcellular localizations.

TRMs are not likely amenable to mutant analysis due to the number of family members suggestive of functional redundancy, but for all other TTP components tested, including TON1, FASS, PP2AA and PP2AC, complete or partial depletion leads to defects in PPB formation and misorientation of division planes. Up to now, *TON1* and *FASS* were the only genes whose mutations induced complete absence of PPBs^{8–12}, and both proteins colocalize at the PPB and cortical division zone. Here we show that genetic depletion of scaffolding and catalytic PP2A subunits also induces prominent defects in PPB formation and division plane set-up. Although a general reduction in PP2A activity is expected to have pleiotropic effects, it is likely that the PP2A activity specifically associated to the TTP complex is causal to the observed defects. What could be the substrate(s) of such phosphatase activity? An obvious candidate is the TON1

phosphoprotein³⁵, which directly interacts and co-localizes with FASS on microtubules, and whose depletion induces the exact same phenotype as the FASS mutation. The TTP complex could also participate in a pathway involved in the regulation of components of the microtubule nucleation machinery, as recent results established a role for FASS in controlling the ratio of in-bundle microtubule nucleation to branched nucleation points¹². Another potential target is the cortical division zone marker AtTAN, whose persistence at the cortex during preprophase depends on PP2A activity³⁶. Several kinases may act to counteract the TTP phosphatase activity since reversible protein phosphorylation/dephosphorylation has an important role in cortical microtubular organization^{37,38}. One of these could be the cyclin-dependent kinase CDKA1, previously shown to decorate the late PPB and to induce its disassembly^{39–42}, and which co-purified with TON1 in TAP experiments⁴³. Aurora kinases, which in animals are functional antagonists of PP2A at the centrosome and microtubule plus ends^{44–46}, are also putative candidates as suggested by recent results pointing to a function for Aurora kinases in premitotic division plane orientation and cortical cytoskeleton organization^{47,48}.

Besides a role in PPB formation, the TTP complex likely has a role in interphase cortical array organization as well. First, the TTP complex is robustly isolated from non-synchronised suspension cells where mitotic cells are a minority. Second, transcription of *TTP* genes are generally not cell-cycle regulated, with the exception of few *TRMs*¹⁵. Furthermore, introducing *pp2aa* or *pp2ac* mutations in weak *fass* alleles that fail to produce PPBs but are still capable of residual interphasic cell elongation, results in a strong *fass* phenotype and suppresses this residual interphasic growth. Finally, some TRM proteins have been shown to specifically function in cell elongation^{15,49}.

Strikingly, all components of the plant TTP complex identified here show various levels of sequence similarity with animal counterparts localizing transiently or permanently to centrosomes and centrioles/basal bodies. This ranges from whole protein similarity between PP2A subunits (including FASS), to domains conserved between TON1 and FOP, and to conserved motifs between TRM proteins and CAP350 (refs 11,16,17). All available studies point to centrosomal functions for these proteins, either mitotic (RSA1 (ref. 17)), centrosomal and/or ciliary (FOP/FOR20 (ref. 50), CAP350 (ref. 16)) or linked with cell-cycle progression (FOP⁵¹). However, and although animal counterparts of the three core TTP components were recently co-purified in human cells⁵², the function of the TTP network as a whole has not been formally studied in any system apart from plants. The diversity of TRMs in plants, as opposed to the single CAP350 in animals, may reflect the divergence of the two kingdoms with respect to spatial distribution of the nucleation process and organization of the cytoskeleton.

Methods

Plant material and growth conditions. *Arabidopsis* seedlings were grown on half-strength Murashige and Skoog (MS) medium pH 5.7 containing 1% sucrose (w/v), 0.05% MES (w/v), 0.01% (w/v) myo-inositol and 0.8% agar at 21 °C in a 16-h light/8-h dark regime. The PSB-D *Arabidopsis thaliana* cell suspension cultures expressing the GS-tag fusion proteins were maintained in 50 ml of MSMO medium (4.43 g l⁻¹ MSMO (Sigma-Aldrich), 30 g l⁻¹ sucrose, 0.5 mg l⁻¹ α -naphthaleneacetic acid, 0.05 mg l⁻¹ kinetin, pH 5.7, adjusted with 1 M KOH) at 25 °C in the dark by gentle agitation (130 r.p.m.). Every 7 days, the cells were subcultured in fresh medium at a 1:6 dilution⁴³. *Nicotiana benthamiana* plants were grown in the growth chamber under 16 h of light, a diurnal temperature of 25 °C and a nocturnal temperature of 20 °C.

