The localization of PHRAGMOPLAST ORIENTING KINESIN1 at the division site depends on the microtubule-binding proteins TANGLED1 and AUXIN-INDUCED IN ROOT CULTURES9 in Arabidopsis

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Abstract
Proper plant growth and development require spatial coordination of cell divisions. Two unrelated microtubule-binding proteins, TANGLED1 (TAN1) and AUXIN-INDUCED IN ROOT CULTURES9 (AIR9), are together required for normal growth and division plane orientation in Arabidopsis (Arabidopsis thaliana). The tan1 air9 double mutant has synthetic growth and division plane orientation defects, while single mutants lack obvious defects. Here we show that the division site-localized protein, PHRAGMOPLAST ORIENTING KINESIN1 (POK1), was aberrantly lost from the division site during metaphase and telophase in the tan1 air9 mutant. Since TAN1 and POK1 interact via the first 132 amino acids of TAN1 (TAN11–132), we assessed the localization and function of TAN11–132 in the tan1 air9 mutant. TAN11–132 rescued tan1 air9 mutant phenotypes and localized to the division site during telophase. However, replacing six amino-acid residues within TAN11–132, which disrupted the POK1–TAN1 interaction in the yeast-two-hybrid system, caused loss of both rescue and division site localization of TAN11–132 in the tan1 air9 mutant. Full-length TAN1 with the same alanine substitutions had defects in phragmoplast guidance and reduced TAN1 and POK1 localization at the division site but rescued most tan1 air9 mutant phenotypes. Together, these data suggest that TAN1 and AIR9 are required for POK1 localization, and yet unknown proteins may stabilize TAN1–POK1 interactions.

Introduction
Division plane orientation is important for many aspects of plant, microbial, and animal development, particularly growth and patterning. Division plane orientation is especially relevant for plant cells, which are encased in cell walls and unable to migrate (Facette et al., 2018; Rasmussen and...
The phragmoplast expands by the forming cell plate (McMichael and Bednarek, 2013; Smertenko et al., 2018). The phragmoplast microtubules are tracks for the movement of vesicles containing cell wall materials toward the cell cortex, and the cell plate and associated membranes fuse with the mother cell membranes at the cell plate fusion site previously specified by the PPB (van Oostende-Triplet et al., 2017).

**Background:** Unlike animal cells, plant cells cannot move due to their semi-rigid cell walls. The correct positioning of the new cell wall is important for overall plant growth and development. Three microtubule-binding proteins are involved in division plane orientation: TANGLED1 (TAN1), AUXIN-INDUCED IN ROOT CULTURES9 (AIR9), and PHRAGMOPHAST ORIENTING KINESIN1 (POK1). These proteins localize as a ring at the edge of the cell where the new cell wall will insert during cell division, at a position called the division site. Here, we focused on how TAN1 and POK1 interactions promote their localization to the division site, and their function in plant growth and division plane positioning.

**Question:** How do TAN1 and AIR9 contribute to POK1 localization, and how does POK1 localization affect new cell wall placement?

**Findings:** TAN1 and AIR9 together maintain POK1 at the division site in Arabidopsis thaliana. When mutant versions of TAN1 that no longer interact with POK1 were transformed into the tan1 air9 double mutant, POK1 and TAN1 localization was partially disrupted and cell wall placement defects occurred. This suggests that POK1 interaction with TAN1 is important for their correct division site localization and new cell wall placement.

**Next steps:** This work strongly suggests that yet unknown proteins mediate TAN1 and POK1 interaction. Discovering what those proteins are, and how AIR9 contributes to division plane positioning are next. Understanding how plants position their division plane will contribute to understanding plant growth and has the long-term potential to contribute to next-generation crop development.
mutant also fails to maintain TAN1 at the division site after entry into metaphase (Lipka et al., 2014). This suggests that TAN1 maintenance at the division site after metaphase is dependent on POK1 and POK2.

In Arabidopsis, the tan1 mutant has minor phenotypic differences compared with wild-type (WT) plants (Walker et al., 2007). However, the tan1 auxin-induced-in-root-cultures9 (air9) double mutant, which has no obvious defects (Buschmann et al., 2015), resulted in a synthetic phenotype consisting of reduced root growth, increased root cell file rotation, and phragmoplast guidance defects (Mir et al., 2018). TAN1 and AIR9 are unrelated microtubule-binding proteins that both localize to the division site (Buschmann et al., 2006; Walker et al., 2007). Both TAN1 and AIR9 colocalize with the PPB. TAN1 remains at the division site throughout cell division, while AIR9 is lost from the division site upon PPB disassembly and then reappears at the division site during cytokinesis when the phragmoplast contacts the cortex. When full-length TAN1 fused to YELLOW FLUORESCENT PROTEIN (TAN1-YFP) and driven either by the constitutive viral cauliflower mosaic CaMV35S promoter (p35S: TAN1-YFP) or the native promoter with the fluorescent protein as either an N- or C-terminal fusion (pTAN1: TAN1–YFP or pTAN1: CFP–TAN1) was transformed into the tan1 air9 double mutant, the phenotype was rescued such that plants looked similar to and grew as well as WT plants (Mir et al., 2018; Mills and Rasmussen, 2022).

TAN1 is an intrinsically disordered protein with no well-defined domains. It was divided into five conserved regions based on alignments of amino acid similarity across plant species. Region I, which covers the first approximately 130 amino acids of TAN1, is the most highly conserved and mediates TAN1 localization to the division site during telophase. This approximately 130 amino acid region also mediates interactions between TAN1 and POK1 in the yeast two-hybrid system (Rasmussen et al., 2011). When TAN1 missing the sequence for the first approximately 130 amino acids was transformed into the tan1 air9 double mutant, no rescue was observed (Mir et al., 2018). This suggests that the first approximately 130 amino acids of the TAN1 protein are critical for function in root growth and division plane positioning.

Here, we show that both AIR9 and TAN1 are required for POK1 to remain at the division site after PPB disassembly. We identified TAN1–POK1 interaction motifs within the first 132 amino acids using the yeast two-hybrid system. Interestingly, the first 132 amino acids of TAN1 (TAN11–132) are sufficient to rescue the tan1 air9 double mutant, but not when a TAN1–POK1 interaction motif was disrupted. We found that when full-length TAN1 with the same mutated motif was used, substantial rescue was observed, except for defects in phragmoplast guidance and loss of POK1 and TAN1 at the division site during metaphase and telophase. Together, this suggests that interactions between POK1 and AIR9, and TAN1 and POK1, as well as other yet unknown proteins, are important for division plane orientation and plant growth.

**Results**

Either TAN1 or AIR9 is sufficient to recruit and maintain POK1 at the division site

To understand how known division site-localized proteins interact at the division site, we examined POK1 fused to YFP–POK1 (Lipka et al., 2014) localization in WT, tan1 air9 double mutant and single mutant Arabidopsis plants expressing the microtubule marker UBX10:mScarlet-MAP4 (Pan et al., 2020). Our hypothesis was that POK1 localization would not be contingent on TAN1 or AIR9 and would therefore be unaltered in the tan1 air9 double mutant. In contrast to our hypothesis, YFP–POK1 was lost from the division site during metaphase and telophase and accumulated less frequently during prophase and prophase in WT cells. YFP–POK1 colocalized with PPBs in 71% of prophase/prophase cells (n = 50/70 cells, 15 plants, Figure 1A) consistent with previous observations (Lipka et al., 2014; Schaefer et al., 2017). In the tan1 air9 double mutant cells, YFP–POK1 colocalized with 50% of PPBs during prophase/prophase, which was not significantly different from the colocalization rate in WT cells (n = 27/54 cells, 15 plants, Figure 1, D and M; Fisher's exact test, P = 0.0165, not significant with Bonferroni correction). In WT cells, YFP–POK1 remained at the division site in all observed metaphase (n = 13/13 cells, Figure 1B) and telophase cells (n = 31/31 cells, Figure 1C), similar to previous studies (Lipka et al., 2014). In rare instances, YFP–POK1 also accumulated in the phragmoplast midline in WT cells (13%, n = 4/31, 11 plants, Figure 1M). In contrast, in tan1 air9 mutants, YFP–POK1 was lost from the division site in metaphase (n = 0/21 cells, Figure 1, E and M) and telophase (n = 0/44, Figure 1F). Interestingly, in tan1 air9 double mutant cells, although YFP–POK1 did not accumulate at the division site, it accumulated at the phragmoplast midline in 77% of cells (n = 34/44), which was a significantly greater midline accumulation frequency than the 13% observed in WT plants (n = 4/31 cells, Fisher's exact test, P < 0.00001). Together, this shows that POK1 is not maintained at the division site after PPB disassembly and that instead it accumulates in the phragmoplast midline. We hypothesize that mislocalized phragmoplast midline accumulation of YFP–POK1 in the tan1 air9 mutant occurs because YFP–POK1 is not maintained at the division site.

Next, we examined YFP–POK1 localization in tan1 and air9 single mutants. YFP–POK1 localized to the division site during all mitotic stages, but aberrantly accumulated in the phragmoplast midline in the tan1 single mutant. Similar to in WT plants, YFP–POK1 colocalized with PPBs during prophase or prophase in the tan1 mutant (Figure 1G) and air9 mutant (Figure 1H), and remained at the division site during metaphase and telophase (Figure 1, G–L and M). In the tan1 single mutant, YFP–POK1 localized both to the division site and the phragmoplast midline in 44% of
TAN1 and AIR9 together promote POK1 maintenance at the division site. YFP–POK1 localization in Col-0 WT, tan1 single mutant, air9 single mutant, and tan1 air9 double mutant plants expressing UBQ10:mScarlet-MAP4 to mark microtubules and pPOK1:YFP–POK1. Scale bars = 10 μm. A–C, YFP–POK1 localization in Col-0 WT plants. A, YFP–POK1 localization during preprophase/prophase. B, YFP–POK1 was maintained at the division site in metaphase and anaphase cells in Col-0 plants. C, YFP–POK1 remains clearly visible at the division site in Col-0 telophase cells. D–F, YFP–POK1 localization in tan1 air9 double mutant plants. D, YFP–POK1 localization during preprophase/prophase. E, YFP–POK1 was lost from the division site upon entry into metaphase. F, In tan1 air9 telophase cells YFP–POK1 was absent from the division site and accumulated in the phragmoplast midline. G–I, YFP–POK1 localization in tan1 single mutant plants. G, YFP–POK1 localization in the tan1 single mutant during preprophase/prophase. H, YFP–POK1 was maintained at the division site in metaphase and anaphase cells in tan1 plants. I, YFP–POK1 remains clearly visible at the division site in tan1 telophase cells. J–L, YFP–POK1 localization in air9 single mutant plants. J, YFP–POK1 localization during preprophase/prophase. K, YFP–POK1 was maintained at the division site in metaphase and anaphase cells in air9 plants. L, YFP–POK1 remains clearly visible at the division site in air9 telophase. M, YFP–POK1 subcellular localization in WT, tan1 air9, tan1, and air9 plants in preprophase/prophase, metaphase, and telophase cells. N > 14 plants for each genotype. Statistically significant differences in localization compared to WT plants were determined using Fisher’s exact test with Bonferroni correction for four sample types. Asterisks indicate significant differences. YFP–POK1 colocalized with the PPB in 71% of WT cells (50/70), 50% of tan1 air9 cells (27/54), 64% of tan1 cells (54/85), and 64% of air9 cells (46/72). YFP–POK1 colocalization with the PPB was not significantly different in tan1 air9, tan1, and air9 plants compared to in WT plants. YFP–POK1 was maintained at the division site in all metaphase cells of WT (13/13 cells), tan1 (17/17 cells), and air9 (24/24 cells) plants. In tan1 air9 double mutant plants, YFP–POK1 was not maintained at the division site during metaphase (0/21 cells, P < 0.00001). During telophase, YFP–POK1 accumulated at the division site only in 87% of WT cells (27/31). In 13% of WT telophase cells, YFP–POK1 accumulated at the division site and in the phragmoplast midline (4/31 cells). YFP–POK1 localization in telophase cells of air9 single mutant plants was not significantly different from that in WT plants, with 90% of cells (36/40) accumulating YFP–POK1 at the division site only and 10% (4/40) accumulating YFP–POK1.
telophase cells (Figure 1I, n = 12/27), which is a significantly greater midline accumulation rate compared to that in WT plants (13%, n = 4/31, 10 plants, Fisher’s exact test, \( P = 0.0094 \)) or the air9 single mutant (Figure 1L, 10%, n = 4/40). Aberrant phragmoplast midline accumulation of YFP–POK1 in the tan1 single mutant suggested that the POK1–TAN1 interaction might be required to maintain POK1 at the division site. This prompted us to examine their interaction more closely.