T-DNA insertion mutants for *PP2AC3* (SALK_069250) and *PP2AC4* (SALK_035009) were obtained from the Nottingham Arabidopsis Stock Centre⁵³. *pp2aa* mutants were the kind gift of A. Delong²⁵. The *ton1* mutant carries a deletion affecting both *TON1a* and *TON1b* genes¹¹. *fass* mutants in this study have

been renamed for clarity: *fass-5*, *fass-13*, and *fass-14* alleles correspond to *ton2-5*, *ton2-13* and *ton2-14* in Camilleri *et al.*¹⁰, and *fass-15* corresponds to *ton2-15* in Kirik *et al.*¹²

Construct design. FASS, TON1, PP2AA1, TRM8, TRM1 as well as truncated versions of TON1, TRM1 and FASS open-reading frames were amplified from *Arabidopsis* complementary DNA clones (Columbia ecotype) using specific primers flanked by AttB1 and AttB2 sites (Supplementary Table S4), cloned into the gateway vector pDONR207 using BP recombination (Invitrogen), and sequenced. Expression vectors were obtained after LR recombination (Invitrogen) between these entry vectors and the following destination vectors: pGWB5 and pGWB6 (ref. 54) for expression of C- and N-terminal GFP fusions, respectively, pH7RWG2 and pH7WGR2 (ref. 55) for expression of C- and N-terminal RFP fusions, respectively, pBiFP vectors for split YFP experiments¹¹, and pLex10- and pGADT7-derived plasmids for yeast two-hybrid interaction assays¹⁵.

Genotyping, RT and qPCR analysis. Primers used to genotype *pp2aa* and *pp2ac* mutants are indicated in Supplementary Table S4. Genotyping PCRs were performed on genomic DNA isolated from a rosette leaf⁵⁶.

Quantitative PCR (qPCR) and RT-PCR were done on RNA isolated from 6-day-old *pp2ac3-c4* double mutant seedlings or RNA from PSB-D cell cultures. RNA was extracted using Trizol (TRI Reagent-Molecular Research Center, Inc.) followed by purification using the RNeasy plant mini kit (Qiagen). cDNA was synthesized from 1 μ g of total RNA using the iScript cDNA synthesis kit (Bio-Rad). Five microlitres of ten times diluted cDNA was used for qPCR, performed with the Power SYBR Green mix (Applied Bioscience) on a Roche thermocycler (Roche). Each reaction was done in triplicate and both *ACTIN* and *EIF4A* were used as normalization genes. The RT-PCR to analyse the presence of remaining full-length transcript in the *pp2ac3-c4* double mutant consisted of 26 repeating cycles with an annealing temperature of 55 °C. The RT-PCR analysis to test for the expression of *PP2AA* and *PP2AC* subunits in the PSB-D cell cultures consisted of a touchdown protocol with a gradual decrease of the annealing temperature from 65 to 55 °C during the first 10 cycles followed by 30 cycles with a constant annealing temperature at 55 °C (see Supplementary Table S4 for the primers used).

Two-hybrid assays. For protein interaction assays, the L40 yeast strain was used (*MATa trp1 leu2 his3 ade2 LYS2::lexA-HIS3 URA3::lexA-lacZ*). Yeast samples transformed with each bait construct (along with empty prey vector) were plated on minimal medium lacking tryptophan, leucine and histidine with increasing concentration of 3-amino-1,2,4-triazole (0–200 mM) to determine the levels of background self-activation of the *HIS3* gene. The lowest concentration of 3-amino-1,2,4-triazole that inhibited growth was used to study pairwise interactions in yeast containing both bait and prey vectors. A LexA-FASS fusion protein was used as a bait to screen a custom cDNA library (Invitrogen) prepared from equal amount of Poly-A⁺ RNA of various Col0 *Arabidopsis* samples (young *in vitro*-grown seedlings, roots, siliques and seeds, cell-culture suspension, flower buds). Experimental procedures for screening, quantitative β -Gal assays and plasmid isolation were performed according to the manufacturer's user guide.