Amino acids 1–132 of TAN1 rescue the tan1 air9 double mutant

POK1 interacts with both full-length TAN1 and the first 132 amino acids of TAN1 using the yeast two-hybrid system (Rasmussen et al., 2011). TAN1 missing the first 126 amino acids failed to rescue the tan1 air9 double mutant, suggesting that this part of the protein is critical for TAN1 function (Mir et al., 2018). To test the function of this region of the protein in Arabidopsis, the TAN1 coding sequence for the first 132 amino acids was fused to YFP (TAN11–132-YFP) driven by the cauliflower mosaic p35S promoter and was then transformed into the tan1 air9 double mutant. We used p35S:TAN1-YFP in the tan1 air9 double mutant as our benchmark for rescue, as its ability to rescue the tan1 air9 double mutant was demonstrated previously (Mir et al., 2018). The progeny of several independent p35S:TAN11–132-YFP lines rescued the tan1 air9 double mutant, as described in more detail below. Overall root patterning of the p35S:TAN11–132-YFP or full-length p35S:TAN1-YFP was restored, while untransformed tan1 air9 double mutant roots had misoriented divisions (Figure 2A; Supplemental Figure S1). Cell file rotation, which skews left and has large variance in the tan1 air9 double mutant (Figure 2, B and C), was significantly rescued in both p35S:TAN11–132-YFP and p35S:TAN1-YFP tan1 air9 lines (n = 37 and 41 plants, respectively), compared to the untransformed tan1 air9 control (Levene’s test used due to nonnormal distribution, \( P < 0.0001 \)). Root length at 8 days after stratification was also restored (Figure 2D). Interestingly, although TAN11–132-YFP rarely co-localizes with PPBs in WT plants (Rasmussen et al., 2011) or in the tan1 air9 double mutant (10%, n = 9/89 cells, Figure 3A), PPB angles in untransformed tan1 air99 plants had significantly less variance compared to the untransformed control (Figure 2E). Phragmoplast positioning defects of the tan1 air9 double mutant were also significantly rescued by p35S:TAN11–132-YFP. Altogether, p35S:TAN11–132-YFP rescued the phenotypes of the double mutant similar to full-length p35S:TAN1-YFP. This indicates that most functions that affect phenotypes assessed here are encoded by the first section of the TAN1 gene.

Disrupting TAN1–POK1 interaction alters TAN1 and POK1 localization to the division site and reduces tan1 air9 rescue

To further understand how TAN1 functions, we disrupted its ability to interact with the kinesin POK1 using alanine scanning mutagenesis. Alanine scanning mutagenesis was used to replace six amino acids with six alanines across the first approximately 120 amino acids of TAN11–132 (described in the “Materials and methods”). After testing their interaction with POK1 using the yeast two-hybrid system, we identified seven constructs that lost interaction with POK1 (Supplemental Figure S2). Reasoning that highly conserved amino acids would be more likely to play critical roles in the TAN1–POK1 interaction, we selected TAN1(28–33A)1–132 with alanine substitutions, replacing the highly conserved amino acids at positions 28–33 (INKVDK) with six alanines (TAN1(28–33A)1–132) for analysis in Arabidopsis. Our hypothesis was that the mutated form of TAN11–132 (TAN1(28–33A)1–132) would not rescue the tan1 air9 mutant due to lack of POK1 and TAN1 interaction. TAN1(28–33A)1–132 was cloned into a plant transformation vector to generate p35S:TAN1(28–33A)1–132-YFP and transformed into the tan1 air9 double mutant. The p35S:TAN1(28–33A)1–132-YFP construct partially rescued the tan1 air9 double mutant (Figure 4; Supplemental Figure S3). p35S:TAN1(28–33A)1–132-YFP in the tan1 air9 double mutant did not rescue cell file rotation defects (Figure 4, B and D) or phragmoplast angle defects (Figure 4F). However, overall plant growth (Figure 4C) and root length (Figure 4E) showed intermediate rescue compared to unaltered p35S:TAN11–132-YFP in the tan1 air9 double mutant. PPB angles in tan1 air9 double mutants expressing either p35S:TAN1(28–33A)1–132-YFP or p35S:TAN11–132-YFP were similar, suggesting that the TAN1–POK1 interaction may not be required for PPB placement (Figure 4F). These results suggest that the first 132 amino acids of TAN1 perform several vital functions, some of which are contingent or partially contingent on a likely interaction with POK1 in Arabidopsis.

To understand how this mutation within TAN11–132 affected localization, we analyzed TAN1(28–33A)1–132-YFP in the tan1 air9 double mutant. Localization of TAN1(28–33A)1–132-YFP to the division site in the tan1 air9 double mutant was significantly reduced compared to that of unaltered TAN11–132-YFP, which localized to the division site during telophase 100% of the time (n = 58/58 cells, 29 plants, Figure 3E; Rasmussen et al., 2011). TAN1(28–33A)1–132-YFP showed no obvious division site localization 68% of the

Figure 1 (Continued)

at the division site and in the phragmoplast midline. In tan1 single mutant plants, YFP–POK1 was present at the division site in all telophase cells but accumulated more frequently at the division site and phragmoplast midline compared to in WT plants (44%; 12/27 cells; \( P = 0.0094 \)). In tan1 air9 double mutant plants, YFP–POK1 never accumulated at the division site of telophase cells and instead accumulated solely in the phragmoplast midline (34/44 cells, \( P < 0.00001 \)).
time ($n = 15/22$ cells, Figure 3I) or faint division site accumulation in $32\%$ of telophase cells ($n = 7/22$ cells, Figure 3J).

When the fluorescence intensity of TAN1(28–33A)1–132-YFP at the division site during telophase was compared to the cytosolic fluorescence intensity in the same cell, the median ratio was $\sim 1.1$, indicating little preferential accumulation of TAN1(28–33A)1–132-YFP at the division site (Figure 3K). In contrast, the median ratio of unaltered TAN11–132-YFP at the division site was $\sim 1.8$ compared to cytosolic fluorescence, indicating its preferential accumulation at the division site. This suggests that TAN1 requires the motif in amino acids 28–33 to localize properly to the division site during telophase. Our hypothesis is that this reduced localization is due to disruptions in the TAN11–132–POK1 interaction.

Figure 2 p35S:TAN11–132-YFP rescues Arabidopsis tan1 air9 double mutant phenotypes. A, Cell walls stained with PI of tan1 air9 double mutant root tips expressing p35S:TAN1-YFP (left) and p35S:TAN11–132-YFP (middle), and untransformed tan1 air9 double mutant root tips (right). Bars = 25 $\mu$m. B, Maximum projections of 10 1-$\mu$m Z-stacks of PI-stained differentiation zone root cell walls. Scale bars = 50 $\mu$m. C, Cell file rotation angles of tan1 air9 double mutants expressing p35S:TAN1-YFP (left), p35S:TAN11–132-YFP (middle), and untransformed plants (right), $n > 13$ plants for each genotype. Each dot represents an angle measured from the left side of the long axis of the root to the transverse cell wall. Angle variances were compared with Levene's test due to nonnormal distribution. D, Root length measurements from 8 days after stratification of tan1 air9 double mutants expressing p35S:TAN1-YFP (left), p35S:TAN11–132-YFP (middle), and untransformed plants (right), $n > 28$ plants for each genotype, compared by two-tailed $t$ test with Welch's correction. E, PPB and phragmoplast angle measurements in tan1 air9 double mutant cells expressing p35S:TAN1-YFP (left), p35S:TAN11–132-YFP (middle), and untransformed plants (right), $n > 20$ plants for each genotype. Angle variances compared with $F$-test. C–E, Mean and standard deviation are indicated. ns indicates not significant, *$P < 0.01$, **$P < 0.0001$.
Next, we generated a construct that introduced alanines at amino acids 28–33 in full-length YFP–TAN1 constructs (p35S:YFP–TAN1(28–33A)) to assess whether p35S:YFP–TAN1(28–33A) would rescue the tan1 air9 double mutant. In contrast to the modest partial rescue provided by p35S:YFP–TAN1(28–33A)–YFP, full-length p35S:YFP–TAN1(28–33A) significantly rescued the defects in the tan1 air9 double mutant, as described below. First, we assessed whether full-length TAN1(28–33A) interacted with POK1 via the yeast two-hybrid system, and it did not (Supplemental Figure S4). Next, we analyzed rescue in Arabidopsis expressing p35S:YFP–TAN1(28–33A). Most defects, except phragmoplast angle variance (Figure 5; Supplemental Figure S5), were fully rescued in the p35S:YFP–TAN1(28–33A)–YFP, full-length p35S:YFP–TAN1(28–33A) tan1 air9 line, including cell file rotation (Figure 5C), root length (Figure 5D), and PPB angles (Figure 5E). Similar to TAN1–YFP, YFP–TAN1(28–33A) localized to the division site in prophase, prophase, and telophase (Supplemental Figure S6).