Plant transformation. Overnight *Agrobacterium* cultures were pelleted, washed and resuspended in infiltration buffer (13 g l⁻¹ S-medium (Duchefa) and 40 g l⁻¹ sucrose, pH 5.7) to an OD₆₀₀ of 0.5. For co-infiltration experiments, equal volumes of the two (or three or four) cultures of OD₆₀₀ of 0.5 were mixed before agro-infiltration. The inoculum was then delivered to the lamina tissue of *N. benthamiana* leaves by gentle pressure infiltration through the lower epidermis. Negative controls correspond to co-expression of YFP^C-TON1 with the unrelated YFP^N-GLOBOSA protein, or to co-expression of YFP^N-FASS with the unrelated YFP^C-DEFICIENS¹¹ and background levels were checked by expression of YFP^C-TON1 or YFP^N-FASS alone.

Microscopy and immunolocalization. For GFP/RFP imaging, fluorescence was recorded using an inverted Zeiss Observer Z1 spectral confocal laser microscope LSM 710 (ref. 15). Embryo imaging was performed using propidium iodide staining⁵⁷. *Arabidopsis* roots of 3- to 4-day-old seedlings were processed for whole-mount immunolocalization⁵⁸ using the B-5-1-2 monoclonal anti- α -tubulin (Sigma-Aldrich) and the anti-KNOLLE antibody (kind gift of G. Jürgens). Scanning electron micrographs were obtained using the Hirox SH-1500 microscope with fresh *Arabidopsis* seedling samples.

Tandem affinity purification. Transgenes encoding tag fusions were cloned by gateway recombination under control of the constitutive cauliflower tobacco mosaic virus 35S promoter⁴³. Transformation of *Arabidopsis* cell suspension cultures was performed by *Agrobacterium* co-cultivation and transgenic culture regeneration⁴³. TAP of protein complexes was done using the GS tag^{19,20} with minor buffer modifications for purification in the presence of digitonin: crude protein extracts were obtained with the Ultra-Turrax T25 mixer (IKA Works, Wilmington, NC) using 80% HB without 0.1% (v/v) Nonidet P-40 (NP-40).

Afterwards the remaining 20% HB containing 5% digitonin (w/v) was added and the sample incubated for 1 h at 4 °C on a rotating wheel. The cell extract was cleared by two subsequent centrifugation steps at 36,900 × g for 20 min. IgG beads were equilibrated with digitonin-free HB. In both IgG and TEV, wash buffer 0.1% (v/v) NP-40 was replaced by 0.2% (w/v) digitonin.

Following TAP, the protein complexes were precipitated and prepared for mass spectrometry analysis. For analysis through MALDI TOF/TOF (TON1a, TON1b, FASS, TRM19, see Supplementary Table S1 for details), proteins were separated by NuPAGE (Invitrogen)²⁰. Gel lanes were cut in slices, proteins digested by trypsin and peptides isolated⁵⁹. Mass spectra were obtained by a 4800 MALDI TOF/TOF Proteomics Analyzer (AB SCIEX), and MS-based protein homology identification on the basis of the TAIR genomic database⁵⁹. Experimental background proteins were subtracted on the basis of ~40 TAP experiments on wild-type cultures and cultures expressing TAP-tagged GUS, RFP and GFP mock proteins⁵⁹. For analysis through a LTQ Orbitrap Velos mass spectrometer (Thermo, Bremen, Germany) (TON1a, TON1b and FASS, see Supplementary Methods for details), TCA-precipitated proteins were separated in a short run of 7 min on a 4–12% gradient NuPAGE gel (Invitrogen) and visualized with colloidal Coomassie Brilliant Blue staining. The protein gel was washed for 1 h in H₂O, polypeptide disulphide bridges were reduced for 40 min in 25 ml of 6.66 mM DTT in 50 mM NH₄HCO₃ and sequentially the thiol groups were alkylated for 30 min in 25 ml of 55 mM IAM in 50 mM NH₄HCO₃. After washing with H₂O, a broad zone containing the proteins was cut from the protein gel, sliced into 24 gel plugs, and collected together in a single Eppendorf. Gel plugs were washed twice with H₂O, dehydrated with 95% CH₃CN (v/v), rehydrated with H₂O and dehydrated again with 95% CH₃CN (v/v). Dehydrated gel particles were rehydrated in 60 µl digest buffer containing 750 ng trypsin (MS Gold; Promega, Madison, WI), 50 mM NH₄HCO₃ and 10% CH₃CN (v/v) for 30 min at 4 °C. Proteins were digested at 37 °C for 3.5 h. LC-MS/MS runs on LTQ Orbitrap Velos, and peak list generation and submission for protein identification to the TAIR database are described in Supplementary Table S2 (TON1b) and Supplementary Table S3 (TON1a and FASS). Proteins identified with at least two high-confidence peptides were retained. A list of non-specific background proteins was assembled by combining our previous background list⁵⁹ with background proteins from control GS purifications on mock, GFP-GS and GUS-GS cell culture extracts and GFP-GS 6-day-old plant extracts identified with Orbitrap VELOS. To obtain the final list of interactors, these background proteins were subtracted from the list of identified proteins.