To determine if full-length YFP–TAN1(28–33A) had reduced accumulation at the division site during telophase similar to TAN1(28–33A)–YFP, fluorescence intensity levels were measured. During prophase, the YFP–TAN1(28–33A) fluorescence intensity at the division site compared to in the cytosol was comparable to the TAN1-YFP fluorescence intensity ratios. In contrast, the YFP–TAN1(28–33A) fluorescence intensity ratios during telophase were reduced to ~1.6 compared with unaltered TAN1-YFP (~2.1) indicating that YFP–TAN1(28–33A) accumulated less at the division site during telophase (Supplemental Figure S6). Together, these data suggest that TAN1 is recruited to the division site during prophase without an interaction with POK1. Defects in phragmoplast positioning may be due specifically to the disruption of the TAN1–POK1 interaction in the tan1 air9 double mutant. Since AIR9 is missing, TAN1 and POK1 interactions may be more important. Moreover, phragmoplast guidance defects may be due to the lower accumulation of TAN1 at the division site that would normally be mediated by POK1 during telophase.

To better understand how these alanine substitutions affect both POK1 and TAN1 localization, we examined tan1 air9 double mutant plants expressing a microtubule marker (UBQ10:Scarlet-MAP4; Pan et al., 2020), pTAN1:CFP–TAN1(28–33A) and pPOK1:YFP–POK1 (Lipka et al., 2014). Both CFP–TAN1(28–33A) and YFP–POK1 had reduced accumulation at the division site in the tan1 air9 double mutant. CFP–TAN1(28–33A) and YFP–POK1 colocalized with

Figure 3 Division site localization during telophase is common for TAN1(1–132)–YFP but rare for TAN1(28–33A)–YFP in tan1 air9 double mutant cells. A–E, PI-stained tan1 air9 plants expressing p35S:TAN1(1–132)–YFP and CFP–TUBULIN during mitosis (n = 29 plants). The division site is indicated by arrowheads in the YFP panels. Scale bars = 10 μm. A, Rare prophase division site localization of TAN1(1–132)–YFP (10%, n = 9/89 cells), (B) common prophase TAN1(1–132)–YFP nuclear accumulation without division site localization (90%, n = 80/89 cells), (C) no specific TAN1(1–132)–YFP division site accumulation in metaphase (100%, n = 28/28 cells), (D) faint TAN1(1–132)–YFP division site accumulation accompanied by midline accumulation in late anaphase/early telophase (80%, n = 16/20 cells), and (E) TAN1(1–132)–YFP division site accumulation during telophase (100%, n = 58/58 cells). F–H, tan1 air9 plants expressing p35S:TAN1(28–33A)–YFP during mitosis (n = 13 plants). The division site is indicated by arrowheads in the YFP panels. F, No specific TAN1(28–33A)–YFP prophase division site accumulation during prophase (100%, n = 20/20 cells), (G) no specific TAN1(28–33A)–YFP division site accumulation during metaphase (100%, n = 12/12 cells), (H) no TAN1(28–33A)–YFP division site or midline accumulation in late anaphase/early telophase (100%, n = 8/8 cells), (I) no specific TAN1(28–33A)–YFP division site accumulation during telophase (68%, n = 15/22 cells) and (J) faint TAN1(28–33A)–YFP division site accumulation during telophase (32%, n = 7/22 cells). K, Ratio of TAN1(1–132)–YFP (left) or TAN1(28–33A)–YFP (right) fluorescence at the division site to cytosolic fluorescence from tan1 air9 plants expressing p35S:TAN1(1–132)–YFP or p35S:TAN1(28–33A)–YFP during telophase, n > 23 plants for each genotype. Mean and standard deviation are indicated. Asterisks indicate a significant difference as determined by Mann–Whitney U test, P < 0.0001.
the PPB in 41% of cells (n = 32/79), which is significantly less frequent when compared to 72% of cells (n = 58/82 cells, 20 plants) with PPBs in the tan1 air9 mutant containing unaltered CFP–TAN1 and YFP–POK1 (Figure 6, A and I; Fisher’s exact test, \(P = 0.0001\), similar to Lipka et al. (2014) and Schaefer et al., 2017). Unaltered CFP–TAN1 fully rescued the tan1 air9 double mutant (Mills and Rasmussen, 2022), and served here as a control. Unaltered CFP–TAN1 and YFP–POK1 localized and were maintained at the division site similar to in the WT during metaphase (Figure 6B, n = 13/13), while CFP–TAN1(28–33A) and YFP–POK1 in the tan1 air9 mutant were sometimes absent from the division site during metaphase with only 58% of metaphase cells maintaining both proteins at the division site (n = 11/19 cells, Figure 6, F and I). During early telophase, unaltered CFP–TAN1 and YFP–POK1 were always at the division site (n = 14/14, Figure 6C), but CFP–TAN1(28–33A) and YFP–POK1 were maintained at the division site in only 65% of early telophase cells (n = 20/31 cells, Figure 6, G and I). Interestingly, YFP–POK1 accumulated in the phragmoplast midline in 26% of early telophase cells (n = 8/31 cells, Figure 6I) but was not observed in the phragmoplast midline in early telophase cells of plants expressing unaltered CFP–TAN1 (n = 0/14 cells, Figure 6I). During late telophase, when the phragmoplast has contacted the cell cortex in at least one location, CFP–TAN1 and POK1 always localized to the division site (100%, n = 63/63 cells, Figure 6D). Interestingly, although not observed in earlier stages, YFP–POK1 and CFP–TAN1(28–33A) recruitment to the division site increased to 90% of late telophase cells.
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(n = 53/59 cells, Figure 6H). In the remaining cells, neither CFP–TAN1(28–33) nor YFP–POK1 localized to the division site (3%, n = 2/59), or only CFP–TAN1(28–33) accumulated at the division site (7%, n = 4/59 cells). Together, these data suggest that TAN1–POK1 interactions play a critical role in stabilizing them together at the division site. Additionally, it suggests that other, yet unidentified proteins may recruit both TAN1 and POK1 to the division site, particularly during late telophase, in the absence of both AIR9 and the TAN1–POK1 interaction.

Discussion

In tan1 and air9 single mutants, POK1 localizes to the division site and there are no discernable division plane defects (Model in Supplemental Figure S7). However, in the tan1 air9 double mutant, POK1 co-localizes with the PPB but is...
and absent at the division site and phragmoplast midline in a portion of early telophase cells (35%; 11/31 cells; 12/31 cells, 0.001). In early telophase, CFP–TAN1(28–33A) colocalized with the PPB. F, YFP–POK1 and CFP–TAN1(28–33A) were maintained at the division site in most metaphase cells. CFP–TAN1(28–33A) was faint at the division site. G, Both YFP–POK1 and CFP–TAN1(28–33A) were observed at the division site in most early telophase cells. H, YFP–POK1 and CFP–TAN1(28–33A) were recruited to the division site in late telophase. I, Subcellular localization of CFP–TAN1 and CFP–TAN1(28–33A) in tan1 air9 double mutant cells during preprophase/prophase, metaphase, early telophase (telophase cells where the phragmoplast has not yet contacted the cell cortex), and late telophase. N > 19 plants for each genotype. Statistically significant differences were determined using Fisher’s exact test. Asterisks indicate significant differences in POK1 and TAN1(28–33A) localization to the division site only. CFP–TAN1 and YFP–POK1 colocalized with the PPB more frequently (72%, 59/82 cells) than CFP–TAN1(28–33A) and YFP–POK1 (41%; 32/79 cells; 0.001). tan1 air9 plants expressing CFP–TAN1(28–33A) also had more PBPs that only accumulated YFP–POK1 (16%; 13/79 cells; 0.0461) compared to CFP–TAN1-expressing plants (6%, 5/82). In metaphase cells, CFP–TAN1 and YFP–POK1 were maintained at the division site in 100% of observed cells (13/13), compared to CFP–TAN1(28–33A) and YFP–POK1 which accumulated at the division site in 58% of cells (11/19, 0.0104). In early telophase, TAN1(28–33A) and YFP–POK1 accumulation at the division site was reduced (39%; 12/31 cells; 0.0001) compared to CFP–TAN1 and YFP–POK1 accumulation (100%, 14/14 cells). CFP–TAN1(28–33A)-expressing plants also accumulated both CFP–TAN1 and YFP–POK1 at the division site and YFP–POK1 in the phragmoplast midline more frequently in early telophase cells compared to CFP–TAN1 expressing plants (26%; 8/31 cells; 0.0436). CFP–TAN1(28–33A) and YFP–POK1 were also absent at the division site and phragmoplast midline in a portion of early telophase cells (35%; 11/31 cells; 0.098), compared to CFP–TAN1 expressing plants (0/14 cells). CFP–TAN1(28–33A) and YFP–POK1 accumulated at the division site in 71% of late telophase cells (42/59 cells) compared to CFP–TAN1 and YFP–POK1 (60/63, 0.0004). Nineteen percent of late telophase cells accumulated both CFP–TAN1(28–33A) and YFP–POK1 at the division site and YFP–POK1 in the phragmoplast midline (11/59 cells) more frequently compared to CFP–TAN1-expressing plants (3/63 cells; 0.022). In some late telophase cells of TAN1(28–33A) expressing plants CFP–TAN1 (28–33A) accumulated at the division site alone (7%; 4/59 cells; 0.0518) or TAN1(28–33A) and YFP–POK1 failed to accumulate at the division site (3%; 2/59 cells; 0.2318) compared to CFP–TAN1-expressing plants where neither localization pattern was observed in late telophase (0/63 cells). Lost from the division site during metaphase (Model in Figure 7). First, this suggests that TAN1 and AIR9 are not essential for POK1 co-localization with the PPB. Second, it suggests that POK1 is maintained at the division site after PPB disassembly via direct or indirect interactions with TAN1 or AIR9. We provide evidence that TAN1 interacts with POK1 through motifs within the first 132 amino acids of TAN1, as identified using the yeast two-hybrid system. Alignments of TAN1 proteins from representative monocots and dicots, such as Solanum lycopersicum, Oryza sativa, Sorghum bicolor, Zea mays, and Brassica napus, showed that amino acids 28–33 (INKVDK) are highly conserved across plant species (Supplemental Figure S8). Amino acids 30–32 (VDK) are identical and the remaining residues within the motif have similar properties across these plant species. The high degree of conservation suggests that these amino acids are likely important for TAN1 function. When alanine substitutions of these amino acids were introduced into TAN1 and transformed into the Arabidopsis tan1 air9 double mutant, we observed reduced TAN1 and POK1 localization at the division site, as well as defects in phragmoplast positioning. Here, we hypothesize that amino acids 28–33 are essential for TAN1 and POK1 interaction in both the yeast two-hybrid system and in Arabidopsis. In addition to several reports showing that TAN1 and POK1 interact using the yeast two-hybrid system (Mueller et al., 2006; Rasmussen et al., 2011), bimolecular fluorescence complementation has also been used to show TAN1–POK1 interactions in Arabidopsis protoplasts (Lipka et al., 2014). Alanine substitutions at positions 28–33 of TAN1 may disrupt TAN1–POK1 interactions through misfolding that blocks the POK1 interaction site or by affecting the amino acids that directly...
mediate POK1 binding. Regardless of the exact mechanism(s) of POK1–TAN1 physical interactions or the possibility that yeast two-hybrid interactions do not reflect equivalent POK1–TAN1 physical interactions in Arabidopsis, we show that these TAN1 amino acids are involved in mediating TAN1 and POK1 localization to the division site.

We demonstrate that the first region of the TAN1 protein, the first 132 amino acids that primarily accumulate at the division site during telophase (Rasmussen et al., 2011), is both necessary (Mir et al., 2018) and sufficient to largely rescue the tan1 air9 double mutant (Figure 2). This suggests that TAN1_{1-132} and its recruitment to the division site during telophase is critical for correct division plane orientation in the tan1 air9 double mutant. Although full length TAN1 localizes to the division site throughout cell division, the ability of TAN1(28–33A) to rescue the tan1 air9 double mutant suggests that TAN1, and possibly POK1, localization to the PPB and division site during metaphase may not be required.
for division site maintenance in Arabidopsis. Indeed, whether the PPB itself is required for division plane positioning has been raised by analysis of a triple mutant in three closely related genes (TONNEAU RECRUITING MOTIF6 (TRM6), TRM7, and TRM8). The trm678 mutant, which lacks well-defined PPBs, has partially disrupted POK1 recruitment to the division site but only minor defects in division positioning (Schaefer et al., 2017). However, when amino acids critical for TAN1–POK1 interactions in the yeast two-hybrid system are disrupted by transforming TAN1(28–33A)1–132–YFP into the tan1 air9 double mutant, root growth and phragmoplast positioning are disrupted. TAN1(28–33A)1–132–YFP accumulation at the division site during telophase was reduced compared to unaltered TAN11–132–YFP. This suggests that TAN1–POK1 interaction promotes, but is not strictly necessary, for TAN1 recruitment to the division site during telophase.