Protein–protein interaction networks were built with the Cytoscape software⁶⁰.

Blue Native-SDS-PAGE. Total protein extracts from 0.7 g harvested *Arabidopsis* PSB-D suspension cells expressing 35S::TON1b-GS or 35S::GS-TON1a were prepared according to the NativePAGE Novex Bis-Tris gels electrophoresis protocol by Invitrogen with 0.1% benzamide and 2% digitonin added to the BN lysis buffer. Extracted protein complexes were separated under native conditions over a 4–16% NativePAGE Novex Bis-Tris gel (Invitrogen). Lanes were subsequently cut out and incubated with denaturing buffer (500 mM Tris pH 6.8, 66 mM Na₂CO₃, 10% (w/v) glycerol, 2% (w/v) SDS, 2% β-mercaptoethanol) for 1 h under gentle agitation. Denatured first-dimension lanes were placed onto the stacking of a 12% SDS-PAGE gel and the proteins were separated in the second dimension using the Mini-PROTEAN II system (Bio-Rad) for 70 min at 180 V. Proteins in the second-dimension gels were blotted on Immobilon-P membranes (Millipore). Membranes were blocked overnight at 4 °C in 3% (v/v) milk powder dissolved in 25 mM Tris-HCl, pH 8, 150 mM NaCl and 0.1% Triton X-100. Western blots were developed using the Peroxidase Anti-Peroxidase (PAP) antibody (1:2,500) (P1291, Sigma-Aldrich).