Full-length TAN1(28–33A) localizes to the division site throughout cell division and almost fully rescues the tan1 air9 double mutant. TAN1, AIR9, and POK1 colocalize at the PPB independently of one another, which may promote the formation of protein complexes required for division site maintenance. Colocalizing with the PPB may provide an opportunity for nearby proteins to form stabilizing interactions before PPB disassembly. This suggests that recruitment of TAN1 and POK1 to the division site early in cell division may provide another temporally distinct way to promote correct division plane positioning. Phragmoplast positioning defects in TAN1(28–33A)1 tan1 air9 plants may be the result of defects in phragmoplast guidance in cells that lacked TAN1(28–33A) and POK1 at the division site in metaphase or early telophase that were not corrected in late telophase.

The ability of TAN1(28–33A) and POK1 to remain at the division site in some cells after PPB disassembly in the tan1 air9 double mutant suggests that there are other proteins that interact with TAN1 and/or POK1 that help stabilize them at the division site perhaps via the formation of multi-protein complexes. The pleckstrin homology GAPs, PHGAP1, and PHGAP2 (Stöckle et al., 2016); RANGAP1 (Xu et al., 2008); and IQ67 DOMAIN (IQD)6,7,8 proteins (Kumari et al., 2021) are division site-localized proteins that may stabilize TAN1 and POK1 at the division site via their interaction with POK1. PHGAP1, PHGAP2, and RANGAP1 are dependent on POK1 and POK2 for division site recruitment. Like TAN1, RANGAP1 colocalizes with the PPB and remains at the division site throughout cell division (Xu et al., 2008). PHGAP1 and PHGAP2 are uniformly distributed in the cytoplasm and on the plasma membrane in interphase cells and accumulate at the division site during metaphase. These proteins also have their own distinct roles in division site maintenance (Stöckle et al., 2016). PHGAP2 has a likely role in division site establishment by regulating ROP activity (Hwang et al., 2008). RANGAP1 regulation of local RAN-GTP levels has potential roles in microtubule organization and division site identity (Xu et al., 2008). IQD6, IQD7, and IQD8 interact with POK1 and play a role in PPB formation and POK1 recruitment to the division site. The IQD6/78 triple mutant has PPB formation defects and fails to recruit POK1 to the division site in cells lacking PPBs. However, POK1 localization to the division site in the IQD6/78 mutant recovers during telophase to WT levels (Kumari et al., 2021). We speculate that this IQD6–8 independent recruitment may depend on TAN1. Unlike the PHGAPs and RANGAP1, IQD8 localization to the division site is not dependent on POK1 and POK2. This suggests that IQD6–8 proteins work upstream of POK1 to establish the division site and are important for POK1 recruitment to the division site early in cell division. Although TAN1–POK1 interactions become critical for TAN1 and POK1 maintenance at the division site in the absence of AIR9, other division site-localized proteins may provide additional stability and help maintain TAN1 and POK1 at the division site.

How AIR9 stabilizes POK1 at the division site in the absence of TAN1 is less clear. There is no information about whether POK1 and AIR9 interact directly with one another. Additionally, AIR9 localization, in contrast to TAN1 localization, is intermittent at the division site. When expressed in tobacco (Nicotiana tabacum) Bright Yellow2 cells, AIR9 colocalizes with the PPB but is then lost from the division site until late telophase when the phragmoplast contacts the cortex (Buschmann et al., 2006). In Arabidopsis, AIR9 may localize to the division site during metaphase or telophase, but it is difficult to observe because AIR9 also strongly colocalizes with cortical microtubules, which may obscure AIR9 localization in nearby cells (Buschmann et al., 2015). Rather than directly interacting with POK1, AIR9 may recruit other proteins to the division site during preprophase that help maintain POK1 at the division site in the absence of TAN1. One potential candidate is the kinesin-like calmodulin binding protein, KCBP, which interacts with AIR9 (Buschmann et al., 2015). KCBP is a minus-end-directed kinesin (Song et al., 1997) that localizes to the division site in Arabidopsis and moss (Miki et al., 2014; Buschmann et al., 2015). We speculate that other TAN1, AIR9, and POK1 interacting proteins that have not been identified yet may be key for TAN1–POK1 division site maintenance.

POK1 and POK2 have roles in phragmoplast guidance, but POK2 also has a role in phragmoplast dynamics (Lipka et al., 2014; Herrmann et al., 2018). Although POK1 does not frequently accumulate in the phragmoplast midline in WT cells, POK2 showed striking dual localization to both the phragmoplast midline and the division site. Localization of POK2 to the phragmoplast midline required the N-terminal motor domain, while the C-terminal region was localized to the division site (Herrmann et al., 2018). Our hypothesis is that the “default” location of both POK1 and POK2 is at microtubule plus ends at the phragmoplast midline, based on likely or confirmed plus-end-directed motor activity (Chugh et al., 2018). Interactions with division site-localized proteins, such as TAN1 and AIR9, may stabilize or recruit
POK1 and POK2 at the division site away from the phragmoplast midline.

We demonstrate that AIR9 and TAN1 function redundantly to maintain POK1 at the division site to ensure correct cell wall placement. In the absence of AIR9, our data suggest that TAN1–POK1 interaction promotes, but is not required for, the maintenance of both proteins at the division site, and disrupting this interaction partially disrupts their localization to the division site. This also suggests that other TAN1, POK1, and AIR9 interacting proteins are involved with stabilizing TAN1 and POK1 at the division site.