References

- De Smet, I. & Beeckman, T. Asymmetric cell division in land plants and algae: the driving force for differentiation. *Nat. Rev. Mol. Cell Biol.* **12**, 177–188 (2011).
- Rasmussen, C. G., Humphries, J. A. & Smith, L. G. Determination of symmetric and asymmetric division planes in plant cells. *Annu. Rev. Plant Biol.* **62**, 387–409 (2011).
- Mineyuki, Y. The preprophase band of microtubules: its function as a cytokinetic apparatus in higher plants. *Int. Rev. Cytol.* **187**, 1–49 (1999).
- Van Damme, D. Division plane determination during plant somatic cytokinesis. *Curr. Opin. Plant Biol.* **12**, 745–751 (2009).
- Lloyd, C. Dynamic microtubules and the texture of plant cell walls. *Int. Rev. Cell Mol. Biol.* **287**, 287–329 (2011).
- Paradez, A., Wright, A. & Ehrhardt, D. W. Microtubule cortical array organization and plant cell morphogenesis. *Curr. Opin. Plant Biol.* **9**, 571–578 (2006).
- Duroc, Y., Bouchez, D. & Pastuglia, M. in *Advances in Plant Biology 2, The Plant Cytoskeleton* (ed. Liu, B.) 145–185 (Springer, USA, 2010).
- Torres-Ruiz, R. A. & Jürgens, G. Mutations in the FASS gene uncouple pattern formation and morphogenesis in *Arabidopsis* development. *Development* **120**, 2967–2978 (1994).
- Traas, J. *et al.* Normal differentiation patterns in plants lacking microtubular preprophase bands. *Nature* **375**, 676–677 (1995).
- Camilleri, C. *et al.* The *Arabidopsis* TONNEAU2 gene encodes a putative novel PP2A regulatory subunit essential for the control of cortical cytoskeleton. *Plant Cell* **14**, 833–845 (2002).
- Azimzadeh, J. *et al.* *Arabidopsis* TONNEAU1 proteins are essential for preprophase band formation and interact with centrins. *Plant Cell* **20**, 2146–2159 (2008).
- Kirik, A., Ehrhardt, D. W. & Kirik, V. TONNEAU2/FASS regulates the geometry of microtubule nucleation and cortical array organization in interphase *Arabidopsis* cells. *Plant Cell* **24**, 1158–1170 (2012).
- Wright, A. J., Gallagher, K. & Smith, L. G. *discordia1* and *alternative discordia1* function redundantly at the cortical division site to promote preprophase band formation and orient division planes in maize. *Plant Cell* **21**, 234–247 (2009).
- Spinner, L. *et al.* The function of TONNEAU1 in moss reveals ancient mechanisms of division plane specification and cell elongation in land plants. *Development* **137**, 2733–2742 (2010).
- Drevensek, S. *et al.* The *Arabidopsis* TRM1-TON1 interaction reveals a recruitment network common to plant cortical microtubule arrays and eukaryotic centrosomes. *Plant Cell* **24**, 178–191 (2012).
- Yan, X., Habedanck, R. & Nigg, E. A. A complex of two centrosomal proteins, CAP350 and FOP, cooperates with EB1 in microtubule anchoring. *Mol. Biol. Cell* **17**, 634–644 (2006).
- Schlaitz, A. L. *et al.* The *C. elegans* RSA complex localizes protein phosphatase 2A to centrosomes and regulates mitotic spindle assembly. *Cell* **128**, 115–127 (2007).
- Janssens, V., Longin, S. & Goris, J. PP2A holoenzyme assembly: in cauda venenum (the sting is in the tail). *Trends Biochem. Sci.* **33**, 113–121 (2008).
- Burckstummer, T. *et al.* An efficient tandem affinity purification procedure for interaction proteomics in mammalian cells. *Nat. Methods* **3**, 1013–1019 (2006).
- Van Leene, J., Witters, E., Inze, D. & De Jaeger, G. Boosting tandem affinity purification of plant protein complexes. *Trends Plant Sci.* **13**, 517–520 (2008).
- Mikolajka, A. *et al.* Structure of the N-terminal domain of the FOP (FGFR1OP) protein and implications for its dimerization and centrosomal localization. *J. Mol. Biol.* **359**, 863–875 (2006).
- Larsen, P. B. & Cancel, J. D. Enhanced ethylene responsiveness in the *Arabidopsis* eer1 mutant results from a loss-of-function mutation in the protein phosphatase 2A regulatory subunit, RCN1. *Plant J.* **34**, 709–718 (2003).
- Kwak, J. M. *et al.* Disruption of a guard cell-expressed protein phosphatase 2A regulatory subunit, RCN1, confers abscisic acid insensitivity in *Arabidopsis*. *Plant Cell* **14**, 2849–2861 (2002).
- Garbers, C., DeLong, A., Deruere, J., Bernasconi, P. & Soll, D. A mutation in protein phosphatase 2A regulatory subunit A affects auxin transport in *Arabidopsis*. *EMBO J.* **15**, 2115–2124 (1996).
- Zhou, H. W., Nussbaumer, C., Chao, Y. & DeLong, A. Disparate roles for the regulatory A subunit isoforms in *Arabidopsis* protein phosphatase 2A. *Plant Cell* **16**, 709–722 (2004).
- Janski, N. *et al.* The GCP3-interacting proteins GIP1 and GIP2 are required for gamma-tubulin complex protein localization, spindle integrity, and chromosomal stability. *Plant Cell* **24**, 1171–1187 (2012).
- Blakeslee, J. J. *et al.* Specificity of RCN1-mediated protein phosphatase 2A regulation in meristem organization and stress response in roots. *Plant Physiol.* **146**, 539–553 (2008).
- Nakamura, M., Ehrhardt, D. W. & Hashimoto, T. Microtubule and katanin-dependent dynamics of microtubule nucleation complexes in the acentrosomal *Arabidopsis* cortical array. *Nat. Cell Biol.* **12**, 1064–1070 (2010).
- Laubert, M. H. *et al.* The *Arabidopsis* KNOLLE protein is a cytokinesis-specific syntaxin. *J. Cell Biol.* **139**, 1485–1493 (1997).
- Reichardt, I. *et al.* Plant cytokinesis requires de novo secretory trafficking but not endocytosis. *Curr. Biol.* **17**, 2047–2053 (2007).
- Pickett-Heaps, J. D. & Northcote, D. H. Organization of microtubules and endoplasmic reticulum during mitosis and cytokinesis in wheat meristems. *J. Cell Sci.* **1**, 109–120 (1966).
- Xu, Z. *et al.* Structure and function of the PP2A-shugoshin interaction. *Mol. Cell* **35**, 426–441 (2009).
- Perez-Callejon, E. *et al.* Molecular cloning and characterization of two phosphatase 2A catalytic subunit genes from *Arabidopsis thaliana*. *Gene* **209**, 105–112 (1998).
- Ballesteros, I. *et al.* Specialised functions of the PP2A subfamily II catalytic subunits PP2A-C3 and PP2A-C4 in the distribution of auxin fluxes and development in *Arabidopsis*. *Plant J.* **73**, 862–872 (2012).
- Benschop, J. J. *et al.* Quantitative phosphoproteomics of early elicitor signaling in *Arabidopsis*. *Mol. Cell Proteomics* **6**, 1198–1214 (2007).
- Rasmussen, C. G., Sun, B. & Smith, L. G. Tangled localization at the cortical division site of plant cells occurs by several mechanisms. *J. Cell Sci.* **124**, 270–279 (2011).