Materials and methods

Growth conditions, genotyping mutants, and root length measurements

Arabidopsis (A. thaliana) seedlings were grown on 1/2 strength Murashige and Skoog (MS) media (MP Biomedicals; Murashige and Skoog, 1962) containing 0.5 g L⁻¹ morpholineethanesulfonic acid monohydrate (MES-monohydrate, Fisher Scientific, Waltham, MA, USA, CAS 145224-94-8), pH 5.7, and 0.8% agar (Fisher Scientific). Seeds sown on plates were first stratified in the dark at 4°C for 2–5 days, then grown vertically in a growth chamber (Percival) with 16-h light cycles. Root length experiments, tan1 air9 CFP-TUBULIN plants were scanned (Epson) and root lengths were measured using FIJI (ImageJ, http://fiji.sc/) after 8 days. Untransformed tan1 air9 double mutant and air9 single mutant seeds were grown alongside the double mutant seeds expressing the TAN1 constructs in equal numbers on the same plates to ensure plants were grown under the same conditions. After plates were scanned, seedlings were screened by confocal microscopy to identify seedlings expressing YFP translational fusion TAN1 constructs and CFP-TUBULIN, if present in the transgenic lines. At least three biological replicates, grown on separate plates on separate days, and at least 28 plants of each genotype across all replicates were analyzed for each root growth experiment. Welch’s t tests were used to identify whether there were statistically significant differences between replicates before pooling the replicates for analysis. Root lengths were then plotted using Prism (GraphPad). Statistical analysis of root length was performed with Prism (GraphPad) using t tests with Welch’s correction. Welch’s t test (unequal variance t test) is used to test the hypothesis that two populations have equal means. Unlike the Student’s t test, Welch’s t test is often used when two samples have unequal variances or sample sizes. This test was used due to the unequal sample sizes because the plants examined were often segregating for multiple transgenes and had lower sample sizes than control plants such as the air9 single mutant and tan1 air9 double mutant, which either lacked transgenes or were segregating fewer transgenes.

YFP translational fusion TAN1 constructs were analyzed in csh-tan (TAN1, AT3G05330) air9-31 (AIR9, AT2G34680) double mutants in Landsberg erecta (Ler) unless otherwise specified. The pPOK1:YFP–POK1 transgene in Columbia, a kind gift from Sabine Müller (Lipka et al., 2014), was crossed into the tan-mad and air9-5 Columbia/Wassilewskija double mutant previously described (Mir et al., 2018). tan-mad and air9-5 mutants were genotyped with primers ATLP and ATLP (to identify WT TAN1), JL202 and ATLP (to identify T-DNA insertion in TAN1), AIR9-SRP and AIR9-5LP (to identify WT AIR9), and LBb1.3 and AIR9RP (to identify T-DNA insertion in AIR9) and by observation of the tan1 air9 double mutant phenotype (Supplemental Table S1).

Generation of transgenic lines

Agrobacterium tumefaciens-mediated floral dip transformation was used as described (Clough and Bent, 1999). csh-tan air9-31 double mutants were used for all floral dip transformations unless otherwise specified. Transgenic plants were selected on 15 μg mL⁻¹ glufosinate (Finale; Bayer) and screened by microscopy before being transferred to soil and selfed. CFP-TUBULIN was crossed into p35S:TAN1(28–33A)1–132-YFP tan1 air9 plants using tan1 air9 CFP-CFP-TUBULIN plants (Mir et al., 2018), and progeny were subsequently screened by microscopy for CFP and YFP signal. csh-tan1 air9-31 double mutants were confirmed by genotyping with primers ATLP and AtTAN 733-CDS Rw (to identify WT TAN1), AtTAN 733-CDS Rw and Ds5-4 (to identify T-DNA insertion in TAN1), AIR9_cDNA 2230 F and AIR9 gnm7511 R (to identify WT AIR9), and AIR9 gnm7511 R and Ds5-4 (to identify T-DNA insertion in AIR9).

Columbia expressing the microtubule marker UBQ10:mScarlet-MAP4 (Pan et al., 2020), a kind gift from Xue Pan and Zhenbiao Yang (UCR), was crossed to tan-mad and air9-5 Columbia/Wassilewskija double mutants expressing pPOK1:YFP–POK1. Progeny were screened for mScarlet-MAP4 and YFP–POK1 by confocal microscopy and then selfed to recover air9-5 single mutants, tan-mad single mutants, and air9-5 tan-mad double mutants expressing mScarlet-MAP4 and YFP–POK1.

pTAN1:CFP–TAN1 and pTAN1:CFP–TAN1(28–33A) were introduced into air9-5 tan-mad double mutants expressing mScarlet-MAP4 and YFP–POK1 by A. tumefaciens-mediated floral dip transformation. pTAN1:CFP–TAN1 and pTAN1:CFP–TAN1(28–33A) transformants were selected on 100 μg mL⁻¹ gentamicin (Fisher Scientific) and the presence of mScarlet-MAP4, YFP–POK1, and either CFP–TAN1 or CFP–TAN1(28–33A) was confirmed by confocal microscopy. Four independent T2 transformed lines for pTAN1:CFP–TAN1(28–33A) and three independent T2 transformed lines for unaltered pTAN1:CFP–TAN1 were examined for division site localization cell counts.

Plasmid construction

TAN1_132-YFP coding sequences were subcloned by EcoRI and BamHI double digestion from the plasmid pEZRK-LNY-TAN1_132-YFP described previously (Rasmussen et al., 2011).
into the pEZT-NL vector (a kind gift from David Ehrhardt, Carnegie Institute, Stanford University) and selected with glufo sineate (Finale; Bayer). The CFP-TUBULIN (CFP-TUA6) vector was previously described, a kind gift from Viktor Kirik (Kirik et al., 2007).

Six amino acid alanine substitutions were generated using overlapping polymerase chain reaction (PCR) (primers in Supplemental Table S1) beginning at amino acid 10 of TAN1(28–33A). The TAN1(28–33A) YFP coding sequence from plasmids described previously was used as the PCR template (Rasmussen et al., 2011). TAN1(28–33A) was cloned into EcoRI–BamHI double digestion into pEZT-NL. To generate YFP–TAN1(28–33A), alanine substitutions were first introduced into G22672 (TAN1 cDNA in pENTR223, from the Arabidopsis Biological Resource Center) using overlapping PCR with the same primers to generate TAN1(28–33A). Gateway LR reaction (Fisher Scientific) was then used to subclone TAN1(28–33A) into pEarley104 (Earley et al., 2006).

pTAN–CFP–TAN1(28–33A) was generated using overlapping PCR. The TAN1 native promoter was amplified from NptATAN–YFP (Walker et al., 2007) using the primers NptTANSacFor and NptTANcereuleanRev. Cereulan was amplified from the Cereulan CDS in pDONR221P4r/P3r using the primers NptTANcereuleanFor and CereulanpEarleyRev. TAN1(28–33A) in pEarley104 was amplified using CereulanpEarleyFor and pEarleyOCSpStRev. TAN1 native promoter, Cereulan, and TAN1(28–33A) were then combined using overlapping PCR using NptTANSac1 and pEarleyOCSpStRev. pTAN–CFP–TAN1(28–33A) was then subcloned into pHA212G, a kind gift of Meng Chen (UCR), using Sac1 and Pst1 double digest. pTAN–CFP–TAN1 was generated the same way as pTAN–CFP–TAN1(28–33A) except unaltered TAN1 in pEarley104 was amplified using CereulanpEarleyFor and pEarleyOCSpStRev.

Microscopy

An inverted Ti Eclipse (Nikon, Tokyo, Japan) with motorized stage (ASJ Piezo) and spinning-disk confocal microscope (Yokogawa W1) built by Solamere Technology was used with Micromanagement software (micromanagement.org). Solid-state lasers (Obis) and emission filters (Chroma Technology) were used. For CFP translational fusions, an excitation of 445 and emission of 480/40 were used; for YFP translational fusions, an excitation of 514 and emission of 540/30 were used; and for propidium iodide (PI), Alexa-568 goat anti-mouse antibody, and mScarlet-MAP4, an excitation of 561 and emission of 620/60. A 20× objective with a 0.75 numerical aperture and a 60× objective with a 1.2 numerical aperture were used with perfluorocarbon immersion liquid (RIAAA-6788 Cargille). Excitation spectra for mScarlet-MAP4 and YFP–POK1 partially overlapped, which resulted in a faint bleed-through signal in the YFP channel for some dense microtubule structures (e.g. spindles and phragmoplasts). YFP–POK1 colocalization with PPBs was carefully determined based on distinct YFP–POK1 signal and the presence of cytosolic YFP–POK1.

The ratio of the division site versus cytosolic fluorescence intensity was determined by taking the median YFP fluorescence intensity from the center Z-stack of individual cells with PPBs or phragmoplasts. For each cell, the median fluorescence intensity was measured for two cytosolic areas and the division site on each side of the cell using circles with areas of 0.875 μm². The sum of the median intensity at the division site on each side was then divided by the sum of the median intensity of the two cytosolic areas to calculate the ratio of the division site versus cytosolic fluorescence intensity. Fluorescence intensities were measured in FIJI. All plants used for this analysis were grown on the same day and imaged using identical conditions, and at least five plants of each genotype were examined.

Measurements of PPB and phragmoplast angles and cell file rotation

At least three biological replicates, grown on separate plates on separate days, composed of at least 15 plants per genotype for PPB measurements and at least eight plants per genotype for phragmoplast measurements were used to gather angle data. Eight-day-old seedlings were stained with 10-μM PI for 1 min and then destained in distilled water before imaging by confocal microscopy using a 20× or 60× objective. PPB and phragmoplast angles were measured using FIJI. The angle was measured between the left-hand cell wall and the orientation of the PPB or phragmoplast in the root tips of tan1 air9 double mutant plants expressing CFP-TUBULIN or immunostained microtubules (described in the next section). Cell file rotation was examined by measuring from the left-hand side of the transverse cell wall relative to the long axis of the root in images of the differentiation zone stained with PI. The differentiation zone was identified by the presence of root hairs. Prism (GraphPad) and Excel (Microsoft Office) were used to perform statistical analyses and to plot data. F-tests were used to compare normally distributed variances (PPB and phragmoplast angles) and Levene’s tests were used to compare nonnormally distributed variances (cell file rotation angle measurements). The tan1 air9 double mutant has nonnormally distributed cell file twisting because the roots tend to twist to the left (Mir et al., 2018). Genotypes across biological replicates were compared to ensure there were no statistically significant differences between them before pooling data.

Immunostaining

air9, tan1 air9 p35S–TAN1–YFP, tan1 air9 p35S–YFP–TAN1(28–33A), and untransformed tan1 air9 plants were stratified and then grown vertically on 1/2 MS plates in a growth chamber at 22°C with a 16-h/8-h light/dark cycle for 8 days. The seedlings were screened by microscopy for YFP and then fixed and processed for immunofluorescence microscopy using a 1:2,000 dilution of monoclonal anti-tubulin B-5-1-2 antibody (Life Technologies, Carlsbad, CA, USA; 32–2500) followed by 1:2000 dilution of Alexa-568 goat anti-mouse antibody (Thermo Fisher, Waltham, MA,
USA; A-11004) as described previously (Sugimoto et al., 2000).

Yeast two-hybrid
Six alanine substitutions were generated using the overlapping PCR and TAN1 coding sequence in pEZRK-LNY-TAN11–132-YFP as a template beginning at amino acid 10 of TAN1 and continuing through to amino acid 123 according to Russell and Sambrook (2001). All amino acids except substitutions for 64–69 and 106–111 were cloned into the pAS vector (Fan et al., 1997) using EcoRI–BamHI double digestion. pBD-TAN1(28–33A) was generated using primers Ala_05_FOR and Ala_05_REV to perform DpnI-mediated site-directed mutagenesis by PCR (Fisher and Pei, 1997). pBD-TAN1 (Walker et al., 2007) and pAS-TAN11–132 (Rasmussen et al., 2011) were used as positive controls, while pAD-MUT was used as a negative control for testing interactions with pAD-POK1 (Müller et al., 2006). pAD-POK1 and pAS-TAN11–132 constructs were co-transformed into yeast strain YRG2 according to manufacturer instructions (Stratagene). A positive yeast two-hybrid interaction was determined by the presence of growth on plates cultured at 30°C lacking histidine after 3 days. Plates were then scanned (Epson, Nagano, Japan).

Accession numbers
TAN1: AT3G05330, AIR9: AT2G34680, and POK1: AT3G17360

Supplemental data
The following materials are available in the online version of this article.

Supplemental Figure S1. p35S:TAN11–132*YFP tan1 air9 lines show significant rescue compared to untransformed tan1 air9 double mutants

Supplemental Figure S2. Yeast-two-hybrid interactions between POK1 (C-terminal amino acids 1,683–2,066, as previously described; Müller et al., 2006; Rasmussen et al., 2011; Lipka et al., 2014) and TAN11–132 alanine scanning constructs.

Supplemental Figure S3. p35S:TAN1(28–33A)1–132*YFP tan1 air9 lines show variable and incomplete rescue compared to unaltered p35S:TAN11–132*YFP tan1 air9.

Supplemental Figure S4. Yeast-two-hybrid interactions between TAN1 and POK1 (C-terminal amino acids 1,683–2,066, as previously described; Müller et al., 2006; Rasmussen et al., 2011; Lipka et al., 2014) and TAN1(28–33A).

Supplemental Figure S5. p35S:YFP–TAN1(28–33A) tan1 air9 lines show significant rescue compared to untransformed tan1 air9, but less accumulation of YFP–TAN1(28–33A) during telophase.

Supplemental Figure S6. YFP–TAN1(28–33A) localizes to the division site in preprophase or prophase and with reduced fluorescence during telophase in tan1 air9 mutants.

Supplemental Figure S7. A model of POK1 localization in tan1 and air9 single mutants.

Supplemental Figure S8. Alignments of amino acids 1–55 of A. thaliana TAN1 with TAN1 homologs from other plant species.

Supplemental Table S1. Primers used for cloning and genotyping.

Supplemental Data Set 1. Statistical analysis of tables.

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Conflict of interest statement. None declared.

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