37. Baskin, T. & Wilson, J. Inhibitors of protein kinases and phosphatases alter root morphology and disorganize cortical microtubules. *Plant Physiol.* **113**, 493–502 (1997).
38. Ayaydin, F. *et al.* Inhibition of serine/threonine-specific protein phosphatases causes premature activation of cdc2Msf kinase at G2/M transition and early mitotic microtubule organisation in alfalfa. *Plant J.* **23**, 85–96 (2000).
39. Colasanti, J., Cho, S. O., Wick, S. & Sundaresan, V. Localization of the functional p34cdc2 homolog of maize in root tip and stomatal complex cells: Association with predicted division sites. *Plant Cell* **5**, 1101–1111 (1993).
40. Hush, J., Wu, L., John, P. C., Hepler, L. H. & Hepler, P. K. Plant mitosis promoting factor disassembles the microtubule preprophase band and accelerates prophase progression in *Tradescantia*. *Cell Biol. Int.* **20**, 275–287 (1996).
41. Weingartner, M. *et al.* Dynamic recruitment of Cdc2 to specific microtubule structures during mitosis. *Plant Cell* **13**, 1929–1943 (2001).
42. Boruc, J. *et al.* Systematic localization of the *Arabidopsis* core cell cycle proteins reveals novel cell division complexes. *Plant Physiol.* **152**, 553–565 (2010).
43. Van Leene, J. *et al.* A tandem affinity purification-based technology platform to study the cell cycle interactome in *Arabidopsis thaliana*. *Mol. Cell Proteomics* **6**, 1226–1238 (2007).
44. Horn, V. *et al.* Functional interaction of Aurora-A and PP2A during mitosis. *Mol. Biol. Cell* **18**, 1233–1241 (2007).
45. Zimniak, T., Stengl, K., Mechtler, K. & Westermann, S. Phosphoregulation of the budding yeast EB1 homologue Bim1p by Aurora/Ipl1p. *J. Cell Biol.* **186**, 379–391 (2009).
46. Sun, L. *et al.* EB1 promotes Aurora-B kinase activity through blocking its inactivation by protein phosphatase 2A. *Proc. Natl Acad. Sci. USA* **105**, 7153–7158 (2008).
47. Van Damme, D. *et al.* *Arabidopsis* alpha Aurora kinases function in formative cell division plane orientation. *Plant Cell* **23**, 4013–4024 (2011).
48. Petrovskaya, B. *et al.* Plant Aurora kinases play a role in maintenance of primary meristems and control of endoreduplication. *New Phytol.* **193**, 590–604 (2012).
49. Lee, Y. K. *et al.* *LONGIFOLIA1* and *LONGIFOLIA2*, two homologous genes, regulate longitudinal cell elongation in *Arabidopsis*. *Development* **133**, 4305–4314 (2006).
50. Sedjai, F. *et al.* Control of ciliogenesis by FOR20, a novel centrosome and pericentriolar satellite protein. *J. Cell Sci.* **123**, 2391–2401 (2010).
51. Acquaviva, C. *et al.* The centrosomal FOP protein is required for cell cycle progression and survival. *Cell Cycle* **8**, 1217–1227 (2009).
52. Hutchins, J. R. *et al.* Systematic analysis of human protein complexes identifies chromosome segregation proteins. *Science* **328**, 593–599 (2010).
53. Alonso, J. M. *et al.* Genome-wide insertional mutagenesis of *Arabidopsis thaliana*. *Science* **301**, 653–657 (2003).
54. Nakagawa, T. *et al.* Development of series of gateway binary vectors, pGWBs, for realizing efficient construction of fusion genes for plant transformation. *J. Biosci. Bioeng.* **104**, 34–41 (2007).
55. Van Damme, D. *et al.* In vivo dynamics and differential microtubule-binding activities of MAP65 proteins. *Plant Physiol.* **136**, 3956–3967 (2004).
56. Pastuglia, M. *et al.* Gamma-tubulin is essential for microtubule organization and development in *Arabidopsis*. *Plant Cell* **18**, 1412–1425 (2006).
57. Truernit, E. *et al.* High-resolution whole-mount imaging of three-dimensional tissue organization and gene expression enables the study of Phloem development and structure in *Arabidopsis*. *Plant Cell* **20**, 1494–1503 (2008).
58. Truernit, E., Bauby, H., Belcram, K., Barthelemy, J. & Palauqui, J. C. OCTOPUS, a polarly localised membrane-associated protein, regulates phloem differentiation entry in *Arabidopsis thaliana*. *Development* **139**, 1306–1315 (2012).
59. Van Leene, J. *et al.* Targeted interactomics reveals a complex core cell cycle machinery in *Arabidopsis thaliana*. *Mol. Syst. Biol.* **6**, 397 (2010).
60. Cline, M. S. *et al.* Integration of biological networks and gene expression data using Cytoscape. *Nat. Protoc.* **2**, 2366–2382 (2007).

Acknowledgements

We are grateful to A. Delong, G. Jürgens, T. Hashimoto and M-E. Chabouté for sharing published materials, to Magalie Uyttewaal for helpful discussions, to M. Anjuere and P. Grillot for taking care of the plants and to J. Van Leene and L. Vercruysse for help with experiments. L.S. and E.S. were recipient of a PhD fellowship from the French Ministry for Research, A.G. is indebted to the Agency for Innovation by Science and Technology for a predoctoral fellowship, Y.D. was funded by the Agence Nationale de la Recherche (ANR-08-BLAN-0056) and D.V.D. was a postdoctoral fellow of the Research Foundation-Flanders. We thank the 'Région Île-de-France' and 'Conseil Général des Yvelines' for supporting our microscopy platform. This work was supported by a grant from the Agence Nationale de la Recherche (ANR-08-BLAN-0056).

Author contributions

L.S., A.G., G.D.J., D.B., D.V.D. and M.P. designed the research; L.S., A.G., K.B., M.G., M.M., Y.D., D.E., E.S., E.V.D.S., G.P., D.V.D. and M.P. performed research; N.D.W., E.W. and K.G. contributed new analytic tools; L.S., A.G., K.B., M.G., Y.D., D.E., E.S., G.D.J., D.B., D.V.D. and M.P. analysed data; A.G., G.D.J., D.B., D.V.D. and M.P. wrote the paper; D.V.D. and M.P. share senior authorship of this paper.

Additional information

Supplementary Information accompanies this paper at <http://www.nature.com/naturecommunications>

Competing financial interests: The authors declare no competing financial interests.

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How to cite this article: Spinner, L. and Gadeyne, A. *et al.* A protein phosphatase 2A complex spatially controls plant cell division. *Nat. Commun.* 4:1863 doi: 10.1038/ncomms2831 (2013